Volume C, Module 2 Leader’s Guide
Opioids: Basics of Addiction; Treatment with Agonists, Partial Agonists, and Antagonists
Module 2: Training goals

To describe the:
- Key components of opiate addiction and its medical / psychiatric consequences
- Benefits and limitations of methadone as a pharmacotherapy for opiate dependence
- Benefits and limitations of buprenorphine as a pharmacotherapy for opiate dependence
- Benefits and limitations of narcotic antagonists for overdose (naloxone) and relapse prevention (naltrexone) for opiate dependence
Module 2: Workshops

Workshop 1: Opiates: What they are, problems associated with their use, and medical treatment implications

Workshop 2: Opiate addiction treatment with methadone

Workshop 3: Opiate addiction treatment with buprenorphine

Workshop 4: Opiate Antagonist Treatment: Naloxone for overdose, Naltrexone for relapse prevention
Icebreaker: Opiate medication in my country

Does your country use opiate medications, and if so, what type of medication?

What are the main problems in your country regarding the use of these medications?

15 minutes
Workshop 1: Opiates

What they are, problems associated with their use, and medical treatment implications
Pre-assessment

Please respond to the pre-assessment questions in your workbook.

(Your responses are strictly confidential.)

10 minutes
Training objectives

At the end of this training you will understand the:

1. Epidemiology of opiate addiction worldwide and its relationship to infectious diseases
2. Basic neurobiology of opiate addiction
3. Medical / psychiatric co-morbidities and treatment strategies for these disorders used with opiate addicts
4. Key issues in engaging opiate addicts into treatment with low threshold approaches
Opioids Definition

Opioids are natural derivatives of opium or synthetic psychoactive substances that have effects similar to morphine or are capable of converting into a drug having such effects.

(SAMHSA, TIP 43)

Notes:
(source: U.S. Department of Health and Human Services. Substance Abuse and Mental Health Services Administration, Center for Substance Abuse Treatment. Medication-Assisted Treatment for Opioid Addiction in Opioid Treatment Programs: TIP 43. