NURSE CARE MANAGER MODEL OF OFFICE BASED ADDICTION TREATMENT:
CLINICAL GUIDELINES

A COLLABORATIVE CARE APPROACH

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Updated: March 9, 2018
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ACKNOWLEDGMENTS

This manual represents an update to the prior version of Boston Medical Center’s Office Based Addiction Treatment clinical guidelines. It was developed, with permission, from two pre-existing manuals on office based addiction treatment:


DISCLAIMER

Boston Medical Center is pleased to share its Office Based Addiction Treatment Clinical Guidelines with other providers. Although Boston Medical Center has attempted to confirm the accuracy of the information contained in these documents, this information is not a substitute for informed medical decision making by an appropriate, licensed provider. Clinicians must confirm the appropriateness of all treatment that they provide to a patient and are responsible for the health care decisions they make when caring for patients. If clinicians believe that any information included in these guidelines should be revised or clarified, please contact Boston Medical Center at 617-414-7453.

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SPONSORSHIP

This publication has been made possible by the grant support of the Massachusetts Department of Public Health Bureau of Substance Abuse Services (BSAS) to provide expansion of treatment with medication for addiction into community settings.
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INTRODUCTIONS

PURPOSE OF CLINICAL GUIDELINES

The purpose of these clinical guidelines is to provide detailed policies and protocols of the Office Based Addiction Treatment program for the use of buprenorphine (alone and in combination with naloxone) and naltrexone (oral and extended-release injectable formulations) in the treatment of substance use disorders.

These policies and protocols are meant to provide best practice guidelines to clinicians utilizing buprenorphine and/or naltrexone for the management of opioid use disorders and alcohol use disorders in mainstream medical practices, and to expand access to treatment.
INTRODUCTION TO OFFICE BASED ADDICTION TREATMENT

Federal data from the 2016 National Survey on Drug Use and Health indicates that 3.3 million people aged 12 or older in the United States (US) reported nonmedical use of prescription pain medication in the past month and 475,000 reported use of heroin in the past month. Largely driven by opioids, drug overdose is the leading cause of personal injury-related death in the US. Since 1999, the rate of overdose death involving any opioid has quadrupled. From 2000 to 2015 more than half a million people died from drug overdoses. In 2015, there were approximately 1.5 times more deaths in the US related to drug overdose than deaths related to motor vehicle accidents. In the same year, overdose death rates involving a synthetic opioid, such as fentanyl (not including methadone) increased by 72.2%; increased death rates attributed to synthetic opioids were seen across all demographics, regions and in numerous states. In 2016, approximately 20.1 million people in the US met criteria for a substance use disorder; however, only one in 10 of those individuals received any specialized care for their substance use disorder. This treatment gap has been attributed to numerous barriers, such as lack of patient and provider knowledge of evidence-based treatments, limited treatment capacity, stigma, and financial, legislative, and geographic obstacles.

Substance use disorders are a group of chronic medical conditions defined in the Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition (DSM-5) of the American Psychological Association that require long-term treatment and support. The US Food and Drug Administration (FDA) has approved three medications for the treatment of opioid use disorder: (i) oral methadone (full opioid agonist); (ii) oral transmucosal, injectable, and sub-dermal implant buprenorphine (nonselective partial opioid agonist); and (iii) oral formulation and long-acting injectable naltrexone (opioid antagonist). The most effective treatment for opioid use disorder involves medication maintenance for an adequate duration of time; the effectiveness of opioid agonist maintenance for treatment of opioid use disorder has been extensively documented through randomized clinical trials, quasi-experimental designs and program evaluations. There is evidence to support the use of injectable naltrexone to treat opioid use disorder, particularly in special populations, though, in general, treatment outcomes have been inferior to those attained with methadone and buprenorphine maintenance. At sufficiently high doses, opioid agonist maintenance treatment relieves craving for opioids. Continuous, steady-state medication maintenance treatment decreases the interaction between the opioid agonist medication (i.e. methadone or buprenorphine) and the μ-opioid receptors in the brain, blocking or attenuating the euphoric effects of illicit opioids (e.g. heroin).

Buprenorphine/naloxone was the first medication available to treat opioid dependence by prescription in a physician’s office or clinic outside of a traditional Outpatient Treatment Program (OTP). Prior to the advent of buprenorphine/naloxone, methadone was the only medication approved by the FDA to treat opioid use disorder in the US and, still today, can only
be dispensed at licensed methadone maintenance clinics. Unlike methadone, which is a full opioid agonist, buprenorphine is a μ-opioid receptor partial agonist. Due to a slow disassociation from the opioid receptor, the withdrawal syndrome from buprenorphine is milder when compared to that resulting from full opioid agonists (i.e. methadone). Naloxone, an opioid receptor antagonist, was added to buprenorphine to deter misuse (i.e. injection) and diversion. When administered sublingually, naloxone is poorly absorbed and has little to no pharmacological effects.\textsuperscript{15} Buprenorphine without naloxone (mono tablet) is typically only prescribed to women during pregnancy. Nationally, the number of patients receiving treatment with buprenorphine/naloxone has been increasing steadily, with good treatment retention. A recent evaluation of the federal buprenorphine waiver program (DATA 2000) found that of the 433 patients on buprenorphine maintenance interviewed, at a six-month follow-up, 60% were still retained in treatment and another 15% had completed treatment.\textsuperscript{16}

In 2016, over half of individuals aged 12 and older in the US reported drinking alcohol in the past 30 days; one in three reported heavy episodic alcohol use.\textsuperscript{1} In 2016, 15.1 million people in the US had an alcohol use disorder, comprising over 75% of people in the US with a substance use disorder. Alcohol use exists on a spectrum, beginning with abstinence and lower-risk drinking ranging all the way to severe alcohol use disorder or addiction. Typically the severity of consequences positively correlates with consumption. Unhealthy alcohol use is associated with risk of serious chronic health conditions (e.g. liver cirrhosis) as well as risks related to acute intoxication and alcohol withdrawal, such as accidental injury and death.\textsuperscript{17} Excessive alcohol use is the fourth leading cause of preventable death in the US;\textsuperscript{18} between 2006 and 2010, there were 88,000 alcohol-related deaths in the US.\textsuperscript{19}

Naltrexone is a competitive mu, kappa, and delta opioid receptor antagonist that blocks the effects of opioids by competitive binding. Naltrexone is available as an oral tablet that is taken daily, and an extended-release injectable formulation, administered intramuscularly into gluteal muscle every 28 days. The FDA approved the oral formulation of naltrexone for treatment of alcohol use disorder in 2006, and the extended-release version was approved in 2010 for treatment of both alcohol use disorder and opioid dependency following detoxification. The mechanism of action of naltrexone in alcohol use disorder is less clear, but is related to the blockage of opiate receptors related to the rewarding effects of alcohol use and craving.\textsuperscript{20}

Patients with substance use disorders should be offered medication treatment for addiction and psychosocial therapies as part of a comprehensive treatment plan. Like other chronic disease models, substance use disorders can be effectively managed in a primary care or community clinic by incorporating models of care like BMC’s Office Based Addiction Treatment program with nurse care managers, which increases access to evidence-based treatment.
INTRODUCTION TO THE NURSE CARE MANAGEMENT MODEL OF OBAT

In 2003, Boston Medical Center's Office Based Addiction Treatment (OBAT) program was created to address a key barrier to the integration of buprenorphine treatment identified by many office based settings: the lack of structured clinical support for providers. Nationally recognized and replicated, the BMC OBAT program utilizes a collaborative care model that relies on nurse care managers (NCM) to ensure delivery of high-quality addiction treatment while effectively and efficiently utilizing the time of providers waivered to prescribe buprenorphine. Within three years of its formation, BMC's OBAT model was expanded to 14 community health centers and the number of physicians waivered to prescribe buprenorphine in Massachusetts increased by 375%.

Dubbed the "Massachusetts Model," BMC's OBAT program serves as a model of service delivery for facilitating access to life-saving treatment and improving treatment outcomes in patients with opioid use disorders. It has been expanded to include treatments for other substance use disorders, including alcohol use disorder. Key to this model is the interprofessional collaboration between the nurse, provider and other members of the care team to provide comprehensive care during all phases of treatment, including: patient screening, assessment, education, care planning, medication induction, stabilization, and maintenance. Additionally, in this model, NCMs are responsible for ongoing medical management, coordination of follow-up care, telephone monitoring, relapse prevention, overdose education and support for patient self-management. By integrating treatment for substance use disorders within primary care, the substance use disorder can be treated similarly to other chronic medical conditions. Their scope of license and expertise in chronic disease management and patient education makes NCMs ideally suited to deliver ongoing care for substance use disorders. At the same time, nurse care management enables a physician, nurse practitioner, or physician's assistant to safely prescribe medications for addiction treatment to a greater number of patients, increasing overall access to these lifesaving medications.

BMC's OBAT program in adult primary care, our flagship clinic, is the largest office based opioid treatment program of its kind in New England, currently serving over 680 patients. The BMC OBAT NCM model is effective at increasing access to evidence-based treatment for addiction as well as improving the quality of care provided to patients with substance use disorders. Expansion and replication of innovative, evidence-based models, such as ours, are key to increasing access to addiction treatment across the US.

OBAT PHILOSOPHY
A substance use disorder is a chronic medical condition that responds best when treated with evidence-based, patient-centered comprehensive medical care. Patients engaged in OBAT deserve to be treated with dignity and respect. We believe that there should not be unnecessary obstacles to access addiction treatment in the OBAT program. The goal of OBAT is a cessation or reduction in harmful substance cravings and use, active participation and engagement in treatment, restoration of normal physiologic functions, and an improvement in one’s quality of life.
OBAT TEAM REQUIREMENTS

PROVIDERS

BUPRENORPHINE, BUPRENORPHINE/NALOXONE

QUALIFICATIONS: Qualified providers must obtain a waiver of authority to prescribe any medication that is a schedule III, IV, or V and FDA approved for the treatment of opioid dependence for the purpose of detoxification or maintenance treatment of patients with opioid dependence. With DATA 2000, physicians became legally qualified to receive waiver training. In July 2016, the Comprehensive Addiction and Recovery Act was signed into law increasing buprenorphine prescription authority to also include physician assistants (PA) and nurse practitioners (NP).

PHYSICIAN WAIVER ELIGIBILITY: To be eligible for a waiver, providers must have a current state medical license, a valid registration number from the US Drug Enforcement Agency (DEA), completion of eight hours of an approved waiver training course, and one or more of the following:

- Board subspecialty certification for addiction psychiatry (American Board of Medical Specialties), addiction (American Society of Addiction Medicine), or addiction medicine (American Osteopathic Academy of Addiction Medicine)
- OR—
- Participation as an investigator in one or more trials that led to the FDA approval of buprenorphine/naloxone or another schedule III-V narcotic medication used for the maintenance or detoxification treatment of opioid addiction
- OR—
- Other training or experience deemed equivalent by either the state Medical Board or by the Secretary of Health and Human Services (HHS).

NP/PA WAIVER ELIGIBILITY: To be eligible for a waiver, NPs and PAs must complete 24 hours of approved training that covers the following topics: opioid maintenance and detoxification; clinical use of all FDA-approved drugs for opioid use disorder treatment; patient assessment; treatment planning; psychosocial services; staff roles; and diversion control. NPs and PAs who are approved to prescribe buprenorphine must be supervised by or work in collaboration with a qualifying physician if required by law in their state.

REFERRALS: Providers must be able to refer patients to counseling and psychiatric services.
**PATIENT LIMITS:** Buprenorphine is a highly regulated medication with strict oversight. Please refer to [www.buprenorphine.samhsa.gov](http://www.buprenorphine.samhsa.gov) for updated information.

For the most current information available on buprenorphine prescribing and for waiver extension applications, contact:

SAMHSA’s Center for Substance Abuse Treatment (CSAT)
866-BUP-CSAT (866-287-2728)
infobuprenorphine@samhsa.hhs.gov
[www.buprenorphine.samhsa.gov](http://www.buprenorphine.samhsa.gov)

**NALTREXONE**

Naltrexone is not a schedule medication and therefore does not require a special licensure, certification, or waiver to prescribe. Any individual who is licensed to prescribe medication (physician, nurse practitioner or physician assistant) may prescribe and/or administer naltrexone. There is no limit to the number of patients that a provider could legally treat with naltrexone. However, when treating patients with substance use disorders, it is important that providers understand the nature of the underlying disorder, the pharmacological properties of available medications, and the importance of patient selection and monitoring.
NURSE CARE MANAGERS

- Registered Nurse (RN), Bachelor of Science in Nursing (BSN) and Certified Addictions Registered Nurse (CARN) preferred, licensed to practice nursing in the state where they are practicing.

- Recommended to have completed an initial training curriculum covering OBAT with buprenorphine and naltrexone, in-person or online.
  
  - Harvard Medical School Global Academy, Opioid Use Disorder Education Program: [https://globalacademy.hms.harvard.edu](https://globalacademy.hms.harvard.edu)
  
  - Providers Clinical Support System: [https://pcssnow.org/](https://pcssnow.org/)

- Attend ongoing professional education on topics relevant to treatment for substance use disorder (e.g. hepatitis C treatment and management, urine toxicology screening [UTS], relapse prevention, overdose education, motivational interviewing, retention, harm reduction, compassion fatigue, case discussions, materials development, and networking).

RESPONSIBILITIES:

- Provide patient-centered care within the nursing license scope of practice including: initial assessment and intake, induction, stabilization, and maintenance phases of treatment.

- Ensure that state and federal guidelines are followed.

- Collaborate with OBAT provider, social worker/counselors, psychiatrists, pharmacy, primary care provider, and specialty care providers to whom the patient has been referred.

- Coordinate between OBAT provider and pharmacy: assist with prescription processing and refills, prior authorizations, insurance issues etc.

- Complete appropriate documentation in the medical record and comply with state, federal, and departmental policies when sharing/documenting patient care data.
PROGRAM REQUIREMENTS

SAMHSA’S CENTER FOR SUBSTANCE ABUSE TREATMENT (CSAT) DIVISION OF PHARMACOLOGIC THERAPIES

BUPRENORPHINE ADMINISTRATIVE REQUIREMENTS

- Certification, accreditation and waiver approval.
- Maintain accurate provider records.
- Keep records on dispensation of buprenorphine and buprenorphine/naloxone in accordance with DEA regulations for controlled substances as described in 21 CFR 1304.03(b).
- Keep records on prescription and dispensation of medications for the detoxification and maintenance treatment of opioid dependence in accordance with DEA regulations 21 CFR 1304.03(c)
  - Maintain log to include patient identifier, name, dose, and quantity of drug prescribed/dispensed, and date.
  - Requirement may be fulfilled by keeping copies of prescriptions in the patient record. Electronic medical records where the prescription records can be accessed fills this requirement and there is no need to keep copies of the prescriptions in your office.
    - For DATA 2000 compliance, the DEA only needs to review records for medications used in the treatment of opioid dependence; therefore, an option is to keep separate records for these medications to facilitate the review.

PATIENT INITIATION ROADMAP

✓ Initial screening by OBAT staff
✓ Intake performed by OBAT NCM
✓ OBAT provider visit
✓ Induction
✓ Stabilization
✓ Maintenance
CANDIDATES FOR OBAT

- Patient must have a *DSM-5* diagnosis of Opioid Use Disorder or Alcohol Use Disorder. See Appendix 1 & 2.
- Patient is able to come to visits during office hours of operation.
- For patients seeking treatment with agonist medications, they must not have chronic pain requiring ongoing opioid management beyond buprenorphine/naloxone.
- For patients seeking treatment with antagonist medications, they must not have acute/chronic pain issues requiring opioid management.
- Patient is able to be treated in an office based setting safely without harm to self or others.
- Patient should be willing to address use of other harmful and/or illicit substances.
- Patient has been assessed by the treatment team and deemed appropriate for medication treatment in an office based setting.

INITIAL SCREENING

PHONE OR IN PERSON SCREENING BY OBAT STAFF

- See Appendix 3: Telephone Screening.
- *Phone screener includes:* Review of medical, social, and substance use history as well as current use. Demographics, living situation, insurance, safety, and patient’s treatment goals are also explored.
- OBAT team reviews initial screening information and makes decision about potential appropriateness of patient receiving medication for substance use disorder in an office based setting. Appropriate candidates proceed to OBAT intake.
  - We support a low threshold for a trial of medication treatment in OBAT. While screening tools are useful and should be used, it is extremely difficult to distinguish which patients will succeed in an OBAT setting.

OBAT INTAKE
INTAKE PERFORMED BY NURSE CARE MANAGER

See Appendix 4: Nursing Intake for Intake Forms

OBAT NCM Intake Includes:

- Laying the groundwork for a therapeutic relationship with the patient. Assess patient goals for treatment, strengths for obtaining recovery and risks to treatment success.
  - The OBAT NCM values the uniqueness of each individual and helps each person define their own goals.

- Assessment of substance use including substance use history, current status, prior treatment and recovery time.

- Review of medical, mental health and social history. Obtain appropriate signed consent forms to assist with collaboration of care with outside providers and supports.

- Education on medication for addiction treatment: what it is, how it works, medication administration, interactions, side-effects, potential adverse reactions, induction and maintenance processes.

- Reinforce that a substance use disorder is a chronic medical condition that affects numerous aspects of a person’s wellbeing. The OBAT team will support the patient throughout the recovery process, even in the event of a relapse. The patient’s treatment plan will be augmented as necessary to assist the patient in obtaining recovery and achieving identified treatment goals.

- Harm reduction education: overdose prevention education, overdose reversal with naloxone, rescue breathing, ensuring patient has access to naloxone.

- Mandatory screening at time of intake includes:
  - Toxicology screening and pregnancy testing
  - HIV testing strongly recommended
  - Ensure PPD screen is up to date per your institution’s protocol

- Obtain laboratory tests as clinically needed.
  - Consider: complete blood count, comprehensive metabolic panel, hepatic function, pregnancy test, RPR, hepatitis A, B and C serologies

- Review of treatment agreement and program expectations. Patient signs treatment agreement and consents for treatment.
  - Discussion includes:
• Introduction to members of treatment team.
• Patient-centered treatment planning and review.
• Appointment frequency with OBAT NCM and provider.
• Counseling and psychiatric assessment and follow-up if warranted.
• Education regarding medication refills, medication safety and responsibilities for safe storage.
• Review of clinic hours and times available for scheduling visits, including afterhours emergency contact information.
• Review of OBAT goals:
  • Cessation or reduction in harmful substance use.
  • Prevention or reduction of opioid withdrawal symptoms.
  • Prevention or reduction of cravings for opioids and/or alcohol.
  • Restoration of normal physiological functions that may have been disrupted by substance use.
  • Improvement in one’s of quality of life.
• If unable to meet the patient’s needs and the program requirements, the site will assist in referring the patient to another treatment setting that may be better able to meet the patients’ needs.
VISIT WITH OBAT PROVIDER

- Initial provider assessment visit should include:
  - Laying the groundwork for a therapeutic relationship with the patient.
  - Conducting medical, mental health and substance use histories.
  - Physical examination if needed, and review of laboratory test results.
  - Provider confirmation of DSM-5 diagnosis of moderate to severe Opioid Use Disorder or DSM-5 diagnosis of Alcohol Use Disorder.
  - Assessment of appropriateness for medication treatment with either buprenorphine/naloxone or naltrexone.
  - Assessment of appropriateness for receiving treatment for opioid use disorder within an office based setting.

- Follow-up visits with OBAT provider occur a minimum of once every four months.
  - OBAT NCM will manage the patient under the guidance of the OBAT provider with close clinical follow-up and ongoing communication with the provider by telephone, electronic medical record, in person meetings, and team meetings.
  - Many OBAT providers may choose to see patients more frequently. This is particularly true if a patient has difficulty stabilizing in OBAT or if a patient becomes destabilized at any point during the course of their treatment.

- Follow up with primary care provider as warranted based on medical needs. Often the PCP and the OBAT provider are the same, and this will not apply.
TREATMENT AGREEMENT

Goal: Engage patient in the treatment plan, along with the OBAT team. Individualize treatment to meet the needs of the patient. Encourage patient involvement in their treatment.

- See Appendix 9A and 9B for Treatment Agreement Forms.
- Set clear expectations/guidelines.
- Explain treatment agreement verbally and provide in written form, which patient will sign and date. This form will be kept in the patient record. Review each line of the contract, and give a copy to the patient to take home for their review.
  - Encourage patient to ask questions.
  - Review this agreement again with the patient intermittently during the course of treatment and as needed.
  - Provide reassurance about common issues, such as patients’ concerns about entering treatment (provide education around options and support), or the risks of transferring care from one form of medication treatment to another, or patients’ ambivalence about such changes.
- The agreement reinforces that a substance use disorder is a chronic medical condition that affects numerous aspects of a person’s wellbeing. The OBAT team will support the patient throughout the recovery process, even in the event of a relapse. The patient’s treatment plan will be augmented as necessary to assist the patient in obtaining recovery and achieving identified treatment goals.
- Patient is informed of the OBAT program’s 24-hour emergency coverage plan.
- The patient can expect:
  - To be treated with dignity and respect.
  - To be notified if the office is closed and how to seek assistance if needed.
  - That confidentiality will be maintained in compliance with CFR 42.
  - To have a means for contacting a member of the OBAT team or a colleague for emergencies at night, on weekends and/or when the office is closed.
PROGRAM POLICIES AND TREATMENT PROTOCOLS:

- **Clinical Appointment Policy** (See Appendix 10B and 10C): All patients who participate in the OBAT program are required to keep appointments with their primary care providers, OBAT providers, and OBAT NCMs. Appointments with the OBAT team are part of the treatment and should these appointments need to be rescheduled, it is the patient’s responsibility to do so. This does not include random callbacks; please see policy under random call backs.

- **Counseling Policy** (See Appendix 10D). Patients are strongly encouraged to engage in counseling and/or similar intensive recovery support services. If needed, patients should receive assistance with referrals and linkages for counseling and recovery support programs from OBAT staff. Patients are encouraged to attend a minimum of twice monthly counseling visits for the first 12 weeks of treatment. Patients should not be discharged from the OBAT program if they do not comply with this recommendation as these individuals may be at increased risk for relapse. However, patients who do not engage in counseling or outside recovery support services should receive more intensive monitoring from the OBAT team.

- **Behavior Policy** (See Appendix 10E). To provide an optimum treatment environment for all, patients, visitors and staff are expected to maintain appropriate behavior in the clinic and on the OBAT program property.

- **Medication Refill Policy** (See Appendix 10F). Prescriptions are processed following a scheduled office visit or phone encounter. Patients are expected to take their medication as directed and to store it in a safe, secure location. Prescriptions should last until the next scheduled appointment. Patients must keep scheduled appointments to obtain refills.

- **Urine Toxicology Screen Policy** (See Appendix 10G). Toxicology samples are required at each visit in accordance with clinic policies. While sample collection is a monitored process, observed urine screens are discouraged. Any questionable urine is addressed by the OBAT NCM and a repeat sample should be collected that same day.

- Patients struggling with program requirements may be referred to another treatment program or another level of care.
BUPRENORPHINE/NALOXONE PRESCRIPTION POLICIES

The role of the provider/NCM, the office staff, and the patient in the handling of prescriptions:

- Prescriptions are processed following a scheduled office visit or phone encounter.
- Prescriptions will be processed by an OBAT NCM, who will review the medication record, consult with provider, pharmacy and the prescription drug monitoring program (PDMP) if needed, to confirm dosage, refill amounts, and timing of refill.
- OBAT NCM will check insurance coverage, preferred covered medication formulary, and need for prior authorization.
- Following confirmation, the OBAT NCM will generate an electronic prescription under the waived provider’s name, print this electronic prescription and hand deliver to the waived provider for signature.
- Once signed, the OBAT NCM will generate a confidential fax cover sheet, confirm the pharmacy information from the EMR and fax the prescription to the pharmacy on record.
- After the prescription has been faxed, the OBAT NCM will stamp the prescription “faxed,” place the prescriptions in a confidential locked file for 60 days in case of fax error or the need to review the prescription arises.
- All prescriptions are held for 60 days and then destroyed in a secure recycle bin.
- Prescription records are maintained in the electronic medical record for review by clinicians as needed and for DEA regulatory purposes.
  - Of note, under the DEA’s rule, “Electronic Prescriptions for Controlled Substances,” providers have the option to electronically write prescriptions for schedule II-V substances, including buprenorphine. The regulations also allow pharmacies to receive, dispense and archive these prescriptions electronically. Practitioners are only able to issue electronic controlled substance prescriptions when the EMR the provider is using complies with requirements in the final rule. For more information on electronic prescribing, visit https://www.deadiversion.usdoj.gov/ecomm/e_rx/faq/faq.htm

Duration of buprenorphine/naloxone prescription:

- Buprenorphine/naloxone prescriptions generally coincide with visit frequency:
  - At treatment initiation, prescriptions will be written for a maximum of one week with no refills.
  - Following four to six weeks of stabilization, if patient is moving on to the maintenance phase of treatment, prescription refills will increase to two week prescriptions with up to four refills.
• If patient continues to progress, prescriptions will increase to three to four week prescriptions with refills.

• Patients must keep scheduled appointments to obtain prescription refills.

• If a patient begins to struggle in their recovery process or the staff are concerned about giving the patient longer-interval prescriptions, the OBAT team will make the decision to continue with shorter-interval prescriptions as long as is necessary.

• If a patient is homeless, living in an unsafe/unstable setting, the OBAT team will work with the patient to develop a plan that promotes the security of their treatment.

  • Weekly prescriptions with refills, work with shelter staff in an effort to find feasible methods to support the patient and secure their medication if possible.

**Lost, stolen or destroyed buprenorphine/naloxone:**

Special precautions should be taken when a patients reports that their buprenorphine/naloxone has been lost, stolen, or destroyed.

• Buprenorphine/naloxone prescriptions are generally not replaced; patients are informed of this verbally and in writing at the time of intake. However, cases will be reviewed on an individual basis by the OBAT team if requested by the patient.

• If a decision is made to replace the medication, it will be a one-time event and a lost/stolen prescription will not be replaced in the future should this occur.

• If more than a one week supply of replacement prescription is needed, the prescription amount will go back to weekly prescriptions until the team feels it is safe for the patient to be given a larger quantity of medication.

• In all of these events: lost, stolen, destroyed, or damaged medications, prior to receiving a replacement prescription, the patient will be asked to return to the OBAT clinic within 24 hours for an assessment and a toxicology screen. At this time, patients will receive additional education about safe handling and storage of buprenorphine/naloxone by the OBAT NCM and/or provider to prevent these events from reoccurring. The treatment plan should be reviewed, along with prescription interval and frequency of visits to further assess and ensure that there are no additional concerns that should be addressed.

• If the patient continues to experience events of: lost, stolen, damaged or destroyed medications, the team will meet to address this and the potential need to refer the patient to a more structured treatment setting to better safeguard their treatment and recovery.
SAFE AND PROPER STORAGE OF MEDICATION:¹

✓ Use a locked box, bag, or cabinet for safe storage.

✓ Keep medication out of sight/reach of children.

✓ Do not put tablets/films down on counters, sinks, dresser, nightstands or in any public unsecure space.

✓ It is easier for children to put small pieces and crumbs in their mouth.

✓ Always keep buprenorphine/naloxone tablets in a labeled prescription bottle with a child-proof cap.

✓ To prevent breakage of buprenorphine tablets, keep cotton or tissue in the bottle.

✓ Patients should keep medication with an official pharmacy label at all times. Patients may request a second label from pharmacy if they plan to carry the medication on their person.

✓ **Call 911** if an accidental exposure occurs and/or go to the nearest emergency department.

✓ Give all patients a copy of the safety and storage brochure and review the bullet points with them. (See Appendix 11C).

¹ Adapted from: “Protecting Others and Protecting Treatment” STATE OBOT (State Technical Assistance Treatment Expansion Office Based Opioid Treatment of Buprenorphine), and Massachusetts Department of Public Health Bureau of Substance Abuse Services (BSAS). 2016.
Urine Toxicology Screening Policy

- Urine samples are required at each visit.

- All belongings (coats, bags, etc.) are left in the office of the medical assistant or outside the bathroom door. Patients may keep their wallet and cellphone with them.

- No washing of hands until the labeled urine sample is handed to the medical assistant in a bio-hazardous bag.

- No flushing of toilet until urine sample is handed to the gloved medical assistant.

- Observed urine screens are discouraged.
  - As an alternative to observed urines, programs may wish to institute measures to reduce incidents of urine tampering such as: utilizing oral swabs, trash receptacle kept outside of the bathroom, having blue colored toilet water, and assessing characteristics of the urine (e.g. temperature, color, clarity, and creatinine concentration).

- Any questionable urine is repeated the same day.
  - In the event of receiving a questionable urine sample, OBAT NCM addresses concerns with the patient and reinforces that the OBAT team is available to work with the patient throughout their recovery process, especially in the event of recent substance use.
  - OBAT NCM reviews the importance of UTS monitoring and honesty in treatment to ensure that the team has the ability to provide appropriate care.

- Patients will receive a buprenorphine/naloxone prescription refill, naltrexone prescription or extended-release naltrexone injection after an acceptable toxicology sample is obtained.
TREATMENT CONSENTS *(See Appendix 8A – 8E)*

In addition to standard HIPAA laws, federal regulations mandate strict confidentiality for information about patients being treated for substance use disorders (42 CFR Part 2). Additionally, the law requires written patient consent before information about addiction treatment can be disclosed to any other source. For OBAT, this may include any communications with other providers, treatment centers, significant others, or pharmacies.

**SPECIFIC ACTIONS THAT ARE PROHIBITED (WITHOUT CONSENT) INCLUDE THE FOLLOWING:**

- Providing information regarding a patient's past, present, or future participation in addiction treatment.
- Disclosing or transmitting a patient's addiction-related medical records.
- Use of a letterhead that identifies the office as an addiction treatment provider.
- Providing information about those who have applied for treatment or have been interviewed, regardless of whether they actually commenced treatment.
- Providing information about deceased patients.
- Verifying information that inquirers already possess -- in other words, a program can neither confirm nor deny that a patient is/was being treated there (SAMHSA, 1994b).

There are some exceptions to the disclosure laws, such as in the case of medical emergencies or specific legal circumstances. Other than in the case of a medical emergency, you should check with your organization’s legal counsel prior to making disclosures without consent.

**For more information about confidentiality and consents, visit:**

**Legal Action Center**
https://lac.org/resources/substance-use-resources/confidentiality-resources/

**SAMHSA**
https://www.samhsa.gov/about-us/who-we-are/laws-regulations/confidentiality-regulations-faqs
TREATMENT INITIATION, STABILIZATION & MAINTENANCE

CHECKLIST: PRIOR TO BUPRENORPHINE/NALOXONE INDUCTION

✓ Treatment agreement and consents are reviewed and signed.

✓ Reinforce to patient the need for frequent appointment adherence, and establish whether this is realistic; if patient states it is not manageable, this needs to be addressed with the team prior to initiating treatment.

✓ The patient should have counseling or other support services in place or be working towards establishing counseling/support services.

✓ Toxicology screen completed and reviewed by OBAT team.

✓ Pregnancy test for women of childbearing age.
  
  • If positive HCG, OBAT team will immediately assist patient engagement with appropriate OB providers and will manage the patient in OBAT until a warm handoff occurs.
  
  • Buprenorphine is a Category C medication.

✓ If patient presents from detoxification, OBAT team should attempt to obtain discharge paperwork that includes medications administered (i.e. methadone administered in detox may delay induction with partial-agonist or antagonist due to risk of precipitated withdrawal).

✓ NCM consults with waivered provider after initial visit and obtains the prescription from the prescriber.

✓ Schedule induction per protocol in collaboration with patient and team: date, time, prescription, and clinic schedule.

✓ NCM discusses induction plan with patient and faxes prescription to pharmacy for patient to pick up on day of induction.

✓ Patient presents to clinic for induction.
BUPRENORPHINE/NALOXONE INDUCTION

Medication induction is performed by the OBAT NCM either in the office or at home with support by phone. An office induction is preferred to ensure appropriate medication administration technique, adequate medication effect, and to minimize the risk of precipitating withdrawal. Home induction may be appropriate for some patients who are experienced with buprenorphine and its effects on their body.

Buprenorphine/Naloxone Office Induction

Prior to Induction:

- Patient discontinues use of illicit opioids prior to buprenorphine induction to avoid risk of precipitated withdrawal.
- Timeline for opioid discontinuation to be determined as part of induction treatment plan and should be based upon patient’s medical status, current opioid use and opioid dependency.
  - Short-acting opioids: typically discontinue 8-12hrs prior to scheduled induction.
  - Long-acting opioids: typically discontinue 12-24hrs prior to scheduled induction.
  - Methadone: typically discontinue 36-96hrs prior to scheduled induction.
    - Methadone to buprenorphine transfers are especially complex due to the long half-life of methadone and unpredictable metabolic clearance.
    - Please refer to the section on Methadone to buprenorphine transfers in these guidelines, pp. 63-64.

Day 1: Induction

- Patient arrives at clinic in early withdrawal, with prescription medication in hand.
- For patients who are actively using opioids other than buprenorphine, the NCM assesses symptoms with the Clinical Opioid Withdrawal Scale (COWS). If the COWS score is >6-12, the NCM instructs the patient to take the buprenorphine/naloxone as prescribed and per clinic protocol.
- For patients who are self-maintaining with buprenorphine, assessment using COWS may not be necessary. Use clinical judgment and refer to recent urine toxicology.
- NCM supervises medication administration, and educates the patient about the appropriate technique as this is a sublingual/buccal administration that requires being
kept in the mouth for a long period of time for appropriate absorption.

- Buprenorphine 2-4mg initial dose is removed by the patient from their medication bottle, taken transmucosally, observed and under instruction by the OBAT NCM for proper administration.

- NCM reassesses patient after 30-60 minutes, and instructs patient to then take their second dose of 2-4mg as directed if needed, again observed and supervised by the OBAT NCM for proper administration.

- NCM provides written instructions, establishes follow-up plan including same-day telephone check-in and clinic visits.

- The dose will continue to be titrated per prescription instructions and/or until signs and symptoms of withdrawal subside. Typically patients will titrate to 8-12mg by the end of day one; however, this dose may be less or could be higher, and will vary according to a patient’s level of physical opioid dependence at the onset of treatment.

- Update on call provider at the end of the day in case of calls or concerns off hours.

**Day 2 through Day 7:**

- Patient is instructed to take total dose equivalent from day one upon awakening. Patient is then required to check in with the NCM by phone a few hours later or in person if this works better for the patient. If increased symptoms throughout the day, the patient may increase up to 16mg. Daily check-in with a phone note as needed; patient to return to clinic within one week or sooner if needed.

- Patient sees NCM weekly until stable, then every other week, and progresses to monthly as clinically indicated. If a patient requires more support (i.e. homeless), they may present in person for more frequent visits.

**Buprenorphine/Naloxone Home Induction:**

**Prior to Induction:**

- Treatment team has determined that patient is an acceptable candidate for home induction with buprenorphine/naloxone.

  - Appropriate patients should ideally be familiar with buprenorphine/naloxone and not concurrently using other opioids or other sedating medications.

  - Candidates for buprenorphine/naloxone home induction may include: Patients who are self-maintaining on a stable dose buprenorphine/naloxone, patients previously prescribed buprenorphine, patients currently prescribed
buprenorphine/naloxone who are transferring care, patients with a support system that is engaged in their treatment and willing to assist in the process.

- Patient discontinues use of illicit full-agonist opioids prior to buprenorphine/naloxone induction to avoid risk of precipitated withdrawal.

- Verbal and written information provided to the patient about program induction policies, COWS, and medication safety prior to the determined home induction date.

Day 1:

- For patients who are actively using opioids other than buprenorphine, via phone conversation, the NCM and patient assess symptoms using COWS. COWS score of >6-12 achieved.
  
  - COWS may not be necessary for patients self-maintaining on buprenorphine or who are transitioning care.

- NCM reviews proper medication administration with patient. NCM remains on the phone with patient as they self-administer 2-4mg of buprenorphine/naloxone, and remain on the phone until the medication has been fully dissolved.

- NCM provides education and reviews dosing limits and safe storage of medication with the patient.

- Contact with patient should occur during first hour, then every two hours for the next four hours, and then as needed.

- The dose will continue to be titrated per prescription instructions and/or until signs and symptoms of withdrawal subside.

- Update provider on call by end of the day in case of calls or concerns off hours.

Day 2 through Day 7:

- Patient is instructed to take total dose equivalent from day one upon awakening. Patient is then required to check in with the NCM by phone a few hours later. If symptoms increase throughout the day, the patient may increase up to 16mg. Daily check-in with a phone note as needed; patient to return to clinic within one week or sooner if needed.

- Patient sees NCM weekly until stable, then every other week, and progresses to monthly as clinically indicated. If a patient requires more support (i.e. homeless), they may present in person for more frequent visits.
BUPRENORPHINE/NALOXONE STABILIZATION

Goal: Stabilization of dosing, elimination or reduction in use of illicit opioids and other substances, and patient engagement in the treatment plan.

- Target buprenorphine dose should eliminate opioid cravings and withdrawal, with minimal to no side-effects.
- Typical transmucosal buprenorphine/naloxone dose = 8-16mg/day.
- Narcotic blockade is typically reached at 16mg and is recommended in the early stages of recovery. (http://www.naabt.org/education/pharmacology_of_buprenorphine.cfm)
- Divided dosing is especially helpful for patients with chronic pain for dual effectiveness and avoidance of narcotic medications.
- Medication has a long half-life. The majority of patients take buprenorphine twice daily; the prescription may need to be specifically written as twice daily dosing to allow some patients to receive it twice daily while engaged in treatment programs or other settings that require specific instructions on administration.
- Patient returns to clinic after one week for assessment, prescription renewal, toxicology screening, counseling, education, support, and evaluation of medical, mental health and social needs.
- No prescriptions lasting longer than one week are to be given during this phase.
- Refills are permitted, but patient must provide pharmacy information as all prescriptions are faxed or sent electronically to the pharmacies. Patients are never given a hard copy of the prescription.
- Patient sees NCM weekly for four to six weeks until stable. If medication dose is stable, toxicology screens are appropriate and the patient is adherent to the treatment plan, they may progress to the maintenance phase.
BUPRENORPHINE/NALOXONE MAINTENANCE

Once stable, clinic visits every two to four weeks, with refills that coincide with visits.

**Goal:** monthly visits; ultimately, random visits if appropriate for patient; random is more effective in assisting patients (ASAM, White Paper, 2013).

- Many patients will continue making visits more frequently than monthly as patients find these visits important to their recovery process.
- Each decrease in visit frequency requires treatment team review.

Clinic visits to include (See Appendix 6: Nurse Follow-up Form):

- Collection of sample for toxicology screen.

- Assessment of status: medication dose, adherence, tolerance, side effects, cravings and withdrawal; safe storage, recovery, relapse, as well as medical, social and psychiatric issues should all be addressed as indicated.

- Review of treatment plan: visit frequency, counseling, and assess need for additional support.

- OBAT NCM’s notes should be documented in the EMR and available to the entire clinical team, including waived providers.

- Lab testing: if LFTs were elevated at induction, consider rechecking within one to two months or sooner depending on degree of elevation, and regularly monitor thereafter. Elevations are more common in patients with hepatitis C and HIV infection.

- In cases of co-occurring alcohol use disorder, address concerns with patient and consider monitoring with breathalyzer or urine EtG (ethyl glucuronide).
  - Acamprosate (Campral) and disulfiram (Antabuse) may be offered to patients with alcohol use disorder with provider input and agreement.
  - Patients managed on buprenorphine/naloxone cannot be treated with any naltrexone formulation, as these medications are contraindicated.

- Contact other OBAT team members as needed, including OBAT provider and PCP if different and warranted.
  - Waivered OBAT provider visits to occur at least every three to four months.

- Review contact information, including pharmacy at each visit.
• Since buprenorphine is a schedule III substance, a maximum of five refills are permitted for each prescription. Therefore, refills covering up to a six month duration may be provided to the stable patient.

• In addition to office visits, the OBAT NCM performs telephone contact for support as needed.
Naltrexone Initiation: Patient Selection

Candidates for treatment with naltrexone include patients who:

- Are not currently using opioids, but have a history of opioid use disorder and are at risk for relapse.
- Have a high degree of motivation for abstinence from opioids.
- Have been successful on opioid agonists and wish to discontinue agonist therapy.
- Are not interested in agonist/partial agonist therapy to treat their opioid use disorder.
- Certain patients who have not experienced successful treatment with agonist therapy.
- History of alcohol use disorder.

Contraindications:

- Advanced liver disease or acute hepatitis.
- Moderate to severe renal impairment.
- Patients with chronic or acute pain that requires opioid analgesics.
- Patients who are unable to remain opioid free for extended periods of time.
- Patients with advanced psychiatric disease, active suicidal/homicidal ideation, especially if symptoms worsen during withdrawal.
- Patients who are currently opioid dependent, or taking opioids, or have an opioid positive urine screen.
- A patient who fails the naloxone/naltrexone challenge test.
- Patients who have displayed a hypersensitivity to naltrexone, PLG, carboxymethyl cellulose, or any other components of the diluent.

Special considerations:

Pain: chronic pain must be managed with non-opioids. Acute pain requires anesthesia consult. If a patient has a surgical procedure pending, they may want to consider delaying naltrexone treatment until after the procedure.

Cirrhosis: Naltrexone is extensively metabolized through the liver and should not be administered if AST/ALT are more than five times the normal limits.
**Pregnancy:** There has not been sufficient research to assess the safety or efficacy of naltrexone in pregnancy. Naltrexone, both oral and injectable formulations, are Category C medications. The provider would need to evaluate the risk/benefit and appropriate consent of unknown risk should be utilized.

**Breastfeeding:** It is known that naltrexone from the oral formulation passes into breast milk. It is not known if extended-release injectable naltrexone passes into breast milk. In vivo studies indicate potential tumorigenicity. At this time, labeling from the manufacturer advises against breastfeeding while on naltrexone, both with oral and injectable formulations.

**Anemia/Thrombocytopenia:** Administer extended-release injectable naltrexone with caution and observe site for bleeding. Consider oral formulation.

**Obese/large body habitus:** Extended-release injectable naltrexone must be administered IM into gluteal muscle using the contents of the medication package. Alternate treatment may be considered for patients whose body habitus precludes an intramuscular gluteal injection with one of the provided needles. Consider oral formulation.
CHECKLIST: PRIOR TO NALTREXONE INITIATION

✓ Treatment agreement and consents are reviewed and signed.

✓ Reinforce with patient the need for frequent appointment adherence, and establish whether this is realistic. If patient states it is not manageable, this needs to be addressed with the team prior to initiating treatment.

✓ The patient should have counseling or other support services in place or be working towards establishing counseling/support services.

✓ Patient cleared by psychiatry if concerning mental health history.

✓ Labs appropriate: HCG neg. LFTs < 5x normal.

  • If positive HCG, OBAT team will immediately assist patient with engagement to appropriate OB providers.

    ▪ Naltrexone is a Category C medication.

  • If LFTs are >5x normal, appropriate medical evaluation should occur to determine cause of transaminase elevation.

  • Patients may be offered alternative treatment options (i.e. buprenorphine or methadone for OUD, acamprosate for AUD, monitoring in OBAT without MAT, psychotherapy).

  • When deciding to start naltrexone treatment in pregnant patients or those with elevated LFTS, providers should utilize clinical judgment, carefully weighing the risk versus benefit of initiating naltrexone and documenting an informed consent to treat with the patient.

✓ UTS that is negative for opioids.

  • Detoxification from opioids should be completed prior to the administration of naltrexone to prevent precipitated or spontaneous withdrawal. The patient must not be experiencing withdrawal symptoms. Patients should discontinue short-acting opioids at least three to seven days prior to starting naltrexone. If taking long-acting opioids such as methadone or buprenorphine, the patient must be off for at least seven to 10 days.

  • Detoxification from alcohol is not always necessary. However, detoxification from alcohol is recommended prior to naltrexone initiation if a patient has a history of alcohol-related seizures, delirium tremens (DTs), longstanding daily use, presence of withdrawal signs or symptoms, or as otherwise clinically indicated.
• If patient presents from detoxification or other inpatient treatment, OBAT team should attempt to obtain discharge paperwork that includes medications administered (i.e. methadone or buprenorphine administered in detox will delay induction with antagonist due to risk of precipitated withdrawal). This paperwork must be reviewed by the treatment team.

✓ NCM consults with OBAT provider and clinical team after initial visit. After OBAT team review, schedule induction per protocol in collaboration with patient and team: date, time, prescription, and clinic schedule.

✓ NCM discusses the medication initiation plan with the patient and orders the medication. Oral naltrexone tablet prescription may be e-faxed to pharmacy for patient to pick up. Extended-release injectable naltrexone often requires insurance prior authorization and ordering through a specialty pharmacy; this process may take several days and requires thoughtful planning.

✓ Patient presents to clinic for induction/medication initiation.
Naltrexone Initiation

- Patients should be started on the oral form of the medication prior to receiving the extended-release IM injection.
  
  - This is to mitigate allergic reactions, side effects, adverse reactions or any other intolerance of the medication.
  
  - Typically, patients will remain on the oral formulation for a few days before receiving their first extended-release naltrexone injection to assess for side effects and any contraindications.

- Patient to be given emergency card, bracelet and/or dog tag.

- The first naltrexone dose should ideally be observed in clinic.

- A “naloxone challenge” or “naltrexone challenge” should be performed if there is a risk of adverse reaction to naltrexone or a risk of precipitating withdrawal with a long-acting naltrexone formulation.
NALOXONE CHALLENGE

- NCM meets with the patient to assess current status.
- UTS negative for all opioids.
- The patient must be off short acting opioids for at least three to seven days. If taking long acting opioids such as methadone or buprenorphine, the patient must be off for seven to 10 days or longer.
- Obtain baseline vital signs.
- Obtain baseline Clinical Opiate Withdrawal Scale (COWS) score.
  - If the patient is still having signs of opioid withdrawal, even if the toxicology screen is negative, do not perform a naloxone challenge. It will be positive.
- A total of 0.8-1.2mg naloxone hydrochloride should be administered IM. This may be divided into two doses to minimize risk of severe withdrawal.
  - Administer 0.4mg naloxone hydrochloride IM
    - Complete COWS at 15min and again at 30min following injection.
  - If no signs or symptoms of withdrawal, administer a second dose of 0.4 - 0.8mg naloxone hydrochloride IM.
    - Observe and complete COWS at 15min and again at 30min following this second injection.
- **Negative Naloxone Challenge:** No change in subjective or objective signs of withdrawal. Proceed with administering full dose of naltrexone or extended-release injectable naltrexone.
- **Positive Naloxone Challenge:** If the patient experiences any symptoms of withdrawal, immediately stop the naloxone challenge. Do not give any more naloxone. In dependent individuals, naloxone will precipitate withdrawal that usually emerges within 5-10 min and dissipates within 30 min. These symptoms should be mild. The most common early signs of a positive challenge will be the patient reporting an increase in anxiety and an increase in heart rate. Have the patient come back in one to two days and repeat naloxone challenge.
**NALTREXONE CHALLENGE WITH ORAL FORMULATION**

- NCM meets with the patient to assess current status.
- UTS negative for all opioids.
- The patient must be off short acting opioids at least three to seven days, long-acting requires at least seven to 10 days.
- Obtain baseline vital signs.
- Obtain baseline Clinical Opiate Withdrawal Scale (COWS) score.
  - If the patient is still having signs of opioid withdrawal, even if the drug screen is negative, do not perform a naltrexone challenge. It will be positive.
- Observe patient self-administer 25mg naltrexone by mouth. Advise patient to remain in the clinic for 60 minutes to monitor the presence/absence of withdrawal symptoms.
- **Negative Naltrexone Challenge:** No change in subjective or objective signs of withdrawal. Proceed with extended-release naltrexone injection per protocol.
- **Positive Naltrexone Challenge:** If the patient experiences any symptoms of withdrawal, immediately stop the naltrexone challenge. Do not give any more naltrexone. Reassure the patient that the symptoms will begin to dissipate in four to six hours. The most common early signs of a positive challenge will be patient report of increased anxiety and an increase in heart rate. Symptom management with adjunctive medications to occur as appropriate with provider input. Have the patient come back in one to two days for a repeat naltrexone challenge.
EXTENDED-RELEASE INJECTABLE NALTREXONE ADMINISTRATION

For the most up to date information about administration of extended-release injectable naltrexone, please refer to the manufacturer’s safety information, medication guide and prescribing information. For Vivitrol: https://www.vivitrol.com/

- Obtain extended-release injectable naltrexone from pharmacy per written prescriber order. Standard dose is 380mg IM. Do not prepare suspension prior to patient arrival.

- Extended-release injectable naltrexone should be stored in the refrigerator. Prior to preparation, allow the drug to reach room temperature. This takes about 45 mins.

- After meeting with the patient and ensuring continued opioid abstinence, reconstitute and immediately administer medication following the specific detailed directions contained in the extended-release injectable naltrexone medication package insert.
  - UTS negative for all opioids, and/or negative naloxone/naltrexone challenge.

- Extended-release injectable naltrexone should be administered as an intramuscular gluteal injection every 28 days.

Special Notes:

- Unrefrigerated, extended-release injectable naltrexone can be stored at temperatures not exceeding 25 °C (77 °F) for no more than seven days prior to administration. Do not expose unrefrigerated product to temperatures above 25 °C (77 °F). This medication should never be frozen.
  - Mark the medication each time it is removed and returned to the refrigerator.

- A properly mixed suspension will be milky white, will not contain clumps, and will move freely down the walls of the vial.

- Use only the needles specifically designed for administration of extended-release injectable naltrexone. Select the appropriate needle based on patient’s body habitus. Do not make any substitutions for components in the medication carton.

- Administer the suspension by deep intramuscular injection into a gluteal muscle, alternating buttocks per monthly injection. Aspirate for blood before injecting.

- If the needle clogs or blood aspirates, change to the spare needle provided in the carton and administer medication into an adjacent site in the same gluteal region, again aspirating for blood prior to injecting.
• Document administration of extended-release injectable naltrexone and note right or left gluteal injection site.

• Advise patient to contact the OBAT clinic, or go to the emergency department in the event of suspected injection site or other adverse reaction.

**Adverse effects and patient education**

- **Injection Site Reactions**: Providers should be trained in proper techniques for IM injections to prevent problems. Extended-release injectable naltrexone injections may cause pain or tenderness at the injection site, which usually resolves in a few days. More serious reactions such as swelling, erythema, bruising, and pruritus have been reported, generally as the result of an inadvertent subcutaneous injection.

- **Vulnerability to Opioid Overdose**: Following injection with extended-release naltrexone, a patient's opioid tolerance is reduced markedly from baseline prior to treatment. Accordingly, patients are vulnerable to a potentially fatal overdose approaching the end of the dosing interval, if a dose is missed or if treatment is discontinued. Attempting to break through the opioid blockade can also result in fatal overdose. The OBAT NCM should outreach to and attempt to reengage with patients who miss an injection.

- **Hepatic Injury**: There have been cases of hepatitis and clinically significant liver injury associated with extended-release injectable naltrexone. Patients should be made aware of this risk.

- **Depression and Suicide**: In pre-market clinical trials of extended-release injectable naltrexone, reports of depression were overall infrequent but more common in the group that received injectable naltrexone than the group that received the placebo. Patients should be evaluated, monitored and treated appropriately, and families and caregivers should be alerted to the need to monitor patients for depression or suicidality.

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NALTREXONE (ORAL OR INJECTABLE) STABILIZATION

- Patient returns to clinic after one week for assessment, toxicology screening, counseling, education, support, and evaluation of mental health, medical, and other needs.

- If a patient misses an extended-release naltrexone injection, he/she should be instructed to receive the next injection as soon as possible. Reassess patient status prior to administering medication. Consider naloxone/naltrexone challenge if suspected opioid use or if injection has lapsed for an extended period of time. Augment treatment plan as needed.

- Patient sees NCM weekly for four to six weeks until stable. If toxicology screens and breathalyzer/EtG screens are appropriate and the patient is adherent to the treatment plan, they may then progress to the maintenance phase.
NALTREXONE MAINTENANCE

- Once stable, clinic visits every two to four weeks.

- *Goal:* Clinic visits every 28 days, occurring on the date of the patient’s extended-release naltrexone injection.
  - Each decrease in visit frequency requires treatment team review.

**Clinic visits to include (See Appendix 6: Nursing Follow-up Form):**

- Collection of sample for toxicology screening.

- Assessment of status: recovery, relapse, and medical, social and psychiatric issues should be addressed as indicated.
  - For management of pain in patients who are engaged in naltrexone treatment, refer to section titled “Pain Management: Naltrexone” or to Appendix 13.

- Monitor and assess for potential medication side-effects or adverse reactions: injection site reaction, hepatic complications, gastrointestinal distress, depression, eosinophilic pneumonia, etc.

- Review of treatment plan: visit frequency, counseling, and assess need for additional supports.

- If history of alcohol use disorder, address concerns with the patient and consider monitoring with breathalyzer or urine EtG (ethyl glucuronide).
  - Acamprosate (Campral) and disulfiram (Antabuse) may also be offered to patients with problematic alcohol use with provider input and agreement.

- Lab testing: if liver function tests were elevated at induction, consider rechecking within one to two months or sooner depending on degree of elevation. Continue to regularly monitor LFTs thereafter.

- Contact other OBAT team members as needed, including OBAT provider and PCP if different and warranted.

- Review contact information, including specialty pharmacy, at each visit.

- OBAT provider visits at least every three to four months

- In addition to office visits, OBAT NCM performs telephone contact for support as needed.
ADDRESSING SUBSTANCE USE DURING TREATMENT

CLINICAL DECISION-MAKING IN THE EVENT OF SUBSTANCE USE DURING TREATMENT

- OBAT is a harm reduction model and therefore does not recommend automatic discharge for patients who struggle with substance use while engaged in treatment.

- Illicit substance use is typically reported by the patient during an OBAT office visit and then later confirmed by toxicology screening. When substance use is reported during an encounter, the treatment plan can be adjusted at the time of the visit.

- When a patient discloses substance use, the OBAT NCM or provider should assess the circumstances surrounding the incident(s) (i.e. home environment, work environment, role of support persons and persons close to the patient). These details are necessary to effectively adjust the treatment plan to meet the evolving needs of the patient.

- Clinical decisions to augment the treatment plan should be based upon the patient’s overall wellbeing and standing within OBAT, and therefore based upon multiple data points including: patient engagement with the treatment plan, recent toxicology screens, and status of physical, mental, and social health.

- OBAT team communication via in-person team meetings, telephone calls, and the electronic medical record is essential to ensure consistent, high-quality patient care. When a patient is struggling with ongoing substance use during treatment, OBAT NCMs and providers should review the case specifics and utilize shared decision making to adjust the treatment plan to best meet the needs of the patient.

- If illicit or harmful substance use occurs, the treatment plan should first be revised to increase monitoring and supports.

- If the patient complies with an intensified treatment plan and is able to be stabilized, the OBAT team will restructure treatment and continue with medication for addiction treatment within an office based setting.

- In a case of continued substance use despite an intensified treatment plan, a patient may be referred to a higher level of care for added support and safety.
  - When possible, ensure a “warm handoff” to providers at agency/program to which the patient has been referred.

- Clinicians should always carefully weigh the risk versus benefit of continuing treatment in an office based setting prior to referring to another level of care.
• In every instance that the treatment plan is augmented, careful documentation should occur in the electronic medical record to support either continuing the patient in the current treatment program or referring to another level of care.

• Potential situations in which the risk of continuing treatment may outweigh the benefit:
  • Inability to stabilize a patient on buprenorphine treatment within an office based setting (i.e. ongoing opioid use despite adequate buprenorphine/naloxone dosage).
  • Opioid use during the end of an extended-release naltrexone injection dosing interval.
  • Multiple negative buprenorphine UTS results for buprenorphine prescribed patients.
  • Ongoing use of benzodiazepines, cocaine/stimulants, alcohol or other harmful/illicit substances causing impairment, sedation, overdose, medical events, and/or hazardous unsafe behaviors despite interventions by the OBAT team.
  • Repeated incidents of presenting intoxicated, overdose, or hospitalization related to substance use.

• Patients who are referred to a higher level of care or discharged should be reconsidered for OBAT in the future.
REVISION OF TREATMENT PLAN MAY INCLUDE:

- More frequent visits
- Shortened prescription intervals
- Loss of refills
- Confirmation of counseling and team engagement with counselor
- Clinical team meeting with patient
- Referral to relapse prevention groups or individual therapy
- Referral to intensive outpatient program (IOP)
- Psychiatric evaluation and treatment per psychiatric assessment
- Residential treatment
- Involvement of social services
- Increased collaboration with community providers
- Increased engagement with law enforcement
- Family/support involvement

REFERRAL TO HIGHER LEVEL OF CARE MAY INCLUDE:

- Detoxification/crisis stabilization services/transitional support services
- Residential treatment
- Methadone maintenance
- Directly observed buprenorphine/naloxone daily dosing in outpatient treatment program
- Dual Diagnosis
BUPRENORPHINE/NALOXONE: ABERRANT URINE TOXIC SCREEN RESULTS

● Illicit substance use is typically reported by the patient during an OBAT visit and then later confirmed by toxicology screening. When substance use is reported during an encounter, the treatment plan can be adjusted at the time of the visit.

● If a patient does not disclose substance use or inappropriate medication management during the office visit, and the toxicology screen is inappropriate (i.e. positive for illicit substances not reported or negative for buprenorphine), the patient is called by the OBAT NCM within 24 hours of UTS result in an effort to address a potential relapse or medication issue.
NEGATIVE BUPRENORPHINE

- Buprenorphine adherence is necessary for successful treatment. Nonadherence may be self-reported or found on a toxicology screen that is negative for buprenorphine.

- There are a variety of reasons why a patient may be nonadherent to buprenorphine and/or have a negative buprenorphine toxicology screen, including:
  - Self-increasing buprenorphine dose or a lost/stolen/destroyed medication, and therefore running out of medication early.
  - Forms of diversion. Diversion may be well intended, such as sharing of medication with a friend, or more salacious such as selling medication for profit.
  - Submission of a tampered urine sample.
  - If a patient’s dose is less than 4-6mg, the toxicology sample may need to be sent for confirmatory testing due to the test cut-off limits and therefore an inability to react positive to buprenorphine despite adherence to treatment.

- In all cases of reported nonadherence to buprenorphine, the OBAT NCM should assess and reinforce medication adherence, including total daily buprenorphine dose, dosing schedule, proper administration, and safe storage.
  - OBAT NCM and treatment team should consider that the patient’s buprenorphine/naloxone dose may need to be adjusted (i.e. increased if struggling, decreased if taking less than prescribed). The entire OBAT team should be consulted prior to adjusting a patient’s medication dose.

- If the patient provides an adequate explanation regarding a negative buprenorphine/naloxone result to the OBAT NCM, the OBAT NCM will establish a follow-up plan that includes a return to weekly clinic visits and weekly prescriptions.

- If the patient is unable to provide an adequate explanation regarding a negative buprenorphine/naloxone result, they should return to the clinic within 24 hours.
  - At return visit, a repeat toxicology screen should be obtained and sent for confirmatory testing (i.e. GC/MS) that includes checking for the presence of buprenorphine and buprenorphine’s metabolite, norbuprenorphine.
  - OBAT providers may consider calling the patient between scheduled clinic visits for a “random callback” visit that includes patient assessment, medication count and toxicology screen.
  - If patient denies any reason for negative buprenorphine result, and repeat is again negative, the entire OBAT team should meet to discuss the next best steps. If diversion is likely or if the risk outweighs the benefit of continuing treatment in an OBAT setting, the patient may be referred to a higher level of care.
POSITIVE OPIOIDS

- Report of opioid use or positive opioid toxicology screen result is addressed by OBAT NCM during visit or through subsequent phone conversation and the treatment plan is intensified accordingly to meet the needs of the patient.

  - If patient has a positive opioid UTS and does not report opioid use at the visit and denies use when addressed on the phone with OBAT NCM, the patient should return to the clinic for a repeat visit as soon as possible for assessment, within 24-48 hours.

  - A report of opioid use or a positive opioid UTS will result in intensification of the treatment plan potentially including: increased frequency of clinic visits, confirm attendance and increase frequency of counseling, encourage meetings, provide education on relapse prevention and overdose and send naloxone prescription to specified pharmacy. This includes the patient returning to weekly clinic visits until stable.

  - If the patient has ongoing reports of opioid use or positive opioid screens, and the OBAT team determines that the risk outweighs the benefit of continuing treatment in current setting, the patient will be assisted with transfer to higher level of care. Patient may return to OBAT at a later date after the OBAT team and patient meet to assess how to assist them differently in their treatment moving forward.
POLYSUBSTANCE USE:

COCAINE

- Report of cocaine use or positive cocaine UTS result is addressed by OBAT NCM during visit or through subsequent phone conversation and the treatment plan is intensified accordingly to meet the needs of the patient.
  
  - If patient has a positive UTS and does not report cocaine use at the visit and denies use when addressed on the phone with OBAT NCM, the patient should return to the clinic for repeat visit for assessment as soon as possible, within 24-48 hours.

- A report of cocaine use or a positive cocaine UTS will result in intensification of the treatment plan, relapse prevention education and support. This includes the patient returning to weekly clinic visits until stable.
  
  - Contingency management combined with psychosocial support (CBT, counseling) has been shown to be an effective strategy for decreasing stimulant misuse and should be considered when possible.

AMPHETAMINES

- Report of illicit amphetamine use or positive amphetamine UTS result is addressed by OBAT CM during visit or through subsequent phone conversation and the treatment plan is intensified accordingly to meet the needs of the patient.
  
  - If patient has a positive UTS and does not report amphetamine use at visit and denies use when addressed on the phone with OBAT NCM, patient must return to clinic for repeat as soon as possible, within 24-48 hours.

  - If a patient denies amphetamine use, the OBAT team should consider sending the toxicology screen for confirmatory testing to identify specific metabolites as false-positive amphetamine results are common and can occur due to cross-reactivity from commonly prescribed and over-the-counter medications.

- A report of illicit amphetamine use or a positive amphetamine UTS will result in intensification of the treatment plan, relapse prevention education and support. This includes the patient returning to weekly clinic visits until stable.
  
  - If patient reports struggling with attention deficit and/or hyperactivity, offer patient referral to psychiatry for evaluation.

  - If patient reports diagnosis of ADHD and is requesting amphetamine medications, patient should undergo neuro-psych evaluation for a proper diagnosis.

  - Check prescription drug monitoring program (PDMP) to search for prescription(s) not reported.
- Contingency management combined with psychosocial support has been shown to be an effective strategy for decreasing stimulant misuse and should be considered.

**BENZODIAZEPINES**

- Report of illicit benzodiazepine use or positive benzodiazepine UTS result is addressed by OBAT NCM during visit or through subsequent phone conversation and the treatment plan is intensified accordingly to meet the needs of the patient.
  - If patient has a positive UTS and does not admit to use at the visit and denies use when addressed on the phone with OBAT NCM, the patient must return to clinic for repeat as soon as possible, within 24-48 hours.
- A report of illicit benzodiazepine use or a positive benzodiazepine UTS will result in intensification of the treatment plan, relapse prevention education, and overdose prevention education. This includes the patient returning to weekly clinic visits until stable.
  - If patient reports struggling with anxiety, offer referral to psychiatry for evaluation. Providers should make every effort to stay clear of medications with potential for misuse.
  - Check the PDMP to search for unreported prescribed medications.
  - Toxicology samples will be sent for confirmatory testing and identification of the medication in instances of ongoing positive benzodiazepine screens.
  - Ongoing benzodiazepine misuse despite an intensified treatment plan may result in referral to a higher level of care. Patient may return to OBAT at a later date after the OBAT team and patient meet to assess how to assist them differently in their treatment moving forward.

**ALCOHOL**

- In addition to frequent clinic visits, patients with concerning alcohol use or co-morbid alcohol use disorder should be monitored via breathalyzer or urine EtG testing.
- A report of risky alcohol use or positive urine EtG result is addressed by OBAT NCM during visit or through subsequent phone conversation and the treatment plan is intensified accordingly to meet the needs of the patient.
- Patients struggling with alcohol use and/or cravings may be offered acamprosate (Campral) or disulfiram (Antabuse) with provider input and agreement. Patients managed on buprenorphine/naloxone cannot be treated with any naltrexone formulation, as these
medications are contraindicated. See Appendix 8A for Consent to Treat with Disulfiram form.

- Patients presenting to clinic impaired from recent alcohol use or with a positive breathalyzer screen will require treatment plan revision. Additionally, a safety assessment should be completed, which may include ability to care for any accompanying children or alternate transportation home.

- Ongoing alcohol misuse, presenting to clinic impaired or noted ED visits or hospital events for ETOH intoxication/use may result in referral to a higher level of care. Patient may return to OBAT at a later date after the OBAT team and patient meet to assess how to assist them differently in their treatment moving forward.
NALTREXONE RELAPSE & ABERRANT URINE TOXICOLOGY SCREEN RESULTS

- As previously stated, illicit substance use is typically reported by the patient during an OBAT office visit and then later confirmed by toxicology screening. When substance use is reported during an encounter, the treatment plan can be adjusted at the time of the visit.

- In all cases of an unexplained toxicology results (i.e. patient did not report substance use at visit), the patient is called by the OBAT NCM within 24 hours of result in an effort to address a potential relapse.

- OBAT clinicians should always carefully weigh the risk versus benefit of continuing treatment with naltrexone in current treatment setting prior to referring to another level of care. In cases of patients using opioids while engaged in naltrexone treatment, rather than referring to another level of care, it may be appropriate to discuss transition to buprenorphine/naloxone due to buprenorphine’s partial-agonist effects of increased assistance managing opioid cravings, withdrawal, and protection against fatal overdose.
**OPIOID USE:**

- Report of opioid use or positive opioid UTS result is addressed by OBAT NCM during visit or through subsequent phone conversation and the treatment plan is intensified accordingly to meet the needs of the patient.
  - If patient has a positive UTS and does not report opioid use at visit and denies use when addressed on the phone with OBAT NCM, the patient must return to clinic for repeat as soon as possible, within 24-48 hours.

- Intensify OBAT plan including: increased frequency of clinic visits, confirm attendance and increase frequency of counseling, encourage meetings, provide relapse prevention education and overdose prevention education and a prescription for naloxone. This includes the patient returning to weekly clinic visits until stable.

- Educate the patient about the increased sensitivity to opioids and the consequential increased risk of a fatal overdose in the event of a relapse. Reduced tolerance is especially concerning at the end of a dosing interval. However, an attempt to overcome the opioid blockade effect of extended-release injectable naltrexone is possible at any point and is extremely dangerous with the potential to cause respiratory arrest and circulatory collapse.

- If relapse occurs towards the end of naltrexone interval (within a week of injection due date), restart the patient on naltrexone only after obtaining a UTS negative for opioids and a successful naloxone/naltrexone challenge has been performed. Do not administer naltrexone if there is any chance opioids are on board.

- With opioid use during naltrexone treatment, a clinical assessment should always occur to evaluate if continuing with naltrexone treatment is in the best interest of the patient or if a different medication or level of care should be considered.

- If the patient is unable to abstain from using opioids for a long enough period of time to safely be restarted on naltrexone, referral to a higher level of care (detox, residential, buprenorphine/naloxone treatment, methadone maintenance) should occur. Patient may return to OBAT for naltrexone treatment at a later date after the team and patient meet to assess how to assist them differently in their treatment moving forward.
**ALCOHOL USE:**

- For patients with known alcohol use disorder or concerning alcohol use, monitoring via urine EtG and/or breathalyzer screening is advised.

- A report of risky alcohol use or positive urine EtG result is addressed by OBAT NCM during visit or through subsequent phone conversation and the treatment plan is intensified accordingly to meet the needs of the patient.

- OBAT providers should consider checking LFTs more regularly in patients consuming alcohol while on naltrexone treatment since this medication is highly metabolized through the hepatic system.

- Patients struggling with alcohol use and/or cravings may be offered concomitant acamprosate (Campral) or disulfiram (Antabuse) with provider input and agreement. See Appendix 8A for Consent to Treat with Disulfiram form.

- Patients presenting to clinic impaired from recent alcohol use or with a positive breathalyzer screen will require treatment plan revision. Additionally, a safety assessment should be completed, which may include ability to care for any accompanying children or alternate transportation home.

- Ongoing alcohol misuse, presenting to clinic impaired, or noted ED visits or hospital events for ETOH intoxication/use may result in referral to a higher level of care. Patient may return to OBAT at a later date after the OBAT team and patient meet to assess how to assist them differently in their treatment moving forward.
POLYSUBSTANCE USE:

COCAINE

- Report of cocaine use or positive cocaine UTS result is addressed by OBAT NCM during visit or through subsequent phone conversation and the treatment plan is intensified accordingly to meet the needs of the patient.
  - If patient has a positive UTS and does not report cocaine use at the visit and denies use when addressed on the phone with OBAT NCM, the patient must return to clinic for repeat as soon as possible, within 24-48 hours.
- A report of cocaine use or a positive cocaine UTS will result in intensification of the treatment plan, relapse prevention education and support. This includes the patient returning to weekly OBAT visits until stable.
  - Contingency management combined with psychosocial support has been shown to be an effective strategy for decreasing stimulant misuse and should be considered.

AMPETAMINES

- Report of illicit amphetamine use or positive amphetamine UTS result is addressed by OBAT NCM during visit or through subsequent phone conversation and the treatment plan is intensified accordingly to meet the needs of the patient.
  - If patient has a positive UTS and does not report amphetamine use at the visit and denies use when addressed on the phone with OBAT NCM, the patient must return to clinic for repeat assessment and UTS as soon as possible, within 24-48 hours.
  - If a patient denies amphetamine use, the OBAT team should consider sending the toxicology screen for confirmatory testing to identify specific metabolites as false-positive amphetamine results are common and can occur due to cross-reactivity from commonly prescribed and over-the-counter medications.
- A report of illicit amphetamine use or a positive amphetamine UTS will result in intensification of the treatment plan, relapse prevention education and support. This includes the patient returning to weekly clinic visits until stable.
  - If patient reports struggling with attention deficit and/or hyperactivity, offer patient referral to psychiatry for evaluation.
  - If patient reports diagnosis of ADHD and is requesting amphetamine medications, patient will be required to undergo neuro-psych evaluation for a proper diagnosis.
  - Check PDMP to search for unreported prescriptions.
Contingency management combined with psychosocial support has been shown to be an effective strategy for decreasing stimulant misuse and should be considered.

**Benzodiazepines**

- Report of illicit benzodiazepine use or positive benzodiazepine UTS result is addressed by OBAT NCM during visit or through subsequent phone conversation and the treatment plan is intensified accordingly to meet the needs of the patient.
  - If patient has a positive UTS and does not report benzodiazepine use at the visit and denies use when addressed on the phone with OBAT NCM, the patient must return to clinic for repeat as soon as possible, within 24-48 hours.
  - A report of illicit benzodiazepine use or a positive benzodiazepine UTS will result in intensification of the treatment plan, relapse prevention education, and overdose prevention education. This includes the patient returning to weekly clinic visits until stable.
  - If patient reports struggling with anxiety, offer patient referral to psychiatry for evaluation. Every effort should be made to stay clear of medications with potential for misuse.
  - Check PDMP to search for unreported prescriptions.
  - Urine samples will be sent for confirmatory testing and identification of the benzodiazepine for instances of ongoing positive benzodiazepine screens.
  - Ongoing positive illicit benzodiazepine UTS will result in referral to a higher level of care. Patient may return to OBAT for naltrexone treatment at a later date after the team and patient meet to assess how to assist them differently in their treatment moving forward.
**Presenting Impaired**

Any patient who presents to the clinic intoxicated (i.e. under the influence of alcohol or any other substance) will require urgent team assessment, safety assessment, and revision of treatment plan. Additionally, if a patient presents intoxicated is accompanied by a child or other dependent, please refer to your institution’s policies regarding safety concerns and mandated reporting.

**Diversion**

In cases of suspected diversion, the patient should be asked to come into the clinic for an urgent assessment. This assessment should include a discussion with the patient about treatment team concerns, toxicology testing and a medication count. When possible, confirmatory testing (i.e. via GC/MS) is recommended to confirm presence of buprenorphine and its metabolite norbuprenorphine.

Any patient known to be diverting buprenorphine will be evaluated by the treatment team to discuss appropriate next steps and possibly discharged from the OBAT program (e.g. patient will be seamlessly transitioned to methadone maintenance or another higher level of care).
**Buprenorphine/Naloxone Tapering**

- A substance use disorder is a chronic and complex condition, therefore enforcing a predefined treatment duration is not recommended nor advised.

- Some patients may choose to taper off of buprenorphine/naloxone. These patients will continue to be supported by the OBAT team and receive assistance with dose decreases, and management of withdrawal symptoms. The taper duration is individualized to the patient and should be continually adjusted to meet the patient’s needs.
  - Buprenorphine/naloxone should be tapered over days, weeks, or months, depending on patient’s tolerance of symptoms.

- Upon abrupt buprenorphine discontinuation, withdrawal syndrome may occur.
  - Subjective withdrawal symptoms typically begin within the first three days.
  - Autonomic withdrawal signs (lacrimation, rhinorrhea, tremors, chills, gooseflesh).
  - General complaints include: restless leg, insomnia, anxiety, abdominal distress.

- Protracted abstinence syndrome can occur and persist for months or years following discontinuation of the medication. It is important to respond to patient’s protracted withdrawal symptoms to support their recovery process and avoid relapse.

- Tapering/discharge from program or initiation of intensive treatment plan should be considered for the following cases:
  - Negative buprenorphine screens.
  - Ongoing opioid use or use of other illicit drugs and the risk of continuing treatment outweighs the benefit.
  - Patient presents to OBAT clinic impaired, incidence of overdose, or hospitalization related to substance use and the risk of continuing treatment outweighs the benefit.
  - Multiple missed appointments or inability to contact patient:
    - Address with treatment team and document in EMR. If unable to reach patient, refills should be canceled in hopes this will bring the patient back in to care.

- Patients who are referred to a higher level of care or discharged, will be reconsidered for OBAT in the future.
**Naltrexone Discontinuation**

- There is no withdrawal syndrome associated with naltrexone discontinuation.

- Some patients may choose to discontinue naltrexone. These patients may continue to be supported by the OBAT team and receive assistance with their recovery in terms of monitoring and clinical management. Patients choosing to discontinue naltrexone should be encouraged to continue psychosocial therapies and mutual-help groups.

- Some patients may stop naltrexone due to side-effects or adverse reactions. In this case, alternative treatment strategies should be discussed.

- Naltrexone discontinuation/discharge from program or initiation of intensive treatment plan should be considered for the following cases:
  
  - **Opioid use**: Ongoing opioid use and the risk of continuing treatment outweighs the benefit. Consider discontinuing naltrexone treatment sooner if opioid use is occurring towards the end of the extended-release naltrexone dosing interval as this places the patient at increased risk for fatal overdose.

  - **Alcohol use**: Patients presenting to clinic smelling of alcohol, positive breathalyzer, provides reports of ongoing ETOH use, or noted ED admissions for ETOH use and the risk of continuing treatment outweighs the benefit.

  - Ongoing use of other substances and the risk of continuing treatment outweighs the benefit.

  - Patient presents to OBAT clinic impaired or incidence of overdose, or hospitalization related to substance use and the risk of continuing treatment outweighs the benefit.

- Multiple missed appointments or inability to contact patient:

  - Initially if a patient misses an extended-release naltrexone injection, he/she should be instructed to receive the next injection as soon as possible. Reassess patient status prior to administering medication. Naloxone/naltrexone challenge if risk of precipitating withdrawal. Augment treatment plan as needed.

  - Multiple missed appointments should be addressed with the patient and the treatment team. Risk may outweigh benefits of continuing naltrexone treatment; document in electronic medical record.

- Patients who are referred to a higher level of care or discharged will be reconsidered for OBAT in the future.
OBAT DISCHARGE

- If a patient is discharged from OBAT they are welcome to re-engage, except if there are administrative or safety concerns connected with the discharge.

- Examples of administrative and safety issues: violence or criminal activity on hospital/clinic grounds; police report or other documentation of patient selling prescribed medication; inappropriate behavior in a clinic setting; threatening safety of staff or other patients.
SPECIFIC POPULATIONS

METHADONE TO BUPRENOPHRINE TRANSFERS

Potential benefits of transitioning to buprenorphine/naloxone:

- Decreased risk of overdose with partial agonist medication
- Opportunity to receive integrated care for OUD in a less restrictive, office based setting.

Considerations when transitioning from methadone to buprenorphine/naloxone:

- The transition from methadone to buprenorphine/naloxone can be very difficult for the patient and there is the risk of discomfort, destabilization and relapse. It is critical that OBAT NCM works with methadone clinic staff to coordinate the methadone taper and transition to buprenorphine/naloxone. The patient must be supported during the process.

- Establish with both patient and methadone clinic that if the transition to buprenorphine/naloxone is unsuccessful (e.g. patient begins to experience withdrawal that interferes with functioning or leads to return to use, or patient does not tolerate the medication), the patient may return to methadone treatment without a gap in treatment.

- Educate patients regarding appropriate methadone dose levels for transferring to buprenorphine/naloxone. To decrease the level of physical opioid dependence and minimize the chance for precipitated withdrawal, most patients will need to have their dose tapered to 30mg before beginning buprenorphine/naloxone treatment.

Recommendations:

- A methadone dose of 20-30mg/day for one to two weeks prior to transition to buprenorphine/naloxone is optimal but not always necessary.

- Alternate approach: taper methadone dose to the point of patient discomfort; with objective withdrawal symptom documentation via COWS, buprenorphine/naloxone can then be initiated.

- Inpatient detoxification is another option to assist a patient in the transition from methadone to buprenorphine/naloxone.

- Advise patient to arrange for time off of work during the transition. Family support with childcare and other responsibilities may be necessary, as discomfort may last one to two weeks.
- It is not necessary to begin with buprenorphine mono-tablet (Subutex) before initiating buprenorphine/naloxone, provided that patient is in an adequate state of withdrawal.

- Timing for last methadone dose/first buprenorphine/naloxone dose is difficult to predict.
  - Generally, at least 36-96 hours after last methadone dose, but utilizing clinical assessment and judgment is essential.
  - Long half-life of methadone (storage in body tissues, especially liver) causes unpredictable clearance.
  - Initiation of buprenorphine/naloxone should be guided by withdrawal symptoms objectively documented with a COWS score of 13-15, rather than by time since last methadone dose.

- Close monitoring and small amounts of clonidine, anxiolytics (including benzodiazepines), sedative/hypnotics, bentyl, trazadone, and NSAIDs may be used to manage distressing withdrawal symptoms and continued during induction, if approved by provider.

- More intensive stabilization support may be needed, such as telephone contact up to three times daily until maintenance dosing is attained. Frequent visits, adequate supports, and a supportive environment can assist with the transition.

- Providers should be experienced in induction prior to transitioning a patient from methadone maintenance to buprenorphine.

- Having the patient go to an inpatient detoxification to make this transition can be a safer, more effective way to get the patient from methadone maintenance to buprenorphine/naloxone.

**Induction recommendations:**

- Once COWS score of 13-15 is documented, start buprenorphine/naloxone at 2mg/0.5mg sublingually as prescribed.

- Continue to dose patient as prescribed until physical withdrawal symptoms have been reduced to manageable levels or are absent. Patients transitioning from methadone may require higher dosing initially and then can taper down over time.

- Continue induction according to patient’s prescription order, assessing symptoms of withdrawal and cravings.

- Symptom management with adjunctive medications as appropriate with provider input.

- Support and access to providers is critical in assisting patients with a successful transition.
Buprenorphine to Naltrexone Transfers

Transitioning from Buprenorphine/Naloxone to Naltrexone

There have been several observation pilot studies conducted to explore the transition from buprenorphine to naltrexone. The vast majority were not randomized controlled trials.

- Per Mannelli et al. (2012), “Taken together, published clinical practice recommends induction to full dose naltrexone five to seven days after buprenorphine discontinuation... The studies we have reviewed here show the feasibility of transferring opioid dependent patients from buprenorphine to naltrexone in a shorter time, if an inpatient treatment option is available.”

- A study by Kosten et al. (1993) found that administration of very low-dose oral naltrexone (1mg) did not induce significant withdrawal in buprenorphine-treated opioid dependent individuals. In participants who discontinued buprenorphine/naloxone and were given naltrexone 1mg titrated to full dose, naltrexone maintenance could be initiated in about half with only a small proportion remaining in treatment after two weeks.

- Sigmon et al. (2009) conducted a pilot study of 15 opioid dependent individuals enrolled to complete buprenorphine/naloxone stabilization, a two-week buprenorphine/naloxone taper, and naltrexone induction once urine levels of buprenorphine/naloxone were undetectable. Overall, rates of abstinence were high during the stabilization and taper periods and decreased markedly following taper off of buprenorphine/naloxone. The authors concluded that while a two-week taper may be appropriate for a subset of individuals, it is unlikely to be sufficient for the majority of individuals with opioid use disorders.

- Inpatient study by Clark et al., a small group of heroin users and buprenorphine-treated patients tapered buprenorphine in two to four days, combined with increasing doses of naltrexone. Following buprenorphine discontinuation, patients received naltrexone 50mg and were discharged. Higher withdrawal discomfort was reported in the initial two days of treatment. All patients completed the protocol. Results: 33% of patients were still taking naltrexone after four weeks, but overall opioid use was reduced by 50% or more compared with treatment admission.

Benefits of transitioning from buprenorphine/naloxone to naltrexone:

- Naltrexone is a long-acting medication.

- Naltrexone tablets have a half-life of 14hrs and can/should be dosed on a once daily regimen.

- Extended-release injectable formulation is administered every 28 days. Patients receive one injection in the clinic every four weeks, thus reducing the need for self-discipline and the burden of daily medication dosing.
• Naltrexone indication for use includes BOTH prevention of relapse to opioids and assistance with treating alcohol use disorder.
  • Naltrexone mutes the reinforcing effects of alcohol.

• No opioid dependency.
  • Patients may choose to stop naltrexone treatment at any time without having to undergo opioid withdrawal.

• No psychoactive effects.

• Treatment is also provided within an established medical system with integration of addiction treatment alongside medical care with the ability to obtain FDA approved medications for opioid use disorder and alcohol use disorder.
  • Insurance may require use of a specialty pharmacy and prior authorization.

• Antagonist medications such as naltrexone accelerate the opioid agonist detoxification process and are often prescribed post-detoxification to help prevent relapse.

Considerations:

When transitioning from buprenorphine to naltrexone, work with current buprenorphine clinic staff to coordinate the buprenorphine taper with the transition to naltrexone:

• Establish with both patient and buprenorphine clinic that, if the transition to naltrexone fails (e.g. patient begins to experience withdrawal or cravings that interferes with functioning or leads to return to use, patient does not tolerate the medication), the patient may return to buprenorphine treatment without a gap in treatment.

• Long half-life of buprenorphine and slow dissociation for mu opioid receptor causes unpredictable clearance.
  • Timing for last buprenorphine dose/first naltrexone dose is difficult to predict.

  • The limited amount of available data suggests that patients may do best when tapered to 2-4mg of buprenorphine/naloxone daily for one week, waiting five to seven days between the last dose of buprenorphine/naloxone and the first dose of naltrexone, and then starting with low dose naltrexone by mouth.

  • The tapering and transitioning period will include discomfort and increased risk for relapse. Please support patients during this process.

  • Educate patients regarding appropriate buprenorphine dose levels for transferring to naltrexone. To decrease the level of physical opioid dependence and minimize
the chance for severe precipitated withdrawal, most patients will need to have their dose tapered to 2mg before beginning naltrexone treatment.

- Advise patient to arrange for time off work during the transition, family support with childcare and other responsibilities as discomfort may last several days.

- Initiation of naltrexone should be guided by patient motivation, clinical judgment, and UTS result negative for ALL opioids rather than by last buprenorphine dose, family pressure, or law enforcement desire for patient to be on antagonist treatment.

- Withdrawal signs and symptoms will occur, causing patient discomfort.
  
  - Intensive stabilization and support may be needed (e.g. telephone contact up to three times daily until free of withdrawal signs/symptoms and patient stable). Frequent visits, adequate supports, supportive environment to assist in the transition.

  - Clonidine, anxiolytics (including benzodiazepines), and NSAIDs may be used to manage distressing withdrawal symptoms and continued during induction if prescribed by provider and closely monitored.

- Begin with naltrexone tablets before administering extended-release injectable naltrexone.

**Suggested Buprenorphine to Naltrexone Protocol:**

- Patient to reduce daily buprenorphine dose to 2mg for one week.

- Establish last dose date with patient. Five to seven days after final buprenorphine dose, patient to come to clinic with naltrexone tablet prescription bottle for naltrexone induction appointment with OBAT NCM.

- UTS negative for all opioids.

- Negative naloxone/naltrexone challenge.
  
  - It is recommended to initiate naltrexone treatment with the oral naltrexone formulation versus the extended-release injectable formulation to mitigate allergic reactions, side-effects and adverse reactions.

- Symptom management with adjunctive medications to occur as appropriate with provider input.

- Support and access to providers is critical in assisting patients with making this transition and not jeopardizing relapse.
PATIENTS WITH HIV

- Naltrexone: almost no interaction with antiretroviral medications.

- Buprenorphine/naloxone: does not interfere with clinical response to antiretroviral medications.

- Side effects from drug interactions between HIV medications and buprenorphine/naloxone are less severe/significant than those experienced with methadone.

- Reassure patients that treatment for their opioid dependence will not interfere with treatment for their HIV disease management.

Considerations:

- Protease inhibitors may increase buprenorphine/naloxone levels; however, no clinically significant increases or toxicities have been observed, with a few exceptions:
  - Atazanavir and atazanavir/ritonavir have been found to cause significant increases in buprenorphine/naloxone levels, with subsequent sedation and cognitive impairment.
  - Decrease the buprenorphine/naloxone dosage until the symptoms disappear.

- Non-nucleoside reverse transcriptase inhibitors (NNRTIs) may decrease buprenorphine/naloxone levels and cause withdrawal symptoms.
  - May need to increase the buprenorphine/naloxone dose.

- Buprenorphine/naloxone may slightly increase protease inhibitor levels.

- Initiation of medication-assisted opioid treatment during HAART maintenance:
  - Clinical needs should determine treatment selection.
  - With opioid agonists, patients may benefit from a trial of buprenorphine/naloxone because of the more benign drug interaction profile of buprenorphine/naloxone compared with methadone.

Initiation of HAART during buprenorphine/naloxone maintenance:

- Continue usual buprenorphine/naloxone dose.

- Advise patient of possible side effects.

- Efavirenz (Sustiva): withdrawal symptoms. Increase buprenorphine/naloxone dose accordingly.
**Patients with Hepatitis C**

**Buprenorphine**

- Both buprenorphine and naloxone are extensively metabolized by the liver.

- Most recent guidelines indicate that there are minimal concerns co-managing HCV and opioid use disorders utilizing buprenorphine/naloxone.\(^{27}\)
  
  - Current data suggests that liver injury from buprenorphine occurs rarely, however patients with hepatitis C are at higher risk to elevations in transaminases and reversible hepatic injury. Most of the evidence suggests that these elevations are related to underlying liver disease and not buprenorphine exposure. Serious hepatic injury is rare.
  
  - Buprenorphine maintenance may have an indirect beneficial effect on liver health via reduction of illicit opioid use.

- A single-dose study of 43 patients compared buprenorphine/naloxone exposure in healthy individuals to persons with mild, moderate, or severe hepatic impairment. Study results indicate that individuals with more advanced hepatic impairment experience higher peak exposure levels of naloxone vs. buprenorphine when compared to healthy subjects.\(^{15}\)
  
  - Dose adjustment may be required for some patients with severe liver disease.
  
  - May consider mono-tablet in some cases of severe liver disease.

- There are a small number of case reports of intravenous use of buprenorphine/naloxone by patients with hepatitis C resulting in increased alanine aminotransferase levels to 30-50x normal.\(^{28}\)
  
  - Case reports of seven patients with hepatitis C using buprenorphine/naloxone who had increased ALT 39x normal.\(^{29}\)
    
    - All continued buprenorphine/naloxone; 50% dose reduction in three patients.
    
    - All recovered without any clinical complications.

- When initiating buprenorphine/naloxone treatment it is important to do baseline hepatic testing and then to retest transaminases as needed based on clinical assessment.
Naltrexone

- Naltrexone is extensively metabolized through the liver and clinical judgment should be used prior to administration in cases of advanced liver disease or acute hepatitis.

- AST and ALT should both be less than 5x the upper limit of normal at treatment initiation.

- Draw follow-up AST and ALT eight to 12 weeks after initiation of naltrexone. At present there is no empirical evidence to support frequency of monitoring; clinical discretion should be used to guide frequency.27

  - Cases of hepatitis and clinically significant liver dysfunction were observed in association with extended-release injectable naltrexone treatment during the clinical development program and in the post-marketing period.

  - A randomized, double-blind, placebo-controlled trial of 624 individuals with alcohol dependence (DSM-IV) and recent heavy drinking, designed to assess the hepatic safety of injectable naltrexone found no difference in hepatic function at six months between participants on injectable naltrexone at the FDA approved dose (380mg) compared to those receiving a placebo.30

  - In a study of 250 participants (89% had history of HCV) at six month follow-up, elevations in AST, ALT, and GGT greater than three times the upper limit of normal were not statistically different in patients treated with injectable naltrexone compared with the placebo.31 The majority of participants with liver enzyme level elevations greater than three times the upper limit of normal had chronic HCV infection.

  - Discontinue use of extended-release injectable naltrexone in the event of symptoms or signs of acute hepatitis (e.g. abdominal pain, nausea, vomiting, fever, dark urine, clay-colored stools, jaundice, or icterus; or ALT or AST levels greater than 10x the upper limit of normal).27

    - If there is no evidence that liver enzyme elevation is related to medication, NCM or provider can restart once ALT and AST levels fall below 10x the upper limit of normal.

- For all patients prescribed either buprenorphine/naloxone or naltrexone, hepatic enzymes should be monitored at regular intervals throughout the course of treatment.

- Patients should receive education about the signs/symptoms of liver inflammation and be advised to report these signs/symptoms to the clinical team or present to emergency department for evaluation if they are present.
**PREGNANCY AND BREASTFEEDING**

**Opioid use disorder in pregnancy is considered high-risk.**

- Educate pregnant patients on the benefits of maintaining opioid replacement during pregnancy.
  
  - Decreased risk for relapse and therefore reduced complications from illicit opioid use.
  
  - Constant levels of fetal opioid exposure result in reduced risk for adverse fetal outcomes related to multiple withdrawals.
  
  - Decreased rate of adverse fetal outcomes such as low birth weight or preterm delivery.

- Incidence of neonatal abstinence syndrome is 47%.
  
  - To date, research indicates that there is no significant correlation between buprenorphine dose and the incidence of NAS. Therefore, it is most important to support and stabilize the mother on a dose of buprenorphine that eliminates or minimizes opioid craving and withdrawal.

- Both methadone and buprenorphine (both combo and mono-tablet formulations) are Category C in pregnancy.

- There is more data and clinical experience utilizing methadone in pregnancy.

- In 2012, the American College of Obstetricians and Gynecologists (ACOG) concluded that there is evidence to support the use of buprenorphine as a potential first-line medication for opioid dependent women.

- A longitudinal study of 73 children evaluated at 24 months (n = 24 exposed to buprenorphine in utero, and n = 19 exposed to methadone in utero, n = 30 non-exposed controls) found no differences between groups in temperament or neurological development during the first two years of life.\(^{32}\)

- A double blind randomized controlled trial of 175 pregnant women with opioid use disorder treated with buprenorphine or methadone maintenance compared maternal and neonatal outcomes between the two groups. A total of 131 neonates were born to mothers followed through the end of their pregnancy (58 exposed to buprenorphine and 73 exposed to methadone).\(^{33}\)
Neonates in the buprenorphine group required significantly less morphine (mean dose, 1.1 mg vs. 10.4 mg) than neonates in the methadone group. They also had significantly shorter hospital stays (10.0 days vs. 17.5 days) and significantly shorter duration of treatment for neonatal abstinence syndrome (4.1 days vs. 9.9 days). The two groups did not vary with regard to maternal or neonatal adverse events.

**Buprenorphine in Pregnancy:** Due to a lack of safety data on buprenorphine/naloxone maintenance in pregnancy it is typical that pregnant women with opioid use disorder are either started on methadone or the buprenorphine mono product or are switched from buprenorphine/naloxone to the mono product (buprenorphine) or methadone.

- For several decades in the absence of additional safety data, two principles have guided the recommendation to use the mono product over buprenorphine/naloxone: (i) pregnant women should limit exposure to exogenous compounds, and (ii) animal studies have suggested the possibility that naloxone could cause maternal and fetal hormonal changes.

- Additional research is still needed; a recent review comprised of preliminary findings from seven previously published studies found no evidence of adverse maternal or neonatal outcomes related to use of buprenorphine/naloxone compared to buprenorphine alone (mono product) or methadone. Currently, many providers use buprenorphine/naloxone for the treatment of opioid use disorder during pregnancy without complication or notable adverse events.

**Buprenorphine Protocol for Pregnant Women**

- Provide smaller prescriptions to limit diversion potential and promote safety.
- Schedule more frequent follow-up visits during pregnancy.
- Refer for high-risk obstetric service if available.
- Minimal information exists on dosing changes by trimester.
- Many require divided dosing during pregnancy.
- Frequent follow up visits should include: assessment, support, UTS, safety assessment, counseling, education and assessing social determinants of health.
- Women should be encouraged to breastfeed provided their UTS are negative for opioids and the mother is not prescribed any other medications that are contraindicated for breastfeeding.
- Breastfeeding women should be maintained on buprenorphine/naloxone.
- Buprenorphine/naloxone is passed into breast milk at 1:1 plasma: milk ratio.
- Because of poor oral bioavailability of buprenorphine/naloxone, the breastfeeding infant is exposed to only 1/10 of buprenorphine/naloxone ingested.

- Breastfeeding during buprenorphine/naloxone use does not suppress neonatal abstinence syndrome. However the close contact afforded by breastfeeding has been shown to assist with symptoms of NAS and enhances maternal-child bonding.

- Cessation of breastfeeding is not associated with onset of neonatal abstinence syndrome.

- **Naltrexone in Pregnancy:** Little research has been conducted to evaluate the safety or efficacy of naltrexone in pregnancy. Naltrexone, in both oral and injectable formulations, is considered a Category C medication.

- **Naltrexone with Breastfeeding:**\(^3\) It is not known if extended-release injectable naltrexone passes into breast milk. It is known that oral naltrexone does pass into breast milk. Due to the potential tumorigenicity shown for naltrexone in animal studies, and because of the serious adverse reactions in nursing infants from injectable naltrexone, a decision should be made to either discontinue the medication or discontinue breastfeeding. Labeling from the manufacturer advises against breastfeeding while taking naltrexone, both with oral and injectable formulations.

**For additional guidance regarding the care of pregnant women with opioid use disorder:**

**SAMHSA.**

*Clinical Guidance for Treating Pregnant and Parenting Women With Opioid Use Disorder and Their Infants.*

This Clinical Guide provides comprehensive, national guidance for management of pregnant and parenting women with opioid use disorder and their infants. The guide helps healthcare professionals and patients determine the most clinically appropriate action for a particular situation and informs individualized treatment decisions.

**Pub id:** SMA18-5054  **Publication Date:** 1/2018

[https://store.samhsa.gov/product/SMA18-5054](https://store.samhsa.gov/product/SMA18-5054)

\(^3\) Adapted from information accessed at: https://www.vivitrol.com/important-safety-information
DUAL DIAGNOSIS

BUPRENORPHINE/NALOXONE

- Buprenorphine/naloxone is metabolized in the liver by the cytochrome P450 3A4 system.

- Clinical experience has not uncovered significant drug-drug interactions with buprenorphine/naloxone.

- Dosing changes are generally not necessary, as opposed to methadone dosing, which is highly influenced by concomitant medication use.

- Reassure patients with comorbid psychiatric conditions that use of buprenorphine/naloxone is not a barrier to treatment of their psychiatric condition.

NALTREXONE

- The cytochrome P450 system is not involved in naltrexone metabolism. In vitro CYP studies have demonstrated that naltrexone is not an inhibitor or inducer of major CYP enzymes.

Dual Diagnosis Treatment in OBAT

- All patients are assessed for psychiatric disorders as a component of OBAT screening procedures.

- After two to three weeks of stabilization, reassess patients for psychiatric symptomatology.

- Substance-induced psychiatric disorders generally resolve within one to two weeks of treatment initiation and cessation of substance use.

- Psychiatric symptoms that persist beyond 30 days after cessation of substance use are suggestive of an independent psychiatric condition. These patients should be offered a referral to behavioral health services for a mental health evaluation.

  - For patients engaged in psychiatry services, obtain patient-signed CFR42 consent for release of information to facilitate coordination of care with mental health providers.

  - Benzodiazepines should be used cautiously with patients receiving buprenorphine/naloxone because of the potential for increased CNS depression, including sedation, respiratory depression and the potential for misuse in the patient with the disease of addiction. Patient history of benzodiazepine misuse should also be explored prior to prescribing.
PAIN MANAGEMENT PROTOCOL: BUPRENORPHINE/NALOXONE

BUPRENORPHINE/NALOXONE PATIENTS REQUIRING SURGERY

Background: These guidelines are designed for patients maintained on buprenorphine or buprenorphine/naloxone undergoing invasive procedures. There is currently a lack of evidence-based studies to direct the management of patients on buprenorphine/naloxone maintenance during the peri-procedure period. Below are guidelines using expert opinions based on pharmacological principles with the intent to avoid under-treatment of acute pain while also avoiding potential opioid withdrawal and disruption of opioid use disorder treatment.

The appropriate treatment of acute pain in patients on buprenorphine/naloxone maintenance includes continuing the patient’s baseline opioid requirements to avoid increased pain sensitivity associated with opioid withdrawal. Thus, daily opioid maintenance treatment requirements must be met before attempting to achieve analgesia. These patients have also been shown to have increased pain sensitivity and cross-tolerance to opioid analgesics, therefore adequate pain control may require higher opioid doses at shorter dosing intervals. All patients on buprenorphine/naloxone maintenance should be co-managed with their buprenorphine/naloxone provider during the pre- and post-procedure period. 35
BUPRENORPHINE: PERI-PROCEDURE MANAGEMENT

RECOMMENDATIONS:

- Daily buprenorphine/naloxone dosing should remain uninterrupted. Patient takes usual buprenorphine/naloxone maintenance dose on the morning of procedure.
  - Because of its high affinity at the opioid receptor, consider fentanyl as the opioid of choice for analgesia during procedures and in PACU for these patients.
- Continue patient’s home dose of buprenorphine/naloxone post-operatively.
  - Consider splitting the patient’s usual buprenorphine/naloxone dose into every eight hour dosing (e.g. 24 mg per day changed to 8mg every 8 hours)
- If further pain control is needed, begin by utilizing multimodal pain management with non-opioids (NSAIDs, acetaminophen, lidocaine patches, etc.).
- Consider the use of local and regional anesthesia as indicated.
- If opioids are needed for breakthrough pain, standard dosing protocols should initially be utilized with careful monitoring and the understanding that patients with a history of OUD may require higher than usual doses due to cross tolerance and increased pain sensitivity.
- Patient Controlled Analgesia’s (PCA) without a basal component may be considered in addition to a patient’s buprenorphine if pain is not adequately captured. If a PCA is utilized, discontinue oral PRN opioids.
- The buprenorphine/naloxone provider should be contacted pre-and post-procedure to assist in ongoing assessment, support, and pain management.
- Schedule patient to be seen by their buprenorphine/naloxone prescriber within one week post-procedure.
BUPRENORPHINE: ACUTE AND CHRONIC PAIN MANAGEMENT

General principles for pain management on buprenorphine/naloxone:

- Patients physically dependent on opioids require maintenance on daily equivalence before any pain relief is achieved with opioid analgesics (the “opioid debt”).

- Evidence-based data now supports continuing patients on their daily maintenance dose of buprenorphine/naloxone during periods of acute pain, rather than discontinuing and later restarting buprenorphine treatment. Maintaining buprenorphine/naloxone has been shown to increase pain control while allowing the patient to remain stabilized on their medication treatment for OUD.

- Reassure patients that their substance use disorder will not be an obstacle to aggressive pain management.

- It is important to include patients in the decision-making process to allay anxiety.

- Establish clear goals for pain management:
  - Pain reduction rather than elimination.
  - Improved function.

- Use multimodal approach to pain management:
  - Consider splitting the patient’s usual buprenorphine/naloxone dose into every eight hour dosing (e.g. 24 mg per day changed to 8mg every 8 hours).
  - Consider modest increase in patient’s buprenorphine/naloxone maintenance dose.
  - Try non-opioids and adjuvant therapies next.
    - Examples include: acupuncture, acupressure, massage, physical therapy, hydrotherapy, mindful meditation, NSAIDs, acetaminophen, topical lidocaine, SSRIs, TCAs, etc.
  - If multimodal therapies do not effectively reduce pain, opioid analgesics are an option.
  - If opioid analgesics are necessary for treatment of chronic pain, buprenorphine/naloxone should be discontinued and methadone maintenance initiated.
Sampling of the Evidence:

- Macintyre et al., (2013) performed a retrospective cohort study comparing pain relief and opioid requirements in the first 24 hours after surgery in 22 patients maintained on buprenorphine and 29 patients maintained on methadone, who were also prescribed patient-controlled analgesia. The study found no significant differences in pain scores, incidence of nausea or vomiting requiring treatment, or sedation between the buprenorphine or methadone maintained patient groups overall. Additionally it was found that buprenorphine maintained patients who were not given their usual buprenorphine dose the day after surgery used significantly more patient-controlled analgesia (P=0.02) compared with those who had received their dose.36

- Kornfield and Manfredi (2010) performed a literature review examining five buprenorphine-maintained patients who underwent seven planned major surgical procedures with high levels of anticipated post-operative pain (right-side colectomy, small bowel resection, L and R knee replacements, bilateral mastectomy, breast reconstruction, X-Stop procedure). In all seven cases, daily buprenorphine maintenance dosing was uninterrupted. Full agonist opioids and non-opioid analgesics were used in conjunction with daily buprenorphine dosing. In all seven surgical cases, good to excellent pain control was achieved.37

- Silca & Rubenstein (2016) presented a case comparing two different outcomes for the same surgical course performed at two different times on the same chronic pain patient. Results showed that pain control was easier to achieve, and functional recovery was greater, when buprenorphine was maintained throughout the perioperative period compared to using a full mu agonist opioid for chronic pain perioperatively.38
PAIN MANAGEMENT PROTOCOL: NALTREXONE

Background:

- These guidelines are designed for patients maintained on naltrexone undergoing invasive procedures. There is currently a lack of evidence-based research to direct the management of patients prescribed naltrexone in the peri-procedure period. Below are guidelines using expert opinions based on pharmacological principles of naltrexone with the intent to avoid under-treatment of acute pain while also avoiding potential disruption of opioid use disorder and/or alcohol use disorder treatment.

- The pain-relieving effects of opioid agonists are blocked while on naltrexone. This includes pure mu agonists, such as methadone or morphine derivatives, partial agonists, as well as mixed agonist/antagonists. In order to overcome the pharmacologic blockade of extended-release injectable naltrexone, extremely high doses of opioids are required to achieve adequate analgesia. This could lead to accidental overdose. It is therefore recommended that non-opioid analgesics be prescribed for pain management in these patients when possible. Non-steroidal anti-inflammatory agents are first-line. Regional nerve blocks and dissociative analgesics such as ketamine have also been recommended. However, expert consultation by an informed, experienced pain specialist should occur.\(^{39}\)

- All OBAT patients receiving naltrexone treatment should be co-managed with their OBAT provider during the pre- and post-procedure period, as well as during periods of acute and chronic pain.
OBAT POLICY FOR NALTREXONE PATIENTS REQUIRING SURGERY:

- Patient to notify OBAT staff of expected procedure ASAP.

- Obtain signed consent for release of information with CFR42 for the surgical/medical team.
  - OBAT team to work with surgical team to manage pre- and post-procedure pain.

- If possible, extended-release naltrexone should be discontinued four to five weeks prior to the planned surgery/procedure date.
  - May bridge patient with oral naltrexone.
  - Oral naltrexone should be discontinued 48-72 hours before the procedure.\(^{38}\)

- Before minor or intermediate elective surgery, the possibility of managing the pain with non-opioids needs to be balanced against the risk of the patient relapsing.

- If a patient is to undergo major surgery where severe post-operative pain is expected, oral naltrexone should be discontinued 72 hours in advance. A degree of resistance to opioid analgesics should be expected, although increased sensitivity is also a possibility.

- Patients should be monitored closely with increased supports throughout the peri-procedure period.

- Patients should be scheduled to see their OBAT provider as soon as possible to have their post-procedure pain managed and to be safely restarted on naltrexone when it is safe to do so.
NALTREXONE: CHRONIC PAIN MANAGEMENT

- Chronic pain requiring opioid medications is a contraindication for naltrexone and should be evaluated as part of OBAT screening process.

- For patients seeking medication treatment for an opioid use disorder who also have severe chronic pain, agonist medications should be considered, including buprenorphine/naloxone or methadone maintenance therapy.

General principles for patients engaged in naltrexone treatment who develop chronic pain:

- Include patients in decision-making process to allay anxiety.

- Establish clear goals for pain management:
  - Pain reduction rather than elimination.
  - Improved function.

- Use multimodal approach to pain management:
  - Try non-opioids initially.
  - Try adjuvant therapies next.

- If pain relief is not sufficient, discuss with patient transfer for agonist therapy: methadone or buprenorphine/naloxone.
NALTREXONE: UNANTICIPATED ACUTE PAIN MANAGEMENT

- If a patient receiving ongoing extended-release naltrexone injections experiences unanticipated severe, acute pain or requires emergent surgery, refer to “Reversal of Extended-Release Injectable Naltrexone.”
- For patients taking oral naltrexone with unanticipated acute pain:
  - Include patients in decision-making process to allay anxiety.
  - Address underlying cause of pain.
- Establish clear goals for pain management:
  - Pain reduction rather than elimination.
  - Improved function.
- Use multimodal approach to pain management:
  - Try non-opioids initially.
  - Try adjuvant therapies next.
- For patients prescribed a formulation of naltrexone, if opioid analgesics are absolutely necessary for treatment of unanticipated acute pain, naltrexone should be discontinued.
- If this occurs, higher than usual doses of opioids may be attempted to overcome naltrexone’s opioid antagonist effects.
- Prescribing opioids to a patient that has been maintained on a formulation of naltrexone must be done with close observation for respiratory depression. Refer to “Reversal of Extended-Release Injectable Naltrexone.”
- Patients should be monitored closely with increased supports throughout the acute pain period.
In an emergency situation for patients receiving extended-release injectable naltrexone, suggestions for pain management include regional analgesia or use of non-opioid analgesics.

- If opioid therapy is required as part of anesthesia or analgesia, patients should be continuously monitored in an anesthesia care setting by persons not involved in the conduct of the surgical or diagnostic procedure.

- The opioid therapy must be provided by individuals specifically trained in the use of anesthetic drugs and the management of the respiratory effects of potent opioids, specifically the establishment and maintenance of a patent airway and assisted ventilation.

- Irrespective of the drug chosen to reverse the extended-release injectable naltrexone blockade, the patient should be monitored closely by appropriately trained personnel in a setting equipped and staffed for cardiopulmonary resuscitation.

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4 Adapted from: https://www.accessdata.fda.gov/drugsatfda_docs/label/2010/021897s015lbl.pdf
### APPENDICES

**APPENDIX 1: DSM-5 CHECKLIST OF DIAGNOSTIC CRITERIA: OPIOID USE DISORDER**

A problematic pattern of opioid use leading to clinically significant impairment or distress, as manifested by at least 2 of the following, occurring within a 12-month period:

<table>
<thead>
<tr>
<th>Diagnostic Criterion</th>
<th>Meets Criterion?</th>
<th>Additional/ Supporting Information</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Opioids are often taken in larger amounts or over a longer period than was intended.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. There is a persistent desire or unsuccessful efforts to cut down or control opioid use.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. A great deal of time is spent in activities necessary to obtain the opioid, use the opioid, or recover from its effects.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Craving, or a strong desire or urge to use opioids.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. Recurrent opioid use resulting in a failure to fulfill major role obligations at work, school, or home.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Continued opioid use despite having persistent or recurrent social or interpersonal problems caused or exacerbated by the effects of opioids.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7. Important social, occupational, or recreational activities are given up or reduced because of opioid use.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8. Recurrent opioid use in situations in which it is physically hazardous.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9. Continued opioid use despite knowledge of having a persistent or recurrent physical or psychological problem that is likely to have been caused or exacerbated by the substance.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10. Tolerance, as defined by either of the following:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>A. A need for markedly increased amounts of opioids to achieve intoxication or desired effect.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>B. A markedly diminished effect with continued use of the same amount of an opioid.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>11. Withdrawal, as manifested by either of the following:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>A. The characteristic opioid withdrawal syndrome</td>
<td></td>
<td></td>
</tr>
<tr>
<td>B. Opioids (or a closely related substance) are taken to relieve or avoid withdrawal symptoms.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Note:** This criterion is not considered to be met for those individuals taking opioids solely under appropriate medical supervision.

**Specify if:**

- **In early remission:** After full criteria for opioid use disorder were previously met, none of the criteria for opioid use disorder have been met for at least 3 months but for less than 12 months (with the exception that Criterion 4, “Craving, or a strong desire or urge to use opioids,” may be met).
- **In sustained remission:** After full criteria for opioid use disorder were previously met, none of the criteria for opioid use disorder have been met at any time during a period of 12 months or longer (with the exception that Criterion 4, “Craving, or a strong desire or urge to use opioids,” may be met).
- **On maintenance therapy:** This additional specifier is used if the individual is taking a prescribed agonist medication such as methadone or buprenorphine and none of the criteria for opioid use disorder have been met for that class of medication (except tolerance to, or withdrawal from, the agonist). This category also applies to those individuals being maintained on a partial agonist, an agonist/antagonist, or a full antagonist such as oral naltrexone or depot naltrexone.
- **In a controlled environment:** This additional specifier is used if the individual is in an environment where access to opioids is restricted.

**Current severity:**

- **Mild:** Presence of 2–3 symptoms. **Code as:** F11.10 (ICD-10)
- **Moderate:** Presence of 4–5 symptoms. **Code as:** F11.20 (ICD-10)
- **Severe:** Presence of 6 or more symptoms. **Code as:** F11.20 (ICD-10)
## Appendix 2: DSM-5 Checklist of Diagnostic Criteria: Alcohol Use Disorder

A problematic pattern of alcohol use leading to clinically significant impairment or distress, as manifested by at least two of the following, occurring within a 12-month period:

<table>
<thead>
<tr>
<th>Diagnostic Criterion</th>
<th>Meets Criterion?</th>
<th>Additional/Supporting Information</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Alcohol is often taken in larger amounts or over a longer period than was intended.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. There is a persistent desire or unsuccessful efforts to cut down or control alcohol use.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. A great deal of time is spent in activities necessary to obtain alcohol, use alcohol, or recover from its effects.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Craving, or a strong desire or urge to use alcohol.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. Recurrent alcohol use resulting in a failure to fulfill major role obligations at work, school, or home.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Continued alcohol use despite having persistent or recurrent social or interpersonal problems caused or exacerbated by the effects of alcohol.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7. Important social, occupational, or recreational activities are given up or reduced because of alcohol use.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8. Recurrent alcohol use in situations in which it is physically hazardous.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9. Alcohol use is continued despite knowledge of having a persistent or recurrent physical or psychological problem that is likely to have been caused or exacerbated by alcohol.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10. Tolerance, as defined by either of the following:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>A. A need for markedly increased amounts of alcohol to achieve intoxication or desired effect.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>B. A markedly diminished effect with continued use of the same amount of alcohol.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>11. Withdrawal, as manifested by either of the following:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>A. The characteristic withdrawal syndrome for alcohol (refer to Criteria A and B of the criteria set for alcohol withdrawal, pp. 499–500).</td>
<td></td>
<td></td>
</tr>
<tr>
<td>B. Alcohol (or a closely related substance, such as a benzodiazepine) is taken to relieve or avoid withdrawal symptoms.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Specify if:

- **In early remission:** After full criteria for alcohol use disorder were previously met, none of the criteria for alcohol use disorder have been met for at least 3 months but for less than 12 months (with the exception that Criterion 4, “Craving, or a strong desire or urge to use alcohol,” may be met).
- **In sustained remission:** After full criteria for alcohol use disorder were previously met, none of the criteria for alcohol use disorder have been met at any time during a period of 12 months or longer (with the exception that Criterion 4, “Craving, or a strong desire or urge to use alcohol,” may be met).
- **In a controlled environment:** This additional specifier is used if the individual is in an environment where access to alcohol is restricted.

**Current severity:**

- **Mild:** Presence of 2–3 symptoms. **Code as:** F10.10 (ICD-10)
- **Moderate:** Presence of 4–5 symptoms. **Code as:** F10.20 (ICD-10)
- **Severe:** Presence of 6 or more symptoms. **Code as:** F10.20 (ICD-10)
APPENDIX 3: TELEPHONE SCREENING

Demographic Info

How did you hear about the hotline?

☐ 1 = Spouse        ☐ 2 = Friend        ☐ 3 = Medical Provider
☐ 4 = Flyer         ☐ 5 = Parent         ☐ 6 = Hotline
☐ 7 = Physician Locator ☐ 8 = Other: ____________

Are you pregnant at this time?

☐ 1 = Yes
☐ 2 = No
☐ 3 = Don’t Know
☐ 4 = Tubal ligation
☐ 5 = Menopause
☐ 6 = History of hysterectomy
☐ 7 = Other

If no, are you on a form of contraception?  ☐ 1 = Yes  ☐ 2 = No

Current Address__________________________________________________________

Phone _______________  Is it ok to leave a message?  ☐ 1 = Yes  ☐ 2 = No

Phone _______________

Emergency Contact ____________________________  Phone ________________

Is the Emergency Contact aware of your addiction?  ☐ 1 = Yes  ☐ 2 = No
### Substance Use History

<table>
<thead>
<tr>
<th>What is your substance of choice?*</th>
<th>Age of initiation</th>
<th>Date of most recent use</th>
<th>Frequency?</th>
<th>Route of administration</th>
<th>Amounts used</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 = heroin</td>
<td>0 IF NEVER</td>
<td>1=12 OR MORE MONTHS AGO</td>
<td>1=LESS THAN</td>
<td>1=ORAL</td>
<td>1=ORAL</td>
</tr>
<tr>
<td>2 = fentanyl</td>
<td>NEVER USED</td>
<td>(SPECIFY DATE)</td>
<td>1/MONTH</td>
<td>2=SMOKING</td>
<td>2=SMOKING</td>
</tr>
<tr>
<td>3 = oxycodone product</td>
<td>2=3-11 MONTHS AGO</td>
<td>2=1-3 TIMES/MONTH</td>
<td>3=INTRANASAL</td>
<td>3=INTRANASAL</td>
<td>3=INTRANASAL</td>
</tr>
<tr>
<td>4 = buprenorphine/naloxone</td>
<td>3=1-2 MONTHS AGO</td>
<td>3=1-2 TIMES/WEEK</td>
<td>4=INTRAVENOUS</td>
<td>4=INTRAVENOUS</td>
<td>4=INTRAVENOUS</td>
</tr>
<tr>
<td>5 = methadone</td>
<td>4=1-3 WEEKS AGO</td>
<td>4=3-6 TIMES/WK</td>
<td>5=Skin Popping</td>
<td>5=Skin Popping</td>
<td>5=Skin Popping</td>
</tr>
<tr>
<td>6 = other opioid</td>
<td>5= USED THIS WEEK</td>
<td>5=DAILY</td>
<td>6=Other</td>
<td>6=Other</td>
<td>6=Other</td>
</tr>
<tr>
<td>7 = benzodiazepines</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8 = alcohol</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>9 = cocaine</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10 = amphetamines</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>11 = tobacco</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>12 = other</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**What substances are you currently using at this time?**
Includes age of first use, date of most recent use, route, frequency, and quantity.

- 1 = heroin
- 2 = fentanyl
- 3 = buprenorphine/naloxone
- 4 = methadone
- 5 = oxycodone product
- 6 = other opioid
- 7 = cocaine
- 8 = benzodiazepines
- 9 = Alcohol
- 10 = Amphetamines
- 11 = Other
- 12 = Nothing
What substances have you used in the past?
Includes age of first use, date of most recent use, route, frequency, and quantity.

- □ 1 = heroin
- □ 2 = fentanyl
- □ 3 = buprenorphine/naloxone
- □ 4 = methadone
- □ 5 = oxycodone product
- □ 6 = other opioid
- □ 7 = cocaine
- □ 8 = benzodiazepines
- □ 9 = Alcohol
- □ 10 = Amphetamines
- □ 11 = Other
- □ 12 = Nothing

Have you ever shared needles? □ 1= Yes □ 2 = No

Have you ever belonged to the needle exchange program? □ 1= Yes □ 2 = No

Have you ever overdosed □ 1= Yes □ 2 = No

Number of lifetime overdoses: ______

Have you ever been hospitalized due to an overdose? □ 1= Yes □ 2 = No

Was naloxone administered? □ 1= Yes □ 2 = No

**Recovery History**

What was the longest period of time that you have been in recovery?
______________________________

When was this? _________________

**Addiction Treatment History**

Have you ever engaged in treatment for a substance use disorder? □ 1= Yes □ 2 = No

If yes, how many times to each type?

- ____ Detoxification Program
- ____ Driving Impaired Program
- ____ Residential (Rehab or Halfway House)
- ____ Methadone Maintenance
- ____ Buprenorphine/naloxone maintenance
- ____ Intensive Outpatient Program
- ____ Naltrexone (oral or injectable)

Do you attend peer support meetings (check all that apply):
☐ 1= AA
☐ 2= NA
☐ 3= Smart Recovery
☐ 4= Other _____________________

How many meetings do you attend each week?
☐ 1 = 1-2 week
☐ 2 = 3-4 week
☐ 3 = 5-6 week
☐ 4 = None
☐ 5 = Other:

Do you have a sponsor?  ☐ 1= Yes   ☐ 2 = No

Do you have any history of any other addictive behaviors such as?
☐ 1 = Gambling
☐ 2 = Sex
☐ 3 = Shopping
☐ 4 = Eating disorder (over eating, bulimia, anorexia)
☐ 5 = Other:
☐ 6 = No

Comments: ____________________________________________________________

Criminal History

Have you ever been incarcerated?  ☐ 1= Yes   ☐ 2 = No

What is the longest period of time you spent in jail/prison? _________

Are you on probation?  ☐ 1= Yes   ☐ 2 = No

Are you on parole?  ☐ 1= Yes   ☐ 2 = No

Are you facing any potential jail time?  ☐ 1= Yes   ☐ 2 = No

Do you have any outstanding legal issues?  ☐ 1= Yes   ☐ 2 = No

If yes, can you tell us about them? _____________________________

Methadone History

Have you ever engaged in a Methadone Maintenance program?  ☐ 1= Yes   ☐ 2 = No

Are you currently on Methadone Maintenance?  ☐ 1= Yes   ☐ 2 = No
If yes to currently on engaged in Methadone treatment:

Where are you engaged in Methadone Maintenance? __________________________
What is the name of your counselor at your Methadone clinic? _________________
How long have you been in your current Methadone Maintenance Program? _____
Are you receiving take-homes? □ 1 = Yes □ 2 = No
If yes, how many? __________

If not currently engaged in methadone treatment:

When were you on Methadone Maintenance? _________________________________
Where were you on Methadone Maintenance? _________________________________
How long were you on Methadone Maintenance? _______________________________
What was your dose? ______
Why did you stop Methadone treatment? ______________________________

_Buprenorphine History_

Have you ever been prescribed buprenorphine/naloxone before?
□ 1 = Yes □ 2 = No
If yes:
  Where were you prescribed buprenorphine/naloxone:_______________________
  When were you prescribed buprenorphine/naloxone? _________________________
  What was your dose? ______________
  Why did you stop taking buprenorphine/naloxone? _________________________

_Naltrexone History_

Have you ever been prescribed naltrexone before?
□ 1 = Yes □ 2 = No
If yes:
  Where were you prescribed naltrexone:___________________________________
When were you prescribed naltrexone? ____________________

Did you ever receive an extended-release naltrexone injection? ________________

Why did you stop naltrexone treatment? ________________________________

**Mental Health History**

Are you currently seeing a psychiatrist, psychologist or counselor for a mental health issue?
☐ 1=Yes  ☐ 2= No

Where do you see your psychiatrist, psychologist or counselor? ________________

What is this individual’s name? __________________________

How often do you see them? ________________

How many times have you seen this person in the last six months? _________ Times.

Are you willing to sign a consent for release of information so that we can communicate with your psychiatrist, psychologist or counselor about your treatment plan?
☐ 1=Yes  ☐ 2=No

Have you ever been hospitalized for mental health issues?
☐ 1=Yes  ☐ 2=No

Have you ever attempted to end your life or to hurt yourself?
☐ 1=Yes  ☐ 2=No

How many times did you try to end your life or to hurt yourself? _________

Do you currently have thoughts about hurting yourself or ending your life?
☐ 1=Yes  ☐ 2=No (If no, skip to homicide question)

    **If yes:**
    Do you currently have a plan for how you would hurt yourself or end your life?
    ☐ 1=Yes  ☐ 2=No

    Do you have the means to carry out your plan?
    ☐ 1=Yes  ☐ 2=No

Have you ever attempted or thought about homicide (killing someone else)?
☐ 1=Yes  ☐ 2=No (If no, skip to health status)

    **If yes:**
Are you presently thinking about killing someone?
☐ 1=Yes  ☐ 2=No

Do you have the means to carry this out?
☐ 1=Yes  ☐ 2=No

Are you willing to Contract for Safety, call 911 etc., per program protocol…

Health status

Have you ever been diagnosed with any medical conditions? Mark all that apply.
☐ 1=Diabetes (specify type): ________________________________
☐ 2=Heart disease (specify type): ____________________________
☐ 3=Cancer (specify type): _________________________________
☐ 4=Asthma
☐ 5=Tuberculosis (TB)
☐ 6=Endocarditis
☐ 7=Skin infection
☐ 8= HIV  ➔ If yes, are you currently in care? ☐ 1= Yes  ☐ 2 = No
☐ 9= Hepatitis A
☐ 10= Hepatitis B  ➔ If yes, have you been treated? ☐ 1= Yes  ☐ 2 = No
☐ 11= Hepatitis C  ➔ If yes, have you been treated? ☐ 1= Yes  ☐ 2 = No
☐ 12= Seizure disorder  ➔ Are you on medications? ☐ 1= Yes  ☐ 2 = No
☐ 13= Head Trauma/Brain Injury
☐ 14= Pancreatic Problems
☐ 15= Other (specify type): ________________________________
☐ 16= None

Have you been tested for HIV?  ☐ 1= Yes  ☐ 2 = No

If yes, did you go back for the results?  ☐ 1= Yes  ☐ 2 = No

If yes, when was the last time you were tested?

Have you ever had surgery?  ☐ 1= Yes  ☐ 2 = No

If yes, why did you have surgery? ______________________________

Do you have any pending surgeries?  ☐ 1= Yes  ☐ 2 = No

If yes, please briefly explain: _________________________________

Pain

Do you have chronic pain?  ☐ 1 = Yes  ☐ 2 = No
Please rate your pain, on a scale from 0 – 10, without any pain medications (prescribed or bought on the street)

____0____1____2____3____4____5____6____7____8____9____10____

Please rate your pain, on a scale from 0 – 10, WITH pain medications (prescribed or bought on the street)

____0____1____2____3____4____5____6____7____8____9____10____

**Health Care Provider Information**

Where do you get most of your healthcare? ____________________________

When was the last time you saw a health care provider?

- □ 1 = Last week
- □ 2 = Last month
- □ 3 = Within the past 3 months
- □ 4 = Within the past 6 months
- □ 5 = Within the past year
- □ 6 = More than 1 year ago

What is the name of your provider? ____________________________

**Employment**

Are you currently employed? □ 1 = Yes □ 2 = No

If yes, what do you do for work? __________________________________________

Are you working full or part time? ____________________________

What days of the week do you work, and how many hours per day do you work?

______________________________________________

**Social Support**

What is your relationship status?

- □ 1 = Single (skip the next question)
- □ 2 = Married
- □ 3 = Long term relationship
- □ 4 = Divorced
- □ 5 = Other ________________________________

Do you live with your partner/significant other? □ 1 = Yes □ 2 = No

Does your partner have a history of substance use disorder? □ 1 = Yes □ 2 = No

Is your partner/significant other currently in treatment? □ 1 = Yes □ 2 = No
How satisfied are you with the support you get from your partner/significant other?
☐ 1 = Very satisfied
☐ 2 = Satisfied
☐ 3 = Fairly satisfied
☐ 4 = Not satisfied
☐ 9 = N/A

Family History

Do any other family members have a history of substance use disorder?
☐ 1 = Yes   ☐ 2 = No

Transportation

How do you get around?
☐ 1 = I drive ➔ Do you have your own car? ☐ 1 = Yes    ☐ 2 = No
☐ 2 = Public Transportation
☐ 3 = Walk
☐ 4 = I get a ride from a family/friend
☐ 5 = Other ______________________

Do you have a valid form of government issued identification?   ☐ 1 = Yes    ☐ 2 = No

How would you get to the OBAT program if you needed to get here?
☐ 1 = I would drive
☐ 2 = Public Transportation
☐ 3 = I would walk
☐ 4 = I get a ride from a family/friend
☐ 5 = Other ______________________

Housing

Have you spent one or more weeks on the street or in a shelter in the last three months?
☐ 1 = Yes    ☐ 2 = No

What type of place are you living in now?
☐ 1 = In a house or apartment you own or rent
☐ 3 = In a house or apartment owned or rented by family or friends
☐ 4 = Hotel
☐ 5 = Alcohol or substance use treatment program
☐ 6 = Shelter
☐ 7 = Street or car
☐ 8 = Other (specify other): ______________________________
☐ 9 = Don’t know
Who do you live with at this time?
☐ 1 = I live alone.
☐ 2 = I live with my partner/significant other
☐ 3 = I live with family members
☐ 4 = I live with friends
☐ 5 = Other: _________________________

Can you tell me what your goals are for treatment?
_____________________________________________________________________________
APPENDIX 4: NURSING INTAKE

Nursing Summary:

Are you pregnant at this time?
☐ 1 = Yes
☐ 2 = No
☐ 3 = Don’t Know
☐ 4 = Tubal ligation
☐ 5 = Menopause
☐ 6 = History of hysterectomy
☐ 7 = Other

If no, are you on birth control?  ☐ 1 = Yes  ☐ 2 = No

If yes, which method of contraception are you currently utilizing? (check all that apply)
☐ Male condoms
☐ Oral contraceptives
☐ Injection (e.g. Depo-Provera)
☐ Hormonal implant
☐ Intrauterine device/contraception (IUD or IUC)
☐ Vaginal ring
☐ Patch
☐ Female barrier method (e.g. diaphragm, female condom)
☐ Rhythm/Fertility Awareness Methods/Withdrawal
☐ Other:

Substance Use History

What substances you currently using at this time?
Includes age of first use, last use, route, frequency, and quantity.
☐ 1 = heroin
☐ 2 = fentanyl
☐ 3 = buprenorphine/naloxone
☐ 4 = methadone
☐ 5 = oxycodone product
☐ 6 = other opioid
☐ 7 = cocaine
☐ 8 = benzodiazepines
☐ 9 = Alcohol
☐ 10 = Amphetamines
☐ 11 = Nicotine
☐ 12 = Other
☐ 13 = Nothing
Do you have any history of any other addictive behaviors such as?
☐ 1 = Gambling  
☐ 2 = Sex  
☐ 3 = Shopping  
☐ 4 = Eating disorder (over eating, bulimia, anorexia)  
☐ 5 = Other:  
☐ 6 = No

Comments:___________________________________________________

**Prior Substance Use Disorder Treatment History**

**Methadone:**

Have you ever been on Methadone Maintenance?  ☐ 1= Yes  ☐ 2 = No

When and where were you on Methadone Maintenance?________________________

What was your dose? ______

Why did you stop Methadone treatment?________________________________

Are you currently on Methadone Maintenance?  ☐ 1= Yes  ☐ 2 = No

What is your dose? ______

Where are you receiving services for your Methadone treatment?_______________

What is the name of your counselor at your Methadone clinic?________________

**Buprenorphine/Naloxone:**

Have you ever been prescribed buprenorphine/naloxone before?
☐ 1= Yes  ☐ 2 = No

If yes, when were you on buprenorphine/naloxone?________________________

What was your dose? ________________

Why did you stop taking buprenorphine/naloxone? ________________

Are you still on buprenorphine/naloxone?  ☐ 1= Yes  ☐ 2 = No

If yes, where/who is prescribing your buprenorphine/naloxone?_______________
What was your dose? ________________

When did you receive your most recent prescription? ________________

Naltrexone:

Have you ever been prescribed naltrexone before?
□ 1= Yes  □ 2 = No

If yes, when were you on naltrexone? ________________________________

Have you ever received an extended-release naltrexone injection? ________________

If yes, when was your most recent injection? ________________________________

Why did you stop naltrexone treatment? ________________________________

Mental Health History

Have you ever been diagnosed with any of the following mental health conditions?
□ 1 = Depression □ 5 = Obsessive Compulsive Disorder (OCD)
□ 2 = Anxiety □ 6 = Post Traumatic Stress Disorder (PTSD)
□ 3 = Bipolar □ 7 = Attention Deficit Disorder
□ 4 = Schizophrenia □ 8 = Anxiety □ 9 = Other: ____________________

Are you currently taking any medication for this/these problem(s)?
□ 1= Yes □ 2 = No

If yes, what medications are you taking? ________________________________

Health status

Have you ever been diagnosed with any medical conditions? Mark all that apply.
□ 1=Diabetes (specify type): ________________________________
□ 2=Heart disease (specify type): ________________________________
□ 3=Cancer (specify type): ________________________________
□ 4=Asthma
□ 5=Tuberculosis (TB)
□ 6=Endocarditis
□ 7=Skin infection
□ 8= HIV ➔ If yes, are you currently in care? □ 1= Yes □ 2 = No
□ 9= Hepatitis A
□ 10= Hepatitis B ➔ If yes, have you been treated? □ 1= Yes □ 2 = No
□ 11= Hepatitis C ➔ If yes, have you been treated? □ 1= Yes □ 2 = No
□ 12= Seizure disorder ➔ Are you on medications? □ 1= Yes □ 2 = No
□ 13= Head Trauma/Brain Injury
☐ 14= Pancreatic Problems
☐ 15= Other (specify type): ________________________________
☐ 16= None

PMH History:
______________________________________________________________________________

Current Medications:
______________________________________________________________________________

Allergies: _________________________________________________________________

Have you been tested for HIV?  ☐ 1= Yes  ☐ 2 = No
   If yes, did you go back for the results?  ☐ 1= Yes  ☐ 2 = No
   If yes, when was the last time you were tested? ____________

Have you been tested for Hepatitis C?  ☐ 1= Yes  ☐ 2 = No
   If yes, did you go back for the results?  ☐ 1= Yes  ☐ 2 = No
   If yes, when was the last time you were tested? ____________

Do you have any pending surgeries? ☐ 1 = Yes  ☐ 2 = No

Pain

Do you have chronic pain?  ☐ 1=Yes  ☐ 2=No

If yes, please explain:
______________________________________________________________________________

Please rate your pain, on a scale from 0 – 10, without any pain medications (prescribed or bought on the street)

   ____0____1____2____3____4____5____6____7____8____9____10____

Can you tell me what your goals are for treatment?
______________________________________________________________________________

Check all appropriate boxes:

☐ OBAT program reviewed with patient including requirements to keep medical and OBAT appointments, urine toxic screens and possible random call backs with medication counts. He /
She is aware of his/her responsibility for their buprenorphine/naloxone medication. Informed to keep medication in a safe undisclosed place, out of reach of children and visitors. Informed to keep medication in a locked storage unit.

☐ OBAT consent and treatment agreement read to and reviewed with the patient. Patient voluntarily signed and dated consent. A copy was given to the patient and the original was placed in the chart. Opportunity for questions provided.

☐ Discussed buprenorphine/naloxone - reviewed medication, potential side effects including elevations in transaminases, potential lethal interaction with benzodiazepine, other sedating medications and ETOH, safe administration and storage. Written information also provided to pt. Patient verbalizes understanding of information provided and wishes to schedule induction phase time and date.

☐ Discussed naltrexone - reviewed potential side effects and adverse reactions including injection site reactions, allergy, pneumonia, increase transaminases, depression, dizziness, opioid blocking effects, and decreased opioid tolerance. Patients need to be opioid free for an extended period of time prior to administration to prevent precipitated or spontaneous withdrawal. Patients who are naltrexone naive will begin with the tablet form of the medication to assess for side effects or adverse reactions. Written info provided to patient. Patient verbalized understanding and wishes to initiate naltrexone treatment.

☐ Contact numbers of medical providers and wallet size buprenorphine/naloxone information given to pt. Patient instructed to give these cards to family members or friends in case patient is ever hospitalized.

☐ Contact numbers of medical providers and wallet size naltrexone information given to pt. Patient instructed to give these cards to family members or friends in case patient is ever hospitalized. Patient also provided with naltrexone medical identification: bracelet and/or dog tag.

☐ Patient has been informed that both buprenorphine/naloxone and naltrexone are Category C medications. Breastfeeding is a currently contraindicated during naltrexone treatment.

☐ Labs sent if indicated may include complete blood count (CBC), Hepatitis A, B, and C serologies, and comprehensive metabolic panel. Required testing includes human chorionic gonadotropin (hCG), urine toxicology screen, and HIV testing strongly recommended.

☐ Overdose education provided. Pt has been trained and has access to use a naloxone rescue kit, if not a prescription is sent to the pharmacy
APPENDIX 5: INDUCTION NOTE

☐ Patient Presents for First Induction

Evaluated using COW scale?  ☐ Yes ☐ No

Scored _________ on COW Scale First Assessment

Patient self-administered ______ mg sl as prescribed

☐ Assessed and instructed patient in proper administration
☐ Patient observed to tolerate medication

Summary 1:
COW Scale First Assessment

Resting Pulse Rate
☐ 0 = pulse rate 80 or below
☐ 1 = pulse rate 80-100
☐ 2 = pulse rate 101–120
☐ 3 = pulse rate greater than 120

Sweating
☐ 0 = no report of chills or flushing
☐ 1 = subjective report of chills or flushing
☐ 2 = flushed or observable moistness on face
☐ 3 = beads of sweat on brow or face
☐ 4 = sweat streaming off face

Restlessness during Assessment
☐ 0 = able to sit still
☐ 1 = reports difficulty sitting still, but is able to do so
☐ 3 = frequent shifting or extraneous movements of legs/arms
☐ 5 = unable to sit still for more than a few seconds

Pupil Size
☐ 0 = pupils pinned or normal size for room light
☐ 1 = pupils possible larger than normal for room light
☐ 2 = pupils moderately dilated
☐ 5 = pupils so dilated that only the rim of the iris is visible

Bone or Joint Aches
☐ 0 = not present
☐ 1 = mild diffuse discomfort
☐ 2 = patient reports severe diffuse aching of joints/muscle
☐ 4 = patient is rubbing joints or muscles and is unable to sit still because of discomfort

**Runny Nose or Tearing**
☐ 0 = not present
☐ 1 = nasal stuffiness or unusually moist eyes
☐ 2 = nose running or tearing
☐ 4 = nose constantly running or tears streaming down cheeks

**GI Upset**
☐ 0 = no GI symptoms
☐ 1 = stomach cramps
☐ 2 = nausea or loose stool
☐ 3 = vomiting or diarrhea
☐ 5 = multiple episodes of diarrhea or vomiting

**Tremor**
☐ 0 = no tremor
☐ 1 = tremor can be felt, but not observed
☐ 2 = slight tremor observable
☐ 4 = gross tremor or muscle twitching

**Yawning**
☐ 0 = no yawning
☐ 1 = yawning once or twice during assessment
☐ 2 = yawning three or more times during assessment
☐ 4 = yawning several times/minute

**Anxiety or Irritability**
☐ 0 = none
☐ 1 = patient reports increasing irritability or anxiousness
☐ 2 = patient obviously irritable/anxious
☐ 4 = patient so irritable or anxious that participation in the assessment is difficult

**Gooseflesh Skin**
☐ 0 = skin is smooth
☐ 3 = piloerection of skin can be felt or hairs standing up on arms
☐ 5 = prominent piloerection

**Total Score ____**

**Score:** 5 – 12 = Mild  
13 – 24 = Moderate  
25 – 36 = Moderately Severe  
More than 36 = Severe Withdrawal
☐ Patient Presents for Second Induction/Re-Evaluation

Evaluated using COWS? ☐ Yes ☐ No

Scored ________ on second COWS assessment

Patient self-administered ______ mg sl as prescribed

☐ Assessed and instructed patient in proper administration
☐ Patient observed to tolerate medication

☐ Patient Presents for Subsequent Induction/Re-Evaluation

Evaluated using COWS? ☐ Yes ☐ No

Scored ________ on second COWS assessment

Patient self-administered ______ mg sl as prescribed

☐ Assessed and instructed patient in proper administration
☐ Patient observed to tolerate medication
APPENDIX 6: NURSING FOLLOW-UP

Buprenorphine/Naloxone Nursing Follow-up Visit:

Visit type:
☐ Scheduled
☐ Call back
☐ Walk-in
☐ Random call back

Reason for visit:

Current dose of buprenorphine/naloxone:
☐ 1 = 2mg  ☐ 4 = 8mg  ☐ 7 = 16mg  ☐ 10 = 28mg
☐ 2 = 4mg  ☐ 5 = 10mg  ☐ 8 = 20mg  ☐ 11 = 32mg
☐ 3 = 6mg  ☐ 6 = 12mg  ☐ 9 = 24mg  ☐ 12 = Other ______

Is patient taking buprenorphine/naloxone as directed?
☐ 1 = Yes  ☐ 2 = No

The patient's dose is:
☐ Stable
☐ Titrating up
☐ Tapering down

How often is patient taking buprenorphine/naloxone?
☐ 1 = single dose  ☐ 2 = divided dose  ☐ 3 = other:

Is patient experiencing?
☐ Cravings
☐ Withdrawal symptoms
☐ Side effects
☐ Other:
☐ Patient denies cravings/withdrawal symptoms

Comments:

Have there been any changes to your medications since your last visit?
☐ 1 = Yes  ☐ 2 = No

If yes, please list: __________________________________________________________

Do you have any active medical issues?  ☐ 1 = Yes  ☐ 2 = No

If yes, please list: __________________________________________________________
PCP Name: ______________________________________________________________

OBAT Provider Name: _________________________________________________

Was the last OBAT provider visit within 4 months? : ___________

When were the patient’s last labs drawn: ___________

Female Patients: Any chance that you are pregnant at this time?
☐ 1 = Yes
☐ 2 = No
☐ 3 = Don’t Know
☐ 4 = Tubal ligation
☐ 5 = Menopause
☐ 6 = History of hysterectomy
☐ 7 = Other

If no, are you on birth control? ☐ 1 = Yes ☐ 2 = No

If yes, which method of birth control are you currently on? (check all that apply)
☐ Male condoms
☐ Oral contraceptives
☐ Injection (e.g. Depo-Provera)
☐ Hormonal implant
☐ Intrauterine device/contraception (IUD or IUC)
☐ Vaginal ring
☐ Patch
☐ Female barrier method (e.g. diaphragm, female condom)
☐ Rhythm/Fertility Awareness Methods/Withdrawal
☐ Other:

Has patient used any substances?
☐ Opioids
☐ Cocaine
☐ THC
☐ ETOH
☐ Benzodiazepines
☐ Amphetamines
☐ Prescribed controlled substance - reason for prescription:
☐ Patient denies all drug use
☐ None
☐ Other:

Comments: _____________________________________________________________
Is patient engaged in counseling? □ 1 = Yes □ 2 = No

Location of counseling: ________________

What is the name of your counselor: ________________

How often is the patient going to counseling?
□ 1 = Once a week
□ 2 = Every other week
□ 3 = Once a month
□ 4 = Every 2-3 months
□ 5 = Other:

Has the patient missed any counseling appointments? □ 1 = Yes □ 2 = No

What is the reason for the missed appointments?

Is the patient seeing a psychiatrist? □ 1 = Yes □ 2 = No

Name of psychiatrist: ________________

How often is the patient seeing a psychiatrist?
□ 1 = Once a week
□ 2 = Every other week
□ 3 = Once a month
□ 4 = Every 2-3 months
□ 5 = Other:

Are you attending peer support meetings? □ 1 = Yes □ 2 = No

If yes, which meetings do you attend (check all that apply)
□ 1 = AA
□ 2 = NA
□ 3 = Smart Recovery
□ 4 = Other: ________________

If yes, how many meetings do you attend each week?
□ 1 = 1-2 week
□ 2 = 3-4 week
□ 3 = 5-6 week
□ 4 = Daily
□ 5 = Other

Are there any changes in your housing status? □ 1 = Yes □ 2 = No

The following portions of the patient's history were reviewed and updated as appropriate:
Medication List
Recent Lab Results
Allergies
Problem List
Other

Recovery education/support conducted during this session? □ 1 = Yes □ 2 = No

Educated/supported the patient in:
□ 1 = Attending meetings
□ 2 = Attending counseling
□ 3 = Addiction behavior
□ 4 = Recovery issues
□ 5 = Relapse prevention
□ 6 = Relationship/family issues
□ 7 = Obtaining a sponsor
□ 8 = Job training
□ 9 = School/vocational training
□ 10 = Other:

Treatment plan reviewed □ 1 = Yes □ 2 = No

Urine toxic screen sent? □ 1 = Yes □ 2 = No

Urine sample sent for confirmatory testing: □ 1 = Yes □ 2 = No

RTC: □ 1 = Scheduled □ 2 = Random call back

Comments:_____________________________________________________________
**Naltrexone Nursing Follow-up Visit:**

Visit type:
- ☐ Scheduled
- ☐ Call back
- ☐ Walk-in
- ☐ Random call back

Patient Receives:
- ☐ Oral naltrexone
- ☐ Extended-release injectable naltrexone

Last injection Date: ______

Last injection location:
- ☐ Right side
- ☐ Left side

Is patient experiencing?
- ☐ Cravings
- ☐ Medication side effects
- ☐ Medication adverse reactions
- ☐ Other:
- ☐ Patient denies cravings/withdrawal symptoms/adverse effects

OBAT Provider Name:_________________________________________

Was the last OBAT provider visit within 4 months? : _________

Female Patients: Any chance that you are pregnant at this time?
- ☐ 1 = Yes
- ☐ 2 = No
- ☐ 3 = Don’t Know
- ☐ 4 = Tubal ligation
- ☐ 5 = Menopause
- ☐ 6 = History of hysterectomy
- ☐ 7 = Other

If no, are you on birth control? ☐ 1= Yes ☐ 2 = No

If yes, which method of birth control are you currently on? (check all that apply)
- ☐ Male condoms
- ☐ Oral contraceptives
- ☐ Shot (e.g. Depo-Provera)
- ☐ Hormonal implant
- ☐ Intrauterine device/contraception (IUD or IUC)
☐ Vaginal ring  
☐ Patch  
☐ Female barrier method (e.g. diaphragm, female condom)  
☐ Rhythm/Fertility Awareness Methods/Withdrawal  
☐ Other:

**Has patient used any substances?**

☐ Opioids  
☐ Cocaine  
☐ THC  
☐ ETOH  
☐ Benzodiazepines  
☐ Amphetamines  
☐ Prescribed controlled substance - reason for prescription:  
☐ Patient denies all drug use  
☐ None  
☐ Other:

**Patient reports the following medical issues:** __________

**Is patient engaged in counseling?**  
☐ 1 = Yes  
☐ 2 = No

**Location of counseling:** ________________

**What is the name of your counselor:** ________________

**How often is the patient going to counseling?**

☐ 1 = Once a week  
☐ 2 = Every other week  
☐ 3 = Once a month  
☐ 4 = Every 2-3 months  
☐ 5 = Other:

**Has the patient missed any counseling appointments?**  
☐ 1 = Yes  
☐ 2 = No

**What is the reason for the missed appointments?** ________________

**Is the patient seeing a psychiatrist?**  
☐ 1 = Yes  
☐ 2 = No

**Name of psychiatrist:** ________________

**How often is the patient seeing a psychiatrist?**

☐ 1 = Once a week  
☐ 2 = Every other week  
☐ 3 = Once a month  
☐ 4 = Every 2-3 months
☐ 5 = Other:

Are you attending peer support meetings?  ☐ 1 = Yes  ☐ 2 = No

If yes, which meetings do you attend (check all that apply)
☐ 1 = AA
☐ 2 = NA
☐ 3 = Smart Recovery
☐ 4 = Other: ___________________________

If yes, how many meetings do you attend each week?
☐ 1 = 1-2 week
☐ 2 = 3-4 week
☐ 3 = 5-6 week
☐ 4 = Daily
☐ 5 = Other

The following portions of the patient’s history were reviewed and updated as appropriate:
☐ Medication List
☐ Recent Lab Results
☐ Allergies
☐ Problem List
☐ Other

Today’s injection was given on the ______:
☐ Right side
☐ Left side

Are there any changes in your housing status?  ☐ 1 = Yes  ☐ 2 = No

Recovery education/support conducted during this session?  ☐ 1 = Yes  ☐ 2 = No

Educated/supported the patient in:
☐ 1 = Attending meetings
☐ 2 = Attending counseling
☐ 3 = Addiction behavior
☐ 4 = Recovery issues
☐ 5 = Relapse prevention
☐ 6 = Relationship/family issues
☐ 7 = Obtaining a sponsor
☐ 8 = Job training
☐ 9 = School/vocational training
☐ 10 = Other:

Treatment plan reviewed  ☐ 1 = Yes  ☐ 2 = No

Urine toxic screen sent?  ☐ 1 = Yes  ☐ 2 = No
Urine sample sent for confirmatory testing: □ 1 = Yes □ 2 = No

RTC: □ 1 = Scheduled □ 2 = Random call back

Comments:_____________________________________________________________
## Intake Checklist

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<td>VISIT WITH OBAT PROVIDER</td>
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<td>CONSENT FOR TREATMENT SIGNED</td>
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<tr>
<td>TREATMENT AGREEMENT SIGNED</td>
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<td>LABS COMPLETED</td>
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<td>HIV TESTING Y / N</td>
<td></td>
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<tr>
<td>UTS OBTAINED</td>
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<tr>
<td>HQN (ALL WOMEN) HCG?</td>
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<tr>
<td>BCP REVIEW (ALL WOMEN &amp; DOCUMENTED IN NOTE)</td>
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<tr>
<td>MEDICATION LIST</td>
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<td>ALLERGIES LIST</td>
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<td>CONSENT FOR COUNSELOR/PSYCHIATRIST</td>
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<td>CONSENT FOR PROBATION/PAROLE OFFICER</td>
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<td>OTHER CONSENT IF NEEDED</td>
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<td>EMERGENCY CONTACT INFO AND CLINIC CONTACT INFO</td>
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<tr>
<td>HCV REFERRAL</td>
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<td>PPD CURRENT</td>
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<td>ORIENTATION TO THE TEAM AND ITS LOCATION.</td>
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<tr>
<td>PROVIDED CONTACT INFORMATION.</td>
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</tbody>
</table>
APPENDIX 8A: TREATMENT CONSENT

Consent for Treatment with Buprenorphine/Naloxone:

Buprenorphine/naloxone is a FDA approved medication for treatment of people with opioid dependence. Qualified providers can treat up to 30 patients for opioid dependence with buprenorphine/naloxone for the first year of practice and then can apply for another waiver to increase to 100 patients, some qualified providers may treat up to 275 patients. Buprenorphine/naloxone can be used for detoxification or for maintenance therapy. Maintenance therapy can continue as long as medically necessary, it is estimated that one will be on buprenorphine/naloxone for at least 6 months.

Buprenorphine/naloxone treatment can result in physical dependence of an opioid. Withdrawal from buprenorphine/naloxone is generally less intense than with heroin or methadone. If buprenorphine/naloxone is suddenly discontinued, some patients have no withdrawal symptoms; others may have symptoms such as muscle aches, stomach cramps, or diarrhea lasting several days. To minimize the possibility of opioid withdrawal, buprenorphine/naloxone should be discontinued gradually over several weeks or more.

If you are dependent on opioid, you should be in as much withdrawal as possible when you take the first dose of buprenorphine/naloxone. If you are not in withdrawal, buprenorphine/naloxone can cause severe opioid withdrawal.

It may take several days to comfortably transition from the opioid that you had been taking to buprenorphine/naloxone. During this time any use of other opioids may cause an increase in symptoms. After becoming stabilized on buprenorphine/naloxone, the use of other opioid will have less effect. Attempts to override the buprenorphine/naloxone by taking more opioids could result in an opioid overdose.

You should not take any other medications without first discussing with your health care provider.

Combining buprenorphine/naloxone with alcohol or other medications may be hazardous. Combining buprenorphine/naloxone with medications such as Klonopin, Valium, Haldol, Librium, Ativan or other sedating medications may result in overdose or death.

The form of buprenorphine that you will be taking (buprenorphine/naloxone) is a combination of buprenorphine with a short acting opioid blocker (Naloxone). If the buprenorphine/naloxone tablet were dissolved and injected by someone taking heroin or another strong opioid (i.e., Morphine), it may cause severe opioid withdrawal.

Buprenorphine/naloxone tablets/film must be held under the tongue until they completely dissolve, buprenorphine/naloxone will not be absorbed from the stomach if it is swallowed.

____________________________________________________________________________
Print Name                                  Sign Name                                  Date

Witness                                        Date
**BOSTON MEDICAL CENTER**

**CONSENT FOR TREATMENT WITH NALTREXONE**

**Oral Naltrexone (Revia) and Extended-Release Injectable Naltrexone**

Naltrexone is a prescription medication that is used to:

- Prevent relapse to opioid use
- Treat alcohol use disorder

You cannot start naltrexone now if you:

- Are currently using opioids
- Are currently having withdrawal from opioid use

It is necessary to stop all drugs/medications that have any opiates/opioids in them 7-10 days before starting naltrexone to avoid getting sick. It is also important that you **NOT** have any opioids (such as: methadone, buprenorphine, heroin, oxycodone, fentanyl, etc.) in your body and **NOT** be currently withdrawing when you begin treatment.

Urine drug screens will be done before each injection to assure abstinence from opioids.

Because extended-release naltrexone is an injection, it cannot be taken out of the body. To make sure you can tolerate the medication, all patients who have never taken this medication must begin with a dose by mouth (tablet form). If you can tolerate the tablet, you can move on to the injection.

A reaction at the site of injection may occur that may be serious. It is important to get medical attention for reactions that get worse or that you are unsure of, including the following:

- Intense pain
- Area feels hard, lumpy
- Swelling, redness and warmth
- Blisters, and/or skin is open

Allergic reactions can happen soon after an injection of naltrexone. Tell your doctor or get immediate medical help if you have any of these symptoms:

- Skin rash
- Chest pain
- Swelling of eyes, mouth, tongue, or face
- Trouble breathing or wheezing
- Dizziness or fainting

Because naltrexone can affect your liver, and blood will be drawn before starting treatment to check the levels and then as needed during treatment to make sure your liver is healthy. If you develop any symptoms during treatment such as:

- Yellowing of the skin or eyes
- Dark urine
- Stomach pain, or loss of appetite
- More tired than normal,
- White stool or diarrhea

You should contact your doctor or be seen by a medical provider and tell them about the medication you are taking.

You may experience depression while on naltrexone. If you develop depression it is important to tell someone and/or alert your medical providers. If you feel like harming yourself or someone else, you should go to your local emergency room or call 911 if you cannot reach your medical providers.
You may develop signs/symptoms of pneumonia on this medication:

- Shortness of breath
- Difficulty breathing
- Wheezing
- Fevers
- Cough that does not go away

If so, please go to your local emergency room or call 911 if you are not physically able to do so.

Dizziness may occur on naltrexone treatment. You should avoid driving, operating heavy or dangerous machinery until you are sure how naltrexone affects you.

Use of large doses of heroin or other opioids (morphine, oxycodone, methadone, codeine, etc.) while on naltrexone could cause serious injury, coma or death.

If you were addicted to opioids before naltrexone, you will be more sensitive to lower doses of opioids and at Risk for an Overdose should you have a relapse.

Relapse to opioids is very dangerous after being on naltrexone. Do not pick up using what you used before starting naltrexone; your body will be more sensitive to opioids. Alert your family, friends, or close contacts that you are on naltrexone and about the risk of an overdose should you have a relapse.

You should carry alert information so others know you are on naltrexone in a medical emergency: medical alert necklace, bracelet and/or emergency card.

For all women of childbearing age: a pregnancy test will be completed before treatment has begun and then before each next injection. If you learn you are pregnant at any time, please alert your medical team.

You will see your treatment team frequently in the beginning and then less frequently as you become more stable. However it is important to be followed closely for support and assessment. During your treatment you should expect the following:

- Urine drug screens at visits
- Clinical check-ins
- Check in: social supports/recovery network
- Provider visits
- Blood work as indicated
- Monthly injections

Naltrexone treatment is only one part of your treatment. It is important that you seek recovery support services along with the medical part of your treatment to assist you in your recovery process.

In an emergency situation if you require pain management with opioid medications it is important that your medical team know that you are on naltrexone. You would require medical management by providers trained in the use of anesthetic drugs and management of potential respiratory effects. Carry emergency contact information with you at all times and have your team contacted if needed to assist in your care.
• Disulfiram (Antabuse) is a medication that is used to help prevent relapse to alcohol.

• The body is not able to process alcohol while taking disulfiram. This includes even very small doses that may be absorbed from perfume, hand sanitizer, food items (dressings, vinegars, marinades, sauces, extracts etc) and alcoholic beverages. It is important to check labels of items that will go in or on your body.

• Disulfiram should NOT be taken if you have consumed alcohol within the past 12 hours.

• An alcohol-disulfiram reaction may include: trouble breathing, throbbing pain in head and neck, nausea, vomiting, sweating, thirst, palpitations, weakness, dizziness, blurred vision and confusion. Severe reactions may involve respiratory failure, heart failure, unconsciousness, seizure and death.

• The larger the dose of the alcohol, the stronger the disulfiram-alcohol effect. The reaction can last from 30 minutes to several hours, or as long as it takes for the alcohol to be metabolized.

• Disulfiram-alcohol reaction may occur for up to 2 weeks after stopping medication.

• This medication can affect your liver. Blood will be drawn before starting treatment, again soon after starting treatment and then as needed to make sure your liver is healthy. Tell your treatment team or seek emergency care if you develop any of these symptoms:
  - Yellowing of the skin or eyes
  - Dark urine
  - White stool or diarrhea
  - Stomach pain, or loss of appetite
  - More tired than normal

• Allergic reactions can happen when taking disulfiram. Alert your treatment team or get immediate medical help if you have any of these symptoms:
  - Skin rash
  - Chest pain
  - Trouble breathing or wheezing
  - Dizziness or fainting
  - Swelling of eyes, mouth, tongue or face

• The most common side-effect of disulfiram is drowsiness, but severe adverse reactions have occurred in some individuals. These include: liver failure, nerve irritation/neuropathy, psychosis, acne, skin rash, impotence, and inflammation of the optic nerve.
- There are some medications that should not be taken with disulfiram (metronidazole, dronabinol, certain cough medicines, others). It is important to let your providers know that you are prescribed disulfiram. Do not change your medications without checking with your provider.

- It is not known if disulfiram is safe during pregnancy or if it can be passed into breast milk. A pregnancy test will be done before treatment has begun. If you learn you are pregnant at any time please alert your medical team. Disulfiram is not recommend while breastfeeding. 

- Store disulfiram at room temperature, in a light-resistant container. Keep all drugs out of the reach of children and pets.

- Relapse to alcohol is very dangerous after being on disulfiram. Alert your family, friends and close contacts that you are on disulfiram and about the risk of a severe reaction should you have a relapse

- Breathalyzers and toxicology screens will be done at each OBAT visit to help assure abstinence from alcohol.

- Disulfiram is only one part of your treatment. It is important that you seek recovery support services along with the medical part of your treatment to assist you in your recovery process

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<thead>
<tr>
<th>Patient Name</th>
<th>Date</th>
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<table>
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<tr>
<th>Provider Name</th>
<th>Date</th>
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APPENDIX 8B: CONSENT FOR RELEASE OF INFORMATION

CONSENT FOR RELEASE OF INFORMATION

I, __________________________________________, BORN ON ______________________
(PATIENT NAME) (PATIENT BIRTH DATE)

AUTHORIZE __________________________________________ to
(CLINIC OR DOCTOR’S NAME)

DISCLOSE TO __________________________________________
(NAME AND LOCATION OF PERSON/ORGANIZATION TO RECEIVE INFORMATION)

THE FOLLOWING INFORMATION: ____________________________________________.

THE PURPOSE OF THIS DISCLOSURE IS: ________________________________________.

THIS AUTHORIZATION EXPIRES ON: _____________, OR

WHENEVER ______________________ IS NO LONGER PROVIDING ME WITH SERVICES.

I understand that my records are protected under the Federal regulations and cannot be disclosed
without my written consent unless otherwise provided for in the regulations. I also understand that I
may revoke this consent at any time except to the extent that action has been taken in reliance on it.

Signature of patient _____________________________ Dated ___________

Signature of witness ______________________________ Dated ____________

ATTENTION RECIPIENT:
Notice Prohibiting Re-disclosure

This information has been disclosed to you from the records protected by Federal confidentiality rules
(42 C.F.R. Part 2). The Federal rules prohibit you from making any further disclosure of this
information unless further disclosure is expressly permitted by the written consent of the person to
whom it pertains or as otherwise permitted by 42 C.F.R. Part 2. A general authorization for the release
of medical or other information is NOT sufficient for this purpose. The Federal rules restrict any use of
this information to criminally investigate or prosecute any alcohol or drug abuse patient.
APPENDIX 8C: APPOINTED PHARMACY CONSENT

APPOINTED PHARMACY CONSENT

(buprenorphine HCl/naloxone HCl dihydrate) sublingual tablet or film
(buprenorphine HCl) sublingual tablet, naltrexone (oral or extended-release injectable)

I ______________________________________________ do hereby: (check all that apply)

□    Patient Name (Print)

□    Authorize ________________________________ at the above address to disclose my treatment for substance use disorder to employees of the pharmacy specified below. Treatment disclosure most often includes, but may not be limited to, discussing my medications with the pharmacist, and faxing/calling in my buprenorphine/naloxone or naltrexone prescriptions directly to the pharmacy.
□    Agree to purchase all buprenorphine/naloxone, and any other medications related to my treatment from the pharmacy specified below.
□    Agree not to use any pharmacy other than the one specified below for the duration of my treatment with the physician specified above, unless specific arrangements have been made with the physician.
□    Agree to make payment arrangements with the pharmacy specified below in advance of treatment, so that my buprenorphine/naloxone prescriptions can be filled and either delivered to the office addressed given above or picked-up by employees of the same.

I understand that I may withdraw this consent at any time, either verbally or in writing except to the extent that action has been taken in reliance on it. This consent will last while I am being treated for opioid dependence by the physician specified above unless I withdraw my consent during treatment. This consent will expire 365 days after I complete my treatment, unless the physician specified above is otherwise notified by me.

I understand that the records to be released may contain information pertaining to psychiatric treatment and/or treatment for alcohol and/or drug dependence. These records may also contain confidential information about communicable diseases including HIV (AIDS) or related illness. I understand that these records are protected by the Code of Federal Regulations Title 42 Part 2 (42 CFR Part 2) which prohibits the recipient of these records from making any further disclosures to third parties without the express written consent of the patient.

I acknowledge that I have been notified of my rights pertaining to the confidentiality of my treatment information/records under 42 CFR Part 2, and I further acknowledge that I understand those rights.

_____________________________________________________________                
Patient Signature                          Date

_____________________________________________________________                
Parent/Guardian Signature       Parent/Guardian Name (Print)       Date

_____________________________________________________________                
Witness Signature                  Witness Name (Print)                   Date
APPOINTED PHARMACY:

NAME____________________________________ PHONE____________________________________

ADDRESS____________________________________________________________________________

CONFIDENTIALITY OF ALCOHOL AND DRUG DEPENDENCE PATIENT RECORDS

THE CONFIDENTIALITY OF ALCOHOL AND DRUG DEPENDENCE PATIENT RECORDS MAINTAINED BY THIS PRACTICE/PROGRAM IS PROTECTED BY FEDERAL LAW AND REGULATIONS. GENERALLY, THE PRACTICE/PROGRAM MAY NOT SAY TO A PERSON OUTSIDE THE PRACTICE/PROGRAM THAT A PATIENT ATTENDS THE PRACTICE/PROGRAM, OR DISCLOSE ANY INFORMATION IDENTIFYING A PATIENT AS BEING ALCOHOL OR DRUG DEPENDENT UNLESS:

1. THE PATIENT CONSENTS IN WRITING;
2. THE DISCLOSURE IS ALLOWED BY A COURT ORDER, OR
3. THE DISCLOSURE IS MADE TO MEDICAL PERSONNEL IN A MEDICAL EMERGENCY OR TO QUALIFIED PERSONNEL FOR RESEARCH, AUDIT, OR PRACTICE/PROGRAM EVALUATION.

VIOLATION OF THE FEDERAL LAW AND REGULATIONS BY A PRACTICE/PROGRAM IS A CRIME. SUSPECTED VIOLATIONS MAY BE REPORTED TO APPROPRIATE AUTHORITIES IN ACCORDANCE WITH FEDERAL REGULATIONS.

FEDERAL LAW AND REGULATIONS DO NOT PROTECT ANY INFORMATION ABOUT A CRIME COMMITTED BY A PATIENT EITHER AT THE PRACTICE/PROGRAM OR AGAINST ANY PERSON WHO WORKS FOR THE PRACTICE/PROGRAM OR ABOUT ANY THREAT TO COMMIT SUCH A CRIME.

FEDERAL LAWS AND REGULATIONS DO NOT PROTECT ANY INFORMATION ABOUT SUSPECTED CHILD ABUSE OR NEGLECT FROM BEING REPORTED UNDER STATE LAW TO APPROPRIATE STATE OR LOCAL AUTHORITIES.
PROGRAMA PARA EL TRATAMIENTO CONTRA ADICCIÓN EN EL CONSULTORIO MÉDICO (OBAT)

Consentimiento para el tratamiento con Buprenorfina en el Boston Medical Center.

Buprenorfina es un medicamento aprobado por la Administración de Drogas y Alimentos (FDA, por sus siglas en inglés) para el tratamiento de personas con adicción a los opioides. Médicos calificados pueden tratar con Buprenorfina hasta 30 pacientes con dependencia a los opioides. La Buprenorfina puede ser utilizada para la desintoxicación o para una terapia de mantenimiento. Esta terapia puede continuar mientras sea clínicamente necesaria, se estima que se estará tomando Buprenorfina, al menos, durante seis (6) meses.

El tratamiento con Buprenorfina puede resultar en una dependencia física a un opioide. La supresión del Buprenorfina, generalmente, es menos intensa que con heroína o metadone. Si el Buprenorfina se descontinúa de repente, es posible que algunos pacientes no presenten síntomas de retirada (“withdrawal”); otros pueden manifestar síntomas como dolores musculares, dolores estomacales, o diarrea durante varios días. Para minimizar la posibilidad de síntomas de retirada de opioides, la Buprenorfina deberá descontinuarse gradualmente durante varias semanas o más.

Si usted es adicto a los opioides, cuando tome la primera dosis de Buprenorfina, deberá estar desintoxicado lo más posible; si no lo está, el Buprenorfina puede causarle consecuencias graves al suprimir el opioide.

Le tomará varios días para acostumbrarse a la transición del opioide tomado y al uso del Buprenorfina. El uso de cualquier otro opioide durante este tiempo, podrá aumentar los síntomas. Una vez estabilizado con la Buprenorfina, el uso de otro opioide tendrá menos efecto. Intentos de hacer caso omiso al Buprenorfina y tomar más opioides pueden resultar en una sobredosis de opioides.

**No debe tomar ningún otro medicamento sin antes consultarlo con su médico.**

Combinar buprenorfina/naloxone con alcohol y otros medicamentos puede ser dañino. Combinar buprenorfina/naloxone con medicinas como Klonopin, Valium, Haldol, Libium, Ativan u otros medicamentos sedantes puede resultar en sobredosis o muerte.

La composición del Buprenorfina (Suboxone) que tomará es una combinación de Buprenorfina con un bloqueador del opioide de rápida acción (Naloxone). Si la tableta de Suboxone estuviera disuelta e inyectada por alguna persona que estuviera inyectándose heroína o cualquier otro opioide fuerte (i.e. Morfina), causaría grave retirada de opioide (grave “withdrawal”).

Las tabletas de Buprenorfina tienen que colocarse bajo la lengua hasta que estén completamente disueltas, el estómago no absorberá las tabletas si se traga la Buprenorfina.

____________________  ______________________  __________
Nombre en letra de molde  Firma  Fecha

__________________________  __________
Testigo  Fecha
APPENDIX 9A: OBAT AGREEMENT

OBAT AGREEMENT

As a patient in the OBAT program, I freely and voluntarily agree to accept this treatment contract, as follows.

I agree to keep all my scheduled appointments with my provider and nurse, and to conduct myself in a courteous manner in the clinic. It is my responsibility to call the clinic if I will be late/early or need to reschedule my appointment.

I agree not to arrive at the clinic intoxicated or under the influence of substances. If I do my treatment plan will be adjusted accordingly.

I agree not to sell, share or give any of my medication to another person. I understand that such mishandling of my medication is a serious violation of this agreement and may result in referral to a higher level of care or discharge.

I agree not to conduct any illegal, threatening, or disruptive activities in the clinic or on the hospital campus, this is grounds for discharge.

I agree not to tamper with urine screens and if I do so, this may be grounds to referral to a more intensive treatment program. I understand that it is best to be honest with my treatment team if I am struggling and understand the team is here to assist me in my treatment.

I agree that my prescriptions can be given to me only at my regularly scheduled times. Missed appointments may result in my not being able to get medication until the next scheduled visit.

I agree that the medication I receive is my responsibility and that I will keep it in a safe and secure place. I agree that lost medication may not be replaced regardless of the reasons for such a loss. My medication should never be kept in public places, and should be out of the reach and site of children at all times. My medication should be kept in a labeled container that displays a prescription label.

I agree that if I obtain medication from any doctors, pharmacies, or other sources that I will inform my physician and/or OBAT nurse immediately.

I understand that mixing buprenorphine with other substances, especially those which can cause sedation such as benzodiazepines or alcohol can be dangerous. I understand that a number of deaths have been reported among persons mixing buprenorphine with sedating substances.

I agree to take my medication as the provider has instructed and not to alter the way I take my medication without first consulting my provider or nurse.

I agree to random call back visits that include toxicology screens and medication counts. I understand that I need to have a working telephone contact. When called for random call backs, I need to respond within 24 hours by telephone.
I agree not to consume poppy seeds while in this treatment program. Poppy seed consumption will not be accepted as a valid reason for a positive opioid screen.

I understand that if I misuse other substances or medications, this issue will be addressed through changes in my treatment plan to assist me. If I continue to struggle with ongoing substance use this could be grounds for transfer to other more intense treatment options.

Positive urine screens for opioids will be evaluated by the treatment team, ongoing positives or missed urines will prompt a team meeting to discuss a potential change in treatment plan including a referral to more intense treatment.

**Urine screens that are negative for buprenorphine will be evaluated by the team and toxicology, and are grounds for intensification of my treatment plan, transfer to another level of care, or discharge.**

OBAT providers will access the Prescription Drug Monitoring Program to review medication profiles on all patients. If patients are found to be accessing prescriptions from other providers, this finding will be reviewed by the OBAT team. If it is determined that the medications obtained by any other providers are in violation of the treatment agreement, the OBAT Team will evaluate the situation, address it with me, and adjust my treatment plan.

I understand that the Office Based Addiction Treatment Program does not have a chain of custody over the urines, the purpose of these tests are for my treatment in OBAT only. If patients have legal or program requirements that require observed urine toxicology testing, this should be done independent of your treatment in OBAT.

If I am female and of child bearing age it is recommended that I utilize contraceptives while on treatment. If I become pregnant while on buprenorphine/naloxone I will alert my health provider immediately so they can assist me in the proper steps and treatment to keep me and my unborn baby safe. This does not mean I will be discharged from treatment, however it may require a change to the “Subutex” tablet which only has buprenorphine.

**Using a new medicine can cause you to react in a number of ways. It is recommended that you do not drive when you first start taking medication until you know how that medication affects you.**

If at any time I am discharged from this program I may be reconsidered at a future time.

I understand that medication alone is not sufficient treatment for my disease, and I agree to participate in the recovery services, as provided, to assist me in my treatment.

I understand that my medical records will be kept in an electronic medical record. These notes will be visible to any healthcare professional involved in my care at this institution. The healthcare providers will only access your medical record if they are involved in your care.

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APPENDIX 9B: SPANISH OBAT AGREEMENT (CONTRATO DE TRATAMIENTO CON BUPRENORFINA)

CONTRATO DE TRATAMIENTO CON OBAT

Como paciente del OBAT, yo libre y voluntariamente acepto este acuerdo para tratamiento, como sigue.

Acepto: asistir, y ser puntual, a todas mis consultas fijadas con mi médico y la enfermera, y ser cortés en la clínica. Es mi responsabilidad llamar a la clínica si llegaré tarde/temprano o si necesito cambiar mi cita.

Acepto: no llegar intoxicado a la clínica o bajo la influencia de narcóticos. En caso contrario, no seré recibido por el médico, ni me será recetado ningún medicamento hasta la próxima cita fijada.

Acepto: no vender, compartir ni dar cualquiera de mis medicamentos a otra persona. Comprendo que la mala administración de mis medicamentos presenta una seria violación al presente contrato, lo cual resultará en referirme a programa de tratamiento más controlado o la terminación del tratamiento sin derecho a apelación.

Acepto: no distribuir, robar, ni realizar ninguna otra actividad ilegal o prejudicial en la clínica y en el hospital o seré dado de alta de inmediato.

Acuerdo: no falsificar los exámenes de orina; en caso contrario, esto será motivo para descontinuar inmediatamente este tratamiento y referirme a un programa de tratamiento más exhaustivo/controlado. Entiendo que es mejor ser honesto con mi equipo de tratamiento y si estoy luchando, entiendo que el equipo está disponible para ayudarme en mi tratamiento.

Acepto: que mis recetas médicos podrán ser entregados, únicamente, en mis horarios regularmente fijados. La falta a las consultas puede resultar en la imposibilidad de obtener medicamentos hasta la próxima consulta fijada.

Acepto: que soy responsable por el medicamento que recibo y que deberá guardarlo en un lugar seguro. Acepto, igualmente, que los medicamentos extraviados no podrán ser reemplazados, sea cual sea la causa de dicho extravío debido al hecho que es una sustancia controlada. Mis medicamentos nunca deben ser guardados en lugares públicos y deben ser guardados lejos del alcance de los niños en todo momento. Mi medicamento debe ser guardado en su botella que muestre el sello con la información de la receta.

Acuerdo: que si obtuviere algún medicamento de otros médicos, farmacias u otras fuentes, deberá informar a mi médico o a la enfermera.
Comprendo que mezclar buprenorfina con otros medicamentos, especialmente con benzodiazepinas y otras drogas puede ser peligroso. Entiendo que ha sido reportado un gran número de muertes de personas que mezclaron buprenorfina con benzodiazepinas.

Acuerdo: tomar los medicamentos como me lo ha indicado el médico, y a no alterar la forma como tomo mis medicinas sin primero consultar con mi médico o la enfermera.

Acepto: visitas para realizar exámenes de orina y a conteos de tabletas al azar. Entiendo que necesito tener un contacto telefónico que funcione. Cuando me llamen al azar, necesito responder en o antes de 24 horas ya que no responder es motivo para darme de alta de la clínica OBAT y para un referido a un nivel de tratamiento más intenso. Las llamadas no respondidas serán consideradas igual que haber obtenido un examen de orina positivo.

Estoy de acuerdo con no consumir semillas de amapola mientras esté en este en el programa de tratamiento. Consumir semillas de amapola puede resultar en una prueba de opioides positiva.

Entiendo que si uso otras sustancias ilegales o medicamentos, esta situación va a ser tratado con cambios en el plan de mi tratamiento a fin de ayudarme a enfrentar esta situación. Si continuó luchando por el uso de las drogas, esto será motivo para pasarme a otras opciones de tratamiento más exhaustivos.

Pruebas de opioides positivas serán evaluadas por el equipo de tratamiento, lo que puede resultar en tratamiento más intensivo.

Los análisis de orina que son negativas para la buprenorfina serán evaluados y positivos para toxicología son motivos para el traslado a otro nivel de atención o para ser dado de alta. 

OBAT periodicamente va a acceder a la sistema estatal de monitoreo de recetas (Prescription Drug Monitoring Program, o PDMP, por sus siglas en inglés) para asegurarse que los pacientes no estén recibiendo otra sustancias controladas de otros proveedores. Si se encuentra que los pacientes acceden recetas de otros proveedores, este resultado será revisado por el equipo de OBAT. Se determina que los medicamentos obtenidos por proveedores fuera del equipo de OBAT constituyen una violación al acuerdo de tratamiento, e equipo de OBAT evaluará la situación y podría resultar en ser dado de alta del Programa OBAT.

Entiendo que el OBAT no tiene una cadena de custodia sobre las pruebas de toxicología en orina. El propósito de estas pruebas de toxicología es para mi tratamiento en OBAT solamente. Si los pacientes tienen requisitos legales o de su programa que requieren pruebas de toxicología en orina observadas, estas deben ser hechas independiente de su tratamiento en OBAT.

Si soy del sexo femenino y en edad para tener hijos (edad reproductiva) es muy recomendable que utilice anticonceptivos durante la administración de buprenorfina/naloxone. Tengo que avisar a mi profesional de salud inmediatamente para que así me
pueda ayudar en los pasos adecuados y el tratamiento para mantenerme a mí y a mi bebé sanos.

Si, en cualquier momento, me dan de alta de este tratamiento, se reconsiderará si el procedimiento en el consultorio médico es la mejor opción para mí en el futuro.

Entiendo que las medicinas solas no son suficiente tratamiento para mi enfermedad, y estoy de acuerdo en participar en educación, consejería y programas de prevención de recaídas según provistos para asistirme en mi tratamiento.

**Entiendo que mi historial, tratamiento e informes médicos serán guardados en los bajo un sistema cerrado de archivos electrónicos confidenciales. Cualquier profesional de la salud que esté participando en mi asistencia médica podrá accesar a estas anotaciones.**

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APPENDIX 10A: TREATMENT PROGRAM REQUIREMENTS

Treatment Program Requirements:

- Patients must keep their scheduled appointments with their OBAT provider.
- Refills will occur at the time of your follow up appointment with the OBAT nurse or provider.
- If an emergency or a schedule change creates a conflict with these appointments patients need to contact the OBAT clinic as soon as possible to address the situation and reschedule the appointment.
- If an emergency arises outside or normal office hours that requires immediate attention from OBAT staff, patients should call the OBAT on call number.
- Patients are required to keep the OBAT clinic updated on all phones numbers and ways to be contacted.
- An OBAT Clinic NCM may call patients for random callbacks and patients must respond by phone within 24 hours of a call and be prepared to come in within 48 hours of a call. If a patient does not respond to a call back the treatment plan may need to be reviewed and changed to better meet your clinical needs.
- Ongoing positive urine screens for opioids will prompt a revision of the treatment plan including referral to more structured treatment options.
- Ongoing struggles with other substances will require a restructured treatment plan potentially including referral to a higher level of care.
- The OBAT clinic must have the name and number of the pharmacy that the patient is using. This information will be kept on file.
- If there are any changes in medications or medical issues including: surgery, medications, hospitalizations, or problems with your OBAT prescription please contact the OBAT nurse.
APPENDIX 10B: CLINICAL APPOINTMENT POLICY

Clinical Appointment Policy:

- All patients who participate in Office Based Addiction Treatment program are required to keep all appointments with their primary care providers, OBAT providers, and OBAT nurses. These appointments are critical to the continuation of care.

- If an appointment cannot be kept, it is the patient’s responsibility to reschedule the appointment. This does not include random callbacks, please see policy under random call backs.

- Patients are expected to make an effort to arrive on time for all scheduled appointments. Appointments with providers may need to be rescheduled if patients arrive late.

- Patients are required to see their OBAT provider at least once every 3-4 months and more frequent if needed per provider, or other medical staff.

- If patients do not show up for medical appointments with their OBAT provider and do not call to inform OBAT staff that they are unable to make the appointment, or arrange for rescheduling, the treatment plan will be revised accordingly.

- If patients continually miss OBAT prescriber appointments and they exceed the four-month visit timeframe, then buprenorphine/naloxone prescriptions may be held until the patient is seen for an office visit by an OBAT provider.
APPENDIX 10C. RANDOM CALLBACK POLICY

Random Callback Policy

- To monitor and verify the proper use of the buprenorphine/naloxone, the OBAT nurse may call the patient sporadically to come in to the clinic for a random toxicology test and a medication count.

- The patient must return this call promptly, and must come to the clinic within 24 hours of the initial call with the medicine bottle and all of the remaining buprenorphine/naloxone tablets or films.

- The patient may be asked to do an observed dose in the clinic observed by the OBAT nurse or provider to further assess adherence.

- For this policy to function, the patient must ensure that we have current and accurate contact information.

- It is the patient’s responsibility to tell the OBAT nurse immediately if there are any changes to this contact information.

- If the patient does not return for a random callback monitoring visit the OBAT Team will meet and reassess the treatment plan with adjustments such as: shorter times between office visits, shorter prescriptions, no refills, etc.
APPENDIX 10D: COUNSELING POLICY

Counseling Policy:

Patients of the Office Based Addiction Treatment (OBAT) program are strongly encouraged to engage in counseling and/or similar intensive recovery support programs. If needed, patients should receive assistance with referrals and linkages for counseling and recovery support services from OBAT staff. Patients are encouraged to attend a minimum of twice monthly counseling visits for the first 12 weeks of treatment. Patients should not be discharged from the OBAT Program if they do not comply with this recommendation as these individuals may be at increased risk for relapse. However, patients who do not engage in counseling or outside recovery support services should continue to receive more intensive monitoring from the OBAT team.

- Patients will agree to sign consent to release information so that OBAT program staff can communicate with the patient’s entire care management team, including those providing outside counseling and recovery support.

- Patients are strongly encouraged to go to weekly or twice monthly counseling (or per the recommendations of the counselor).

- Patients will be expected to discuss their engagement in counseling and other outside recovery services with the OBAT team.

- Groups, IOP’s (Intense Outpatient Programs), Residential, and Halfway houses are methods of treatment that are accepted as counseling.

- If an individual’s counselor or other medical provider recommends that the patient seeks psychiatric evaluation then the patient is required to follow through with this and the decided upon plan of treatment.

- Role of counseling:
  - Educate patient at the onset and ongoing about the importance of adjunct counseling and recovery support and its role. Reinforce that medication alone rarely addresses all aspects of recovery and building recovery capital will improve their chances of success.
  - Educate patients that at the start of treatment, weekly counseling, in the form of either one-on-one or in a group format, is strongly encouraged. Patients are welcome to participate in counseling specific to buprenorphine/naloxone or naltrexone, as they may find it helpful to discuss their treatment openly with others who are engaged in the same treatment.
• Role of self-help peer-support groups

  • Remind patients that recovery is a process that will take a lot of time and commitment. Attending peer-support groups may not be the right treatment modality for them at the start of treatment but something that they may choose later on. They may also decide that peer-support groups are not helpful and prefer other recovery support options. It is important that the patient is empowered and given options.

  • AA, NA and SMART Recovery are examples of self-help treatment options

  • Encourage patients to attend meetings and to keep going, to try different meetings if one does not feel like it “fits.” Encourage patients not to have high expectations, not to focus on what everyone else is or is not doing, to “take what they need and leave the rest.” Remind patients that it often takes some time to build a connection and establish a sense of belonging.

  • Encourage patients to join a home group, to get involved in the meetings (set up, clean up, make the coffee, etc.).

  • For some patients, getting a sponsor, or forming a healthy relationship with another person in recovery, may be a goal they work toward. Patients often report feeling that making this connection is an important piece in one’s recovery process.

  • Hand out AA, NA, SMART Recovery and other meeting books to patients. Assist patients by highlighting some meetings near their work or home at hours that are convenient for them. Contract with them to try a certain number between now and your next visit.

  • Provide patients with websites for NA, AA, Smart Recovery, Emotional Recovery, Online meetings.
APPENDIX 10E: BEHAVIOR POLICY

Behavior Policy:

As a patient in the Office Based Addiction Treatment (OBAT) program, you have made a voluntary decision to participate in this program. We seek to provide an optimum treatment environment for all patients, therefore, patients are expected to maintain appropriate behaviors such as:

- No illegal activities in the clinic environment, or on hospital grounds.
- No disruptive behavior i.e., loud, aggressive behavior, etc. will be tolerated in the clinic.
- No verbal or physical threats towards anyone including: OBAT staff, clerical, pharmacy, other patients, etc. of any kind will be tolerated.
- No possession of weapons or other harmful objects on clinic property.
APPENDIX 10F: MEDICATION ADMINISTRATION POLICY

Medication Administration Policy: Buprenorphine/Naloxone

All patients who participate in the Office Based Addition Treatment program are required to follow the instructions of the OBAT Staff and your provider regarding your buprenorphine/naloxone prescription.

- Patients must take their buprenorphine/naloxone prescription as directed by the prescribing provider.
- Patients should not take more of their prescription without first discussing this with an OBAT nurse.
- Once stabilized, you will receive a prescription with refills.
- Buprenorphine/naloxone is a controlled substance, therefore prescriptions should be filled on the scheduled fill date.
- Patients need to have an identified pharmacy that is kept on file by the OBAT team, should you change the pharmacy, the OBAT team must be notified. Appropriate release should be signed by the patient and kept on file.
- Refills may be canceled if patients do not return for scheduled visits or when randomly requested.
- Patients should have a safe place to store their medication.
- It is strongly advised that patients do not carry the buprenorphine/naloxone on their person, keep it in a vehicle, or bring to work, etc. as it is a controlled medication. Reports of lost/stolen/destroyed medication require a team consult.
- The OBAT team expects that patients will inform their other providers (therapists, counselors, physicians, etc.) that they are taking buprenorphine/naloxone.
- It is strongly advised that patients carry the emergency identification card on buprenorphine/naloxone on their person, and give this card to a provider should they have the need for medical treatment.
- Patients are also expected to disclose to OBAT staff if they are being seen by other providers (pain management specialists, psychiatrists, counselors, physicians, etc.) and whether they have been prescribed medications by these providers.
APPENDIX 10G: URINE TOXICOLOGY POLICY

Urine Toxicology Policy

- Urine samples are required at each visit.

- All belongings (coats, bags, etc.) are left in the office of the medical assistant or outside the bathroom door. Patients may keep their wallet and cellphone with them.

- No washing of hands until the labeled urine sample is handed to the Medical Assistant in a bio-hazardous bag.

- No flushing of toilet until urine sample is handed to the gloved Medical Assistant.

- Any questionable urine is an automatic repeat the same day.
  - In the event of a questionable sample, the patient will meet with the OBAT NCM to discuss any concerns. This may occur before and/or after a repeat toxicology sample is collected.
  - Please remember that if you are struggling, the OBAT team is here to help.

- Patients will receive a buprenorphine/naloxone prescription, naltrexone prescription or extended-release naltrexone injection after an acceptable toxicology sample is obtained.
APPENDIX 11A: PATIENT HANDOUT: PEDIATRIC EXPOSURE TO BUPRENOPHINE/ NALOXONE

ORDER @ HTTP://MASSCLEARINGHOUSE.EHS.STATE.MA.US/PRODUCT/SA1064KIT.HTML

Protecting Others and Protecting Treatment
KEEPING BUPRENOPHINE* SAFE AND AWAY FROM CHILDREN

REGIONAL CENTER FOR POISON CONTROL AND PREVENTION
Serving Massachusetts and Rhode Island
1-800-222-1222

*Some of the brand names are Suboxone, Subutex, Bunavali, Zubsolv
APPENDIX 11B: PATIENT HANDOUT: OVERDOSE EDUCATION

1 Know the Signs of Overdose. Save a Life.

- Signs of opioid overdose may include:
  - Breathing that is slow or shallow — or no breathing at all
  - Very sleepy and not responding to your voice or touch
  - Blue or grayish skin color, with dark lips and fingernails
  - Snoring or gurgling sounds

- If there are symptoms of an overdose:
  - Tap, shake, and shout at the person to get a response
  - If there is still no response, rub knuckles on the breast bone
  - If no or little response, call 911

Opioids include: heroin, codeine, fentanyl, hydrocodone (i.e. Vicodin), hydromorphone, morphine, oxycodone (i.e. OxyContin, Percocet), etc.

2 Call 9-1-1. An Overdose Is a Medical Emergency.

- An opioid overdose can cause a coma or death within minutes. A medication called naloxone (Narcan) can reverse an overdose and save a life.

- When you call 9-1-1:
  - Give the address
  - Tell them it’s an overdose so they can bring naloxone (Narcan). Or say, “My friend is not breathing.”
  - Stay with the person. The 9-1-1 Good Samaritan law provides protection from arrest and prosecution for drug possession.

- While you wait for the ambulance:
  - Do rescue breathing.
  - Give naloxone (Narcan) if you have it.
  - If you have to leave the person for any amount of time, place the person on their side.

- Tell the ambulance staff anything you can about any alcohol or drugs the person has taken. If you cannot stay, leave a note with the information.

3 Do Rescue Breathing if Breathing Is Slowed or Stopped.

- Make sure nothing is in the mouth.
- Tilt head back, lift chin, pinch nose.
- Breathe in mouth once every 5 seconds.
Get Treatment. There is Hope.

You are not alone. The following resources can help you find substance abuse treatment, prevention services, and information.

Massachusetts Substance Abuse Information and Education Helpline
- Free and confidential information and referrals to public and private treatment programs
- Health insurance may not be required
- Translation available in 140 languages
Toll free 1-800-327-5050
Staffed 7 days a week
TTY: Use MassRelay at 711 or 1-800-720-3480
www.helpline-online.com

Massachusetts Health Promotion Clearinghouse
- Resources on prevention and treatment.
(Toll free) 1-800-952-6637
TTY: Use MassRelay at 711 or 1-800-720-3480
www.mass.gov/maclearinghouse

Massachusetts Overdose Prevention Resources
- Free and confidential training on preventing, recognizing, and responding to overdose is available. Training includes rescue breathing and how to use naloxone (Narcan).
- Naloxone (Narcan) is available at specific locations statewide. It is also available at many pharmacies. Ask your pharmacist.
- To find a naloxone (Narcan) site near you call:
Toll free 1-800-327-5050
TTY: Use MassRelay at 711 or 1-800-720-3480
Help is available in over 140 languages.
www.helpline-online.com

For information about available overdose resources visit
www.mass.gov/dph/overdose
**APPENDIX 12A: CLINICAL TOOLS: COWS SCORE**

**Opioid Withdrawal Record (Induction Form)**
(Adapted from Clinical Opioid Withdrawal Scale)

Patient Name_______________________________________________________ Treatment Start Date_____________

Circle the number/description which best corresponds to your patient’s present symptoms

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Baseline Observation</th>
<th>1st Dose Observation</th>
<th>1st Dose, 2nd Observation (if needed)</th>
<th>2nd dose Observation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Administer 1st Dose mg</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Time given ____am/pm</td>
<td>4</td>
<td>4</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>Resting pulse rate ____beats/min</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Measure after patient is sitting lying for 1 min</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 pulse rate 80 or below</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 pulse rate 81-100</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2 pulse rate 101-120</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4 pulse rate greater than 120</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sweating Over past 30 minutes; not accounted for by room temperature or patient activity</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 no report of chills or flushing</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 subjective report of chills or flushing</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2 flushed or observable moistness on face</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 beads of sweat on brow or face</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4 sweat streaming off face</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Restlessness Observation during assessment</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 able to sit still</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 reports difficulty sitting still, but is able to do so</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 frequent shifting or extraneous movements of legs/arms</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5 unable to sit still for more than a few seconds</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pupil size</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 pupils pinned or normal size for room light</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 pupils possibly larger than normal for room light</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2 pupils moderately dilated</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5 pupils so dilated that only the rim of the iris is visible</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

---

**Boston Medical Center OBAT Clinical Guidelines 2018**

Page 135
<table>
<thead>
<tr>
<th></th>
<th>Baseline Observation</th>
<th>1st Dose Observation</th>
<th>1st Dose, 2nd Observation</th>
<th>2nd Dose</th>
<th>2nd Dose Observation</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>GI upset</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Over last 30 minutes</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 no GI symptoms</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>1 stomach cramps</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>2 nausea or loose stool</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 vomiting or diarrhea</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>5 multiple episodes of diarrhea or vomiting</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td><strong>Anxiety or irritability</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 none</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>1 patient reports increasing irritability or anxiousness</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>2 patient obviously irritable/anxious</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4 patient so irritable/anxious that participation in assessment is difficult</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>104232</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Bone or joint aches</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>If patient was having pain previously, gauge the additional component attributed to opioid withdrawal only</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 not present</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>1 mild diffuse discomfort</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>2 patient reports severe diffuse aching of joints/ muscles</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>4 patient is rubbing joints or muscles and is unable to sit still because of discomfort</td>
<td>4</td>
<td>4</td>
<td>4</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td><strong>Yawning</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Observation during assessment</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 no yawning</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>1 yawning once or twice during assessment</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>2 yawning three or more times during assessment</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>4 yawning several times/minute</td>
<td>4</td>
<td>4</td>
<td>4</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td><strong>Runny nose or tearing</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Not accounted for by cold symptoms or allergies</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 not present</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>1 nasal stuffiness or unusually moist eyes</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2 nose running or tearing</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4 nose constantly running or tears streaming down cheeks</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>104242</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Gooseflesh skin</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 skin is smooth</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>3 skin piloerection can be felt or hairs standing up on arms</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5 prominent piloerection</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td><strong>Total Score</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total score is the sum of all 11 items</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• 5-12 = mild</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• 13-24 = moderate</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• 25-36 = moderately severe</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• &gt;36 = severe withdrawal</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**APPENDIX 12B: CLINICAL TOOLS: MULTIDISCIPLINARY APPROACH TO BUPRENORPHINE/NALOXONE MAINTENANCE**

Boston Medical Center’s Multidisciplinary Approach to Buprenorphine/naloxone Maintenance

- **Contact Initiated or provider referral**
- **Chart reviewed by OBAT clinician for appropriateness, is then either referred to another level of care or screened.**
- **Program Coordinator, OBAT Medical Assistant or OBAT NCM screens the patient**
- **OBAT Nurse Manager reviews Initial screen, conducts chart review. Reviews info from other sources i.e., methadone clinic. Approves move forward or recommends other treatment options.**

- **Program coordinator schedules patient for:**
  - OBAT NCM Intake Visit
  - OBAT Provider appointment
  - Addresses counseling needs
  - Collects pharmacy information

- **OBAT Nurse Intake Visit**
  - Nursing assessment
  - Consent, contract reviewed/signed/copies to patient
  - Education
  - Lab work, UTS
  - Post NCM Visit: reviews labs, urine toxicology. Alert OBAT provider of abnormal labs, contacts patient as needed

- **OBAT Provider visit**
  - Provider assessment: DSM5 Confirmed PE if needed
  - Review labs
  - Sign off on appropriateness

- **OBAT Nurse Care Manager:**
  - Generates prescription which is then signed by OBAT provider.
  - Reviews induction plan with patient.
  - Prescription faxed to pharmacy for patient pick up day of induction

- **OBAT Nurse Intake Visit**
  - Induction in person or home
  - Follow by telephone as needed
  - Return for weekly or more frequent if needed x 4-6 weeks
  - Weekly to EOW, Monthly, Random, PRN

- **OBAT Provider**
  - Patient visits at least once every 4 months
  - PCP visits as needed
  - EMR and face to face communication: notes, labs, UTS, prescription dose, visit frequency, treatment plan, acute issues, etc.
  - Phone consultation and support as needed

- **OBAT Nurse Care Manager:**
  - Maintenance
  - Follow-ups/Callbacks/Phone check in
  - Medical issues: surgery, acute/chronic pain, HCV, medication updates, pregnancy
  - Mental health: need for psychiatric evaluation, medication, counseling, intervention

- **Program Coordinator:**
  - Collaboration with counseling
  - Update contact information, pharmacy information
  - Assist with team meetings
  - Assist with patient interventions
  - Assist with insurance issues
  - Assist in development of new treatment plan

---

*Key*
- Green = Patient
- Red = Program Coordinator
- Blue = Nurse
## Appendix 12C: Clinical Tools: Pharmacotherapy for Opioid Use Disorders

<table>
<thead>
<tr>
<th>Indications</th>
<th>Methadone</th>
<th>Buprenorphine/Naloxone</th>
<th>Buprenorphine</th>
<th>Naltrexone</th>
</tr>
</thead>
<tbody>
<tr>
<td>OUD (DSM diagnosis) and patient meets Federal OTP Standards (42 C.F.R. §8.12)</td>
<td>OUD (DSM diagnosis)</td>
<td>OUD or AUD (DSM diagnosis) with:</td>
<td>Prevention of relapse to opioid dependence/use, following opioid detoxification</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Willingness and stability to receive, store, and administer weekly supply of buprenorphine/naloxone</td>
<td></td>
<td>Treatment for alcohol use disorders</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Willingness and stability to receive monthly injections</td>
<td></td>
</tr>
</tbody>
</table>

### Contraindications

- Hypersensitivity
- Chronic pain requiring opioid management beyond buprenorphine.
- Receiving opioid agonists
- Physiologic opioid dependence
- Failed naloxone challenge or naltrexone challenge test
- Positive urine opioid screen
- Acute Hepatitis or liver failure
- Hypersensitivity
- Advanced psychiatric disease, active suicide ideation
- Breastfeeding - oral naltrexone has shown tumorigenicity in animal studies

### Warnings/Precautions

- Concurrent enrollment in another OTP
- Use caution in patients with respiratory, liver, or renal insufficiency
- Concurrent benzodiazepines or other CNS depressants including opioids and active AUD (potential respiratory depression)
- Use of opioid antagonists
- Pregnancy Category C
- Buprenorphine/naloxone may precipitate withdrawal in patients on full agonist opioids
- Use caution in patients with respiratory, liver, or renal insufficiency
- Concurrent benzodiazepines or other CNS depressants, including opioids and active AUD (potential respiratory depression, overdose)
- Use of opioid antagonists (e.g., parenteral naloxone, oral or parenteral nalmefene, naltrexone)
- Pregnancy Category C
- Active liver disease, cirrhosis
- Moderate to severe renal insufficiency: unknown effects
- Thrombocytopenia or coagulation disorders
- Chronic and/or acute pain must be managed with non-opioids
- Large body habitus
- Vulnerability for fatal opioid overdose in case of relapse to opioids
- Pregnancy Category C

### Baseline Evaluation
- Consider baseline electrocardiogram and physical examination for patients at risk for QT prolongation or arrhythmias
- Toxicology screen
- Liver transaminases
- Urine beta-HCG for females
- Toxicology screen
- Liver transaminase levels <5x upper normal limits
- CrCl (estimated or measured) 50 mL/min or greater
- Ensure patient has adequate muscle mass for injection
- Urine beta-HCG for females
- Toxicology screen

### Dosage and Administration
- **Initial dose:** 15-20 mg single dose, maximum 30 mg
- **Daily dose:** Maximum 40 mg/day on first day
- **Usual dosage range for optimal effects:** 60-120 mg/day
- **Titrate carefully, consider methadone’s delayed cumulative effects**
- **Administer orally in single dose**
- **Individualize dosing regimens**
- **Daily visits at OTP clinic, may receive take-home doses per clinic protocol.**

**Sublingual dosing:**
- **Induction:** Pt to present in mild-moderate withdrawal
- **Induction dose:** 2-4mg initial dose, titrate per prescription instructions and/or until withdrawal symptoms subside.
- **Typical Day 1 dose = 8mg**
- **Day 2-7:** Take total dose equivalent from day 1 upon awakening. Check in with clinical team. May titrate up to 16mg.
- **Stabilization/maintenance:** Target dose = 8-16mg (max 24mg daily) may be taken in single or bid dosing regimen.
- **weekly visits/prescriptions until stable, then biweekly, and eventually monthly or random callback basis**
- **To be administered after negative UTS and/or successful naltrexone/naloxone challenge.**
- **Oral:** 25-50mg by mouth daily
- **ER Injectable:** 380 mg every 28 days by deep intramuscular gluteal injection
- **Alternate injection sites**
- **Weekly visits until stable, then biweekly, may progress to clinic visits every 28 days occurring on the date of patient’s extended-release naltrexone injection.**

### Alternative Dosing Schedules
- **Give in divided daily doses based on peak and trough levels that document rapid metabolism that justifies divided doses**
- **Divided dosing helpful for patients with chronic pain for dual effectiveness and avoidance of narcotic medications**
- **Residential programs may require specific Sig**
- **Consider remaining on oral formulation for patients with coagulation disorders, thrombocytopenia or large body habitus**

### Dosing in Special Populations

<table>
<thead>
<tr>
<th>Special Population</th>
<th>Dosage Adjustment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Renal or hepatic impairment</td>
<td>Reduce dose</td>
</tr>
<tr>
<td>Elderly or debilitated</td>
<td>Reduce dose</td>
</tr>
<tr>
<td>Hepatic impairment</td>
<td>Reduce dose</td>
</tr>
<tr>
<td>For concurrent chronic pain</td>
<td>Divide total daily dose into bid, tid, or qid</td>
</tr>
<tr>
<td>Mild renal insufficiency (CrCl 50-80 mL/min)</td>
<td>No dosage adjustment necessary</td>
</tr>
</tbody>
</table>

### Adverse Effects

<table>
<thead>
<tr>
<th>Major Effects</th>
<th>Common Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Respiratory depression, shock, cardiac arrest, prolongation of QTc interval on electrocardiogram and torsades de pointes ventricular tachycardia</td>
<td>Sedation, nausea, vomiting, edema</td>
</tr>
<tr>
<td>Lightheadedness, dizziness, sedation, nausea, vomiting, constipation, edema</td>
<td></td>
</tr>
<tr>
<td>Less common: Sexual dysfunction</td>
<td></td>
</tr>
<tr>
<td>Hepatitis, hepatic failure, respiratory depression (usually when misused intravenously or if combined with other CNS depressants)</td>
<td>Headache, pain, abdominal pain, insomnia, nausea, vomiting, sweating, constipation</td>
</tr>
<tr>
<td>Common: Headache, pain, abdominal pain, insomnia, nausea, vomiting, sweating, constipation</td>
<td></td>
</tr>
<tr>
<td>Sublingual buprenorphine/naloxone film: Oral hypoesthesia, glossodynia, oral mucosal erythema</td>
<td></td>
</tr>
</tbody>
</table>

### Drug Interactions

| Drugs that reduce serum methadone levels: Ascorbic acid, barbiturates, carbamazepine, ethanol (chronic use), interferon, phenytoin, rifampin, efavirenz, nevirapine, other antiretrovirals with CYP3A4 activity | Metabolized in the liver by Cytochrome P450 3A4 system |
| Drugs that increase serum methadone level: Amitriptyline, atazanavir, atazanavir/ritonavir, cimetidine, delavirdine, diazepam, fluconazole, fluvoxamine, ketoconazole, voriconazole | Drugs that reduce serum buprenorphine level: Ascorbic acid, barbiturates, interferon, carbamazepine, ethanol (chronic use), phenytoin, rifampin, efavirenz, nevirapine, other antiretrovirals with CYP3A4 activity |
| Opioid antagonists may precipitate withdrawal                                  | Drugs that increase serum buprenorphine level: Amitriptyline, atazanavir, atazanavir/ritonavir, cimetidine, delavirdine, diazepam, fluconazole, fluvoxamine, ketoconazole, voriconazole |
|                                                                               | Opioid partial agonist: Buprenorphine/naloxone or buprenorphine may precipitate opioid withdrawal |
|                                                                               | Opioid antagonists may precipitate withdrawal |
|                                                                               | Opioid-containing medications, including over the counter preparations |
|                                                                               | Thoridazine (increased lethargy and somnolence) |

**Monitoring**

| | |
|---|---|---|
| Signs of respiratory and CNS depression | Liver function tests prior to initiation and during therapy as needed | Repeat liver transaminase levels at 6 and 12 months and then every 12 months thereafter |
| Frequent toxicology Screening | Frequent toxicology screening | Increase hepatic monitoring in cases of mild to moderate elevation (1-5x normal limits). |
| Frequent toxicology Screening | |

Abbreviations: OUD: opioid use disorder; UTS: urine toxicology screening; Cmax: maximum concentration; CNS: central nervous system; CrCl: creatinine clearance; DSM: Diagnostic and Statistical Manual of Mental Disorders; HCG: human chorionic gonadotropin; m: meter(s); mg: milligram(s); min: minute(s); mL: milliliter(s)

# Appendix 12D: Clinical Tools: Pharmacotherapy for Alcohol Use Disorder

<table>
<thead>
<tr>
<th>Naltrexone Oral</th>
<th>Naltrexone Injectable</th>
<th>Acamprosate</th>
<th>Disulfiram</th>
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</thead>
<tbody>
<tr>
<td><strong>Indications</strong></td>
<td><strong>Indications</strong></td>
<td><strong>Indications</strong></td>
<td><strong>Indications</strong></td>
</tr>
<tr>
<td>AUD (DSM diagnosis) with:</td>
<td>AUD (DSM diagnosis) with:</td>
<td>AUD (DSM diagnosis) with:</td>
<td>AUD (DSM diagnosis) with:</td>
</tr>
<tr>
<td>- Pretreatment abstinence not required but may improve response</td>
<td>- Pretreatment abstinence not required but may improve response</td>
<td>- Abstinence at treatment initiation</td>
<td>- Abstinence &gt;12 hours and BAL=0</td>
</tr>
<tr>
<td>- Initial engagement in addiction-focused Medical Management and/or other recommended psychosocial intervention</td>
<td>- Willingness to receive monthly injections</td>
<td>- Initial engagement in addiction-focused Medical Management and/or other recommended psychosocial intervention</td>
<td>- Combined cocaine dependence</td>
</tr>
<tr>
<td></td>
<td>- Difficulty adhering to an oral regimen</td>
<td></td>
<td>- Previous response to disulfiram</td>
</tr>
<tr>
<td></td>
<td>- Initial engagement in addiction-focused Medical Management and/or other recommended psychosocial intervention</td>
<td></td>
<td>- Capacity to appreciate risks and benefits and to consent to treatment</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>- Initial engagement in addiction-focused Medical Management and/or other recommended psychosocial intervention</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>- Note: More effective with monitored administration (e.g., in clinic, with spouse, with probation officer)</td>
</tr>
</tbody>
</table>


<table>
<thead>
<tr>
<th>Naltrexone Oral</th>
<th>Naltrexone Injectable</th>
<th>Acamprosate</th>
<th>Disulfiram</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Contraindications</strong></td>
<td><strong>Contraindications</strong></td>
<td><strong>Contraindications</strong></td>
<td><strong>Contraindications</strong></td>
</tr>
<tr>
<td>• Receiving opioid agonists</td>
<td>• Receiving opioid agonists</td>
<td>• Hypersensitivity</td>
<td>• Severe cardiovascular, respiratory, or renal disease</td>
</tr>
<tr>
<td>• Physiologic opioid dependence with use within past 7 days</td>
<td>• Physiologic opioid dependence with use within past 7 days</td>
<td>• Severe renal insufficiency (CrCl ≤30 mL/min)</td>
<td>• Severe hepatic dysfunction (i.e., transaminase levels &gt;3 times upper limit of normal or abnormal bilirubin)</td>
</tr>
<tr>
<td>• Acute opioid withdrawal</td>
<td>• Acute opioid withdrawal</td>
<td>• Hypersensitivity</td>
<td>• Severe psychiatric disorders, especially psychotic and cognitive disorders and suicidal ideation</td>
</tr>
<tr>
<td>• Failed naloxone/naltrexone challenge test</td>
<td>• Failed naloxone/naltrexone challenge test</td>
<td>• Severe renal insufficiency (CrCl ≤30 mL/min)</td>
<td>• Poor impulse control</td>
</tr>
<tr>
<td>• Positive urine opioid screen</td>
<td>• Positive urine opioid screen</td>
<td>• Hypersensitivity</td>
<td>• Metronidazole or ketoconazole therapy which already induce a similar reaction to alcohol</td>
</tr>
<tr>
<td>• Acute Hepatitis or liver failure</td>
<td>• Acute Hepatitis or liver failure</td>
<td>• Inadequate muscle mass or body habitus too large for supplied injection needles</td>
<td>• Hypersensitivity</td>
</tr>
<tr>
<td>• Hypersensitivity</td>
<td></td>
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</tbody>
</table>

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<thead>
<tr>
<th>Naltrexone Oral</th>
<th>Naltrexone Injectable</th>
<th>Acamprosate</th>
<th>Disulfiram</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Warnings/Precautions</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Active liver disease</td>
<td>• Active liver disease</td>
<td>• Monitor for emergence of depression or suicidality</td>
<td>• Alcohol-disulfiram reaction; patients must be vigilant to avoid alcohol in all forms including mouthwash, over the counter medications, etc.</td>
</tr>
<tr>
<td>• Severe renal failure</td>
<td>• Uncertain effects (no data) in moderate to severe renal insufficiency</td>
<td>• Reduce dose in patients with renal insufficiency, including elderly</td>
<td>• Pregnancy Category C</td>
</tr>
<tr>
<td>• Breastfeeding – not advised, proven teratogenicity in animal studies</td>
<td>• Injection site reactions</td>
<td>• Pregnancy Category C</td>
<td></td>
</tr>
<tr>
<td>• Acute/Chronic pain</td>
<td>• Use intramuscular injections with caution in patients with thrombocytopenia or coagulation disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Hx severe depression, acute psychiatric illness</td>
<td>• Acute/Chronic pain</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Pregnancy Category C</td>
<td>• Breastfeeding – not advised</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Hx severe depression, acute psychiatric illness</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Pregnancy Category C</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Liver transaminase levels</td>
<td>• Liver transaminase levels</td>
<td>• CrCl (estimated or measured)</td>
<td>• Liver transaminase levels</td>
</tr>
<tr>
<td>• Bilirubin within normal limits</td>
<td>• Bilirubin within normal limits</td>
<td>• Urine beta-HCG for females</td>
<td>• Physical assessment</td>
</tr>
<tr>
<td>• Urine beta-HCG for females</td>
<td>• CrCl (estimated or measured) 50 mL/min or greater</td>
<td></td>
<td>• Psychiatric assessment</td>
</tr>
<tr>
<td>• Toxicology screen</td>
<td>• Ensure patient has adequate muscle mass for injection</td>
<td></td>
<td>• Electrocardiogram if indicated by history of cardiac disease</td>
</tr>
<tr>
<td></td>
<td>• Urine beta-HCG for females</td>
<td></td>
<td>• Verify abstinence with breath or BAL</td>
</tr>
<tr>
<td></td>
<td>• Toxicology screen</td>
<td></td>
<td>• Urine beta-HCG for females</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Naltrexone Oral</th>
<th>Naltrexone Injectable</th>
<th>Acamprosate</th>
<th>Disulfiram</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Dosage and Administration</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• 50-100 mg orally 1 time daily</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• 380 mg 1 time monthly by deep intramuscular injection</td>
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<tr>
<td>• 666 mg orally 3 times daily, preferably with meals</td>
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<tr>
<td>• 250 mg orally 1 time daily (range, 125-500 mg daily)</td>
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<tr>
<td>• 25 mg 1- or 2-time(s) daily with meals to reduce nausea, especially during the first week</td>
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<tr>
<td>• 100 mg on Monday and Wednesday and 150 mg on Friday</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>• Mild renal insufficiency (CrCl 50-80 mL/min): No dosage adjustment necessary</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Uncertain effects (no data) in moderate to severe renal insufficiency</td>
<td></td>
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</tr>
<tr>
<td>• Reduce dose to 125 mg to reduce side effects</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>• For monitored administration, consider giving 500 mg on Monday, Wednesday, and Friday</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>• Hepatic or renal insufficiency: Use caution</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>• Moderate renal insufficiency (CrCl 30-50 mL/min): 333 mg 3 times daily</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Do not administer to patients with severe renal insufficiency (CrCl ≤30 mL/min)</td>
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<thead>
<tr>
<th>Naltrexone Oral</th>
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<th>Disulfiram</th>
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</thead>
<tbody>
<tr>
<td><strong>Adverse Effects</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Common: Nausea</td>
<td>Major: Eosinophilic pneumonia, depression, suicidality</td>
<td>Major: Suicidality 2.4% (vs. 0.8% on placebo during the first year in clinical trials)</td>
<td>Major: Hepatotoxicity, peripheral neuropathy, psychosis, delirium, severe disulfiram- ethanol reaction</td>
</tr>
<tr>
<td>• Other: Headache, dizziness, nervousness, fatigue, insomnia, vomiting, anxiety, somnolence</td>
<td>Common: Injection-site reactions, injection site tenderness, injection site induration, nausea, headache, asthenia</td>
<td>Common: Diarrhea (16%)</td>
<td>Common: Somnolence, metallic taste, headache</td>
</tr>
<tr>
<td>• Opioid-containing medications, including over the counter preparations</td>
<td>Opioid-containing medications, including over the counter preparations</td>
<td>Naltrexone: 33% increase in Cmax of acamprosate (no dosage adjustment is recommended)</td>
<td>Alcohol containing medications, including over the counter preparations</td>
</tr>
<tr>
<td>• Thioridazine (increased lethargy and somnolence)</td>
<td>Thioridazine (increased lethargy and somnolence)</td>
<td>Antidepressants: Weight gain and weight loss more common than with either medication alone</td>
<td>Drug-drug interactions may occur with phenytoin, warfarin, isoniazid, rifampin, diazepam, chlordiazepoxide, imipramine, desipramine, and oral hypoglycemic agents</td>
</tr>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Naltrexone Oral</th>
<th>Naltrexone Injectable</th>
<th>Acamprosate</th>
<th>Disulfiram</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Monitoring</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Repeat liver transaminase levels at 6 and 12 months and then every 12 months thereafter</td>
<td>• Repeat liver transaminase levels at 6 and 12 months and then every 12 months thereafter</td>
<td>• Monitor serum creatinine/CrCl, particularly in the elderly and in patients with renal insufficiency</td>
<td>• Repeat liver transaminase levels within the first month, then monthly for first 3 months, and periodically thereafter as indicated</td>
</tr>
<tr>
<td>• Discontinue medication and consider alternatives if no detectable benefit after an adequate trial (50 mg daily for 3 months)</td>
<td>• Discontinue if there is no detectable benefit within 3 months</td>
<td>• Maintain therapy if relapse occurs</td>
<td>• Consider discontinuation in event of relapse or when patient is not available for supervision and counseling</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Naltrexone Oral</th>
<th>Naltrexone Injectable</th>
<th>Acamprosate</th>
<th>Disulfiram</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient Education</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Discuss compliance enhancing methods</td>
<td>• Report any concerning injection site reactions</td>
<td>• Report any new or worsening depression or suicidal thinking</td>
<td>• Avoid alcohol in food and beverages, including medications</td>
</tr>
<tr>
<td>• Negotiate commitment from the patient regarding monitored ingestion</td>
<td>• Report any new or worsening depression or suicidal thinking</td>
<td></td>
<td>• Avoid disulfiram if alcohol intoxicated</td>
</tr>
<tr>
<td>• Side effects, if any, tend to occur early in treatment and can typically resolve within 1-2 weeks after dosage adjustment</td>
<td>• May cause allergic pneumonia; contact provider if patient develops signs and symptoms of pneumonia</td>
<td></td>
<td>• May cause sedation; caution operating vehicles and hazardous machinery</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Discuss compliance enhancing methods</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Family members should not administer disulfiram without informing patient</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Provide patients with wallet cards that indicate the use of disulfiram</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• If signs and symptoms of acute Hepatitis occur, discontinue naltrexone and contact provider immediately</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Very large doses of opioids may overcome the effects of naltrexone and lead to serious injury, coma, or death</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Small doses of opioids, such as in analgesic, antidiarrheal, or antitussive drugs, may be blocked by naltrexone and fail to produce a therapeutic effect</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Patients who have previously used opioids may be more sensitive to toxic effects of opioids after discontinuation of naltrexone</td>
</tr>
</tbody>
</table>

Abbreviations: AUD: alcohol use disorder; BAL: blood alcohol level; Cmax: maximum concentration; CNS: central nervous system; CrCl: creatinine clearance; DSM: Diagnostic and Statistical Manual of Mental Disorders; HCG: human chorionic gonadotropin; m: meter(s); mg: milligram(s); min: minute(s); mL: milliliter(s)

APPENDIX 13. BOSTON MEDICAL CENTER’S PERIOPERATIVE MANAGEMENT OF NON-PREGNANT PATIENTS ON MAINTENANCE THERAPY FOR OPIOID DEPENDENCE

Buprenorphine, methadone, and naltrexone are pharmacologic therapies indicated for maintenance treatment of opioid use disorder. The appropriate treatment of acute pain in patients on buprenorphine and methadone maintenance includes continuing the patient’s baseline opioid requirements to avoid increased pain sensitivity associated with opioid withdrawal. Thus, daily opioid maintenance treatment requirements must be met before attempting to achieve analgesia. These patients have also been shown to have increased pain sensitivity and cross-tolerance to opioid analgesics, therefore adequate pain control will often necessitate higher opioid doses at shorter dosing intervals. All patients on buprenorphine and methadone maintenance should be co-managed with their buprenorphine or methadone provider during the pre- and post-procedure period. Addiction medicine is available for consultation to assist with recommendations for opioid use disorder management in the postoperative period.

These guidelines are designed for patients maintained on chronic opioids, buprenorphine, methadone or naltrexone therapy undergoing invasive procedures. There is currently a lack of evidence-based studies to direct the management of patients on buprenorphine, methadone, or naltrexone maintenance in the peri-procedural period. Below are guidelines using expert opinion based on pharmacological principles with the intent to avoid sub therapeutic acute pain management while also preventing opioid withdrawal and disruption of opioid use disorder management.

See Table 1 for recommendations for perioperative management.

References:


Other Related Guidelines or Policies: Methadone and Buprenorphine during Pregnancy, Epidural and Intrathecal Analgesia, Sedation and Pain Control – ICU, Pain Management (Adult), Patient-Controlled Analgesia (PCA) - Adult
**Opioid Dependence**

**Pre-operative Pain Recommendations**

**Post-operative Pain Recommendations**

<table>
<thead>
<tr>
<th>Patient Category</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Chronic Pain on Chronic Opioid Therapy</strong></td>
<td>Continue standing opioid dose the day of surgery. Hold any usual PRN breakthrough opioid doses the day of surgery.</td>
<td>Continue equivalent chronic opioid dose (IV if patient strict NPO) with hold parameters for sedation. For acute postoperative pain, utilize multimodal pain management with non-opioid medications (NSAID, acetaminophen, epidural/spinal analgesia, nerve blocks) as indicated. If opioids are required for breakthrough pain, patients with history of chronic opioid use may require higher than usual doses due to cross tolerance. PCA’s may be considered if pain is not adequately captured. This may be utilized with or without a basal component.</td>
</tr>
<tr>
<td><strong>Inclusion:</strong> Patient on chronic opioids &gt; 2 weeks or with other signs of physical dependence. Does not include patients taking occasional or prn opioids for breakthrough pain.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Methadone Maintenance Therapy</strong></td>
<td>Confirm methadone dose with patient’s methadone maintenance treatment program (MMTP). Continue usual dose of methadone the day of surgery. The patient may need to arrange home doses of methadone (“medical take home doses”) with his or her MMTP if they are unable to go to the MMTP on the day of surgery. If this is not possible, the patient should receive his or her usual confirmed methadone dose in the pre-operative area.</td>
<td>Continue usual daily methadone dose. If the patient is strict NPO, they should receive 50%-75% of their usual methadone dose given IV, divided into 2-4 doses/day (e.g. if usual dose is 60 mg PO daily, appropriate IV doses would be approximately 15 mg IV BID or 10 mg IV TID). For acute postoperative pain, utilize multimodal pain management with non-opioid medications (NSAID, acetaminophen, epidural/spinal analgesia, nerve blocks) as indicated. If opioids are required for breakthrough pain, patients with history of opioid use disorder may require higher than usual doses due to cross tolerance and increased pain sensitivity.</td>
</tr>
<tr>
<td>Buprenorphine Maintenance Therapy</td>
<td>PCA’s without basal component may be considered in addition to patient’s methadone if pain is not adequately captured. Remember to discontinue other oral PRN opioids. On discharge, the patient should be given a “last dose letter” addressed to the MMTP and whether any modifications have been made. The discharge case manager and patient may need to arrange for home doses of methadone (“medical take home doses”) with his or her MMTP if he or she is unable to go to the MMTP on the days of after discharge.</td>
<td>Take AM dose of buprenorphine on the day of the procedure. Continue patient’s home dose of buprenorphine post-operatively. Consider splitting patient’s totally daily buprenorphine dose into q8h schedule for better pain coverage. For acute postoperative pain, utilize multimodal pain management with non-opioid medications (NSAID, acetaminophen, epidural/spinal analgesia, nerve blocks) as indicated. If opioids are required for breakthrough pain, patients with history of opioid use disorder may require higher than usual doses due to cross tolerance and increased pain sensitivity. PCA’s without basal component may be considered in addition to patient’s buprenorphine if pain is not adequately captured. Remember to discontinue other oral PRN opioids.</td>
</tr>
<tr>
<td>Naltrexone (oral or depot) Maintenance Therapy</td>
<td>Discontinue oral naltrexone 72 hours before surgery. Discontinue depot naltrexone 1 month prior to elective surgery, if possible.</td>
<td>Utilize multimodal pain management with non-opioid medications (NSAIDs, acetaminophen, epidural/spinal analgesia, nerve blocks) as indicated. If surgery performed emergently or naltrexone was not discontinued prior to surgery, naltrexone should be discontinued postoperatively. If this occurs, higher than usual doses of opioids may be attempted to overcome naltrexone’s opioid antagonist effects. This must be done with close observation for respiratory depression.</td>
</tr>
</tbody>
</table>
APPENDIX 14: RESOURCE LIST

Practice Guidelines for Treating Substance Use Disorders

American Society of Addiction Medicine (ASAM)
*National Practice Guideline for the Use of Medications in the Treatment of Addiction Involving Opioid Use.* Published June 2015.
https://asam.org/resources/guidelines-and-consensus-documents/npg

ASAM

Substance Abuse and Mental Health Services (SAMHSA)
*TIP 63: Medications for Opioid Use Disorders*
This Treatment Improvement Protocol (TIP) reviews use of the three Food and Drug Administration approved medications used to treat opioid use disorder - methadone, naltrexone, and buprenorphine. Other strategies and services needed to support recovery are also addressed.
Pub id: SMA18-5063FULLDOC Publication Date: 2/2018
https://store.samhsa.gov/product/SMA18-5063FULLDOC

SAMHSA
*Technical Assistance Publication Series. TAP 30: Buprenorphine: A Guide for Nurses*
DHHS Pub id: (SMA) 09-4376 Publication Date: 2009

US Department of Veterans Affairs.
*VA/DoD Clinical Practice Guidelines: Management of Substance Use Disorder*
https://www.healthquality.va.gov/guidelines/mh/sud/

Practice Guidelines for Treating SUD in Pregnancy

SAMHSA
*Clinical Guidance for Treating Pregnant and Parenting Women With Opioid Use Disorder and Their Infants.*
This Clinical Guide provides comprehensive, national guidance for management of pregnant and parenting women with opioid use disorder and their infants. The guide helps healthcare professionals and patients determine the most clinically appropriate action for a particular situation and informs individualized treatment decisions.
Pub id: SMA18-5054 Publication Date: 1/2018
https://store.samhsa.gov/product/SMA18-5054
Institute for Health and Recovery, Massachusetts Perinatal Quality Collaborative

*Maternal Opioid Use During Pregnancy toolkit*

This toolkit provides guidance in regards to the medical, psychological and social needs of pregnant women with opioid use disorders. It was developed to help advance clinical interventions by offering screening, treatment engagement and coordinated care throughout the pregnancy and post-delivery.

http://www.healthrecovery.org/maternal-opioid-use/

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**Practice Guidelines for Safe Opioid Prescribing**

**Centers for Disease Control (CDC)**

*Guideline for Prescribing Opioids for Chronic Pain.*

This guideline is intended to improve communication between providers and patients about the risks and benefits of opioid therapy for chronic pain, improve the safety and effectiveness of pain treatment, and reduce the risks associated with long-term opioid therapy.


**Scope of Pain – Safe and Competent Opioid Prescribing Education**

A series of continuing medical and nursing education activities designed to help providers effectively manage patients with chronic pain, and when appropriate, with opioid analgesics.

www.scopeofpain.com/

**My TopCare - Transforming Opioid Prescribing in Primary Care**

Research and services available for prescribers, pharmacists and patients.

T (617) 414-6938  F (617)414-4676  mytopcare.org

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**Organizations for Professional Education and Evidence-Based Addiction Research**

**American Society of Addiction Medicine (ASAM)**

ASAM is a professional medical society that is dedicated to increasing access and improving the quality of addiction treatment, educating physicians and the public, supporting research and prevention, and promoting the appropriate role of providers in the care of patients with addiction.

T (301) 656-3920 | F (301) 656-3815  https://www.asam.org

**International Nurses Society on Addictions (IntNSA)**

Professional for nurses committed to the prevention, intervention, treatment, and management of addictive disorders including alcohol and other drug dependencies, nicotine dependencies, eating disorders, dual and multiple diagnosis, and process addictions such as gambling.

http://www.intnsa.org/

**National Institute on Alcohol Abuse and Alcoholism (NIAAA)**

NIAAA supports, disseminates, and conducts research on the impact of alcohol use on human health and well-being.

https://niaaa.nih.gov/
National Institute on Drug Abuse (NIDA)
NIDA’s mission is to advance science on the causes and consequences of substance misuse and addiction and to apply that knowledge to improve individual and public health.
https://www.drugabuse.gov/

Providers Clinical Support System – Medication Assisted Treatment (PCSS-MAT)
The goal of PCSS-MAT is to make available the most effective medication treatment for addictions to serve patients in a variety of outpatient settings.
T (888) 572-7724 | F (401) 272-0922 pcssmat@aaap.org

Substance Abuse and Mental Health Services Administration (SAMHSA)
SAMHSA is the agency within the U.S. Department of Health and Human Services that leads public health efforts to advance the behavioral health of the nation. SAMHSA's mission is to reduce the impact of substance abuse and mental illness on America's communities.
https://www.samhsa.gov

Treatment Locator

SAMHSA Behavioral Health Treatment Locator - https://findtreatment.samhsa.gov/
NIAAA Alcohol Treatment Navigator: https://alcoholtreatment.niaaa.nih.gov/
HIV, Hepatitis and STD testing locator: https://gettested.cdc.gov/
PrEP locator: https://aidsvu.org/locators/prep-locator/

SAMHSA: Factsheet for “Finding Quality Treatment for Substance Use Disorders”
https://store.samhsa.gov/product/PEP18-TREATMENT-LOC

Hotline/Resources

SAMHSA National Hotline
Also known as, the Treatment Referral Routing Service, this Helpline provides 24-hour free and confidential treatment referral and information about mental health and/or substance use disorders, prevention, and recovery in English and Spanish.
1-800-662-HELP (4357) TTY: 1-800-487-4889
Website: www.samhsa.gov/find-help/national-helpline

Suicide Prevention Lifeline
24-hour, toll-free, confidential suicide prevention hotline available to anyone in suicidal crisis or emotional distress. Your call is routed to the nearest crisis center in the national network of more than 150 crisis centers.
1-800-273-TALK (8255) TTY: 1-800-799-4889
Website: www.suicidepreventionlifeline.org
Veteran's Crisis Line
Connects veterans in crisis (and their families and friends) with qualified, caring Department of Veterans Affairs responders through a confidential, toll-free hotline, online chat, or text.
1-800-273-TALK (8255) TTY: 1-800-799-4889
Website: www.veteranscrisisline.net

CDC-Info
Offers live agents by phone and email to help you find the latest, reliable, and science-based health information on more than 750 health topics.
1-800-CDC-INFO (1-800-232-4636) TTY: 1-888-232-6348
Email: cdcinfo@cdc.gov
Hours: Monday through Friday, 8:00 a.m. to 8:00 p.m. (Eastern Time)

National Association of People With AIDS Hotline
1-240-247-0880
Hours: Monday through Friday, 9:00 a.m. to 5:30 p.m. (Eastern Time)

Hepatitis Resources

CDC
Viral Hepatitis
Information on prevalence, transmission, screening and diagnosis, and treatment guidelines of Viral Hepatitis. Also includes training information, funding opportunities, and legal resources.
https://www.cdc.gov/hepatitis/resources/

National Institute of Diabetes, Digestive Disorders and Kidney Health
Viral Hepatitis
https://www.niddk.nih.gov/health-information/liver-disease/viral-hepatitis

World Health Organization
Hepatitis C
Includes information on prevalence, transmission, prevention, screening and diagnosis, and treatment guidelines of Hepatitis C.
http://www.who.int/mediacentre/factsheets/fs164/en/

HIV Resources

AIDSInfo
Offers information of HIV/AIDS prevention, treatment, research, clinical guidelines and support for persons living with HIV.
https://aidsinfo.nih.gov/

AIDSVU
AIDSVu is an interactive online map illustrating the prevalence of HIV in the United States, alongside social determinants of health. Site also includes treatment locator services.
https://aidsvu.org/
CDC
HIV/AIDS
https://www.cdc.gov/hiv/default.html

HIV.gov
Resource for healthcare providers, individuals with HIV/AIDS, support persons, and the community. Includes information on prevalence, transmission, testing, treatment, living with HIV/AIDS, and the federal response.
https://www.hiv.gov/

National Institute of Health
Center for AIDS Research (CFAR)
https://www.niaid.nih.gov/research/centers-aids-research

Legal Resources

Legal Action Center.  Substance Use: Confidentiality Resources:
http://lac.org/resources/substance-use-resources/confidentiality-resources/

www.ecfr.gov/cgi-bin/text-idx?c=ecfr&tpl=/ecfrbrowse/Title42/42cfr2_main_02.tpl

Harm Reduction Resources

Harm Reduction Coalition:
National advocacy and capacity-building organization that promotes the health and dignity of individuals and communities impacted by substance use. Efforts advance harm reduction policies, practices and programs that address the adverse effects of drug use including overdose, HIV, hepatitis C, addiction, and incarceration.
http://harmreduction.org/

Buprenorphine Prescribing Regulations

SAMHSA

American Society of Addiction Medicine (ASAM)
Summary of the Comprehensive Addiction and Recovery Act (CARA)

SAMHSA’s Center for Substance Abuse Treatment (CSAT)
866-BUP-CSAT (866-287-2728)
www.buprenorphine.samhsa.gov
Waiver Training Courses

PCSS-MAT
Free courses cover medications and treatments of opioid use disorder, and provide the required hours of education needed to obtain the waiver to prescribe buprenorphine for physicians, nurse practitioners, and physician assistants.
https://pcssmat.org/education-training/mat-waiver-training/

ASAM
Free courses cover medications and treatments of opioid use disorder, and provide the required hours of education needed to obtain the waiver to prescribe buprenorphine for physicians, nurse practitioners, and physician assistants.
https://elearning.asam.org/buprenorphine-waiver-course

Clinical Tools, Inc:
Providers may complete an approved training course to achieve the 8/24hrs of education required to obtain a waiver to prescribe buprenorphine. These courses have a registration fee.
https://www.buppractice.com/

Patient Educational Materials:

Harm Reduction Coalition
http://harmreduction.org/our-resources/

National Institute on Alcohol Abuse and Alcoholism

National Institute for Drug Abuse
https://www.drugabuse.gov/publications/orderable

SAMHSA
https://store.samhsa.gov/home

Provider Continuing Education

Harvard Medical School Global Academy, Opioid Use Disorder Education Program.
A series of three 8hr courses comprise the Opioid Use Disorder Education Program (OUDEP), a continuing medical education program intended for nurses, nurse practitioners, physician assistants, physicians, social workers, and other health care providers collaborating to treat patients with substance use disorders.
https://globalacademy.hms.harvard.edu/

- Course One: Understanding Addiction:
  https://globalacademy.hms.harvard.edu/courses/course-v1:HarvardMedGlobalAcademy+OUDEP1+1T2017/about

- Course Two: Identification, Counseling, and Treatment:
  https://globalacademy.hms.harvard.edu/courses/course-v1:HarvardMedGlobalAcademy+OUDEP2+2T2017/about
• Course Three: Collaborative Care Approaches for Management of OUD:
  https://globalacademy.hms.harvard.edu/courses/course-v1:HarvardMedGlobalAcademy+OUDEP3+2T2017/about

Scope of Pain – Safe and Competent Opioid Prescribing Education
A series of continuing medical and nursing education activities designed to help providers
effectively manage patients with chronic pain, when appropriate, with opioid analgesics.
Ongoing live conferences and online trainings, also an “Ask an Expert” online forum.
www.scopeofpain.com/

Extension for Community Healthcare Outcomes (ECHO)
Project ECHO is a collaborative model of medical education and care management that
empowers clinicians everywhere to provide better care to more people, right where they live.
ECHO programs utilize teleconferencing technology to partner specialist mentors with front-line
clinicians to assist with ongoing clinical education.
https://echo.unm.edu/

Providers Clinical Support System
PCSS is funded by SAMHSA and was created in response to the opioid overdose epidemic to
train primary care providers in the evidence-based prevention and treatment of opioid use
disorders and treatment of chronic pain. PCSS is made up of a coalition, led by American
Academy of Addiction Psychiatry, of major healthcare organizations all dedicated to addressing
this healthcare crisis. Through a variety of trainings and a clinical coaching program, PCSS’s
mission is to increase healthcare providers’ knowledge and skills in the prevention,
identification, and treatment of substance use disorders with a focus on opioid use disorders.
https://pcssnow.org/
APPENDIX 15: LIST OF ACRONYMS

BMC: Boston Medical Center
BSAS: Bureau of Substance Abuse Services
CFR-42: Code of Federal Regulations, Title 42
CNS: Central Nervous System
COWS: Clinical Opioid Withdrawal Scale
CSAT: SAMHSA’s Center for Substance Abuse Treatment
CSS: Clinical Stabilization Services (short-term inpatient stabilization)
DEA: US Drug Enforcement Agency
DCF: Department of Children and Families
DSM: Diagnostic and Statistical Manual of Mental Disorders
ETOH: Alcohol
FDA: Food and Drug Administration
GC/MS: Gas Chromatography/Mass Spectrometry
HCG: Human Chorionic Gonadotropin
HIPAA: Health Insurance Portability and Accountability Act
IOP: Intensive Outpatient Program (counseling)
LFT: Liver Function Test
NAS: Neonatal Abstinence Syndrome
NCM: Nurse Care Manager
NSAID: Non-steroidal Anti-inflammatory Drug
NSDUH: National Survey on Drug Use and Health
OBAT: Office Based Addiction Treatment
OUD: Opioid Use Disorder
OTP: Outpatient Treatment Program (daily medication administration treatment)
PCA: Patient Controlled Analgesia
PDMP: Prescription Drug Monitoring Program
STATE-OBAT: State Technical Assistance and Treatment Expansion of Office Based Addiction Treatment with buprenorphine and naltrexone formulation
TSS: Transitional Stabilization Services (inpatient “holding” facility)
UTS: Urine Toxicology Screening
APPENDIX 16: REFERENCES


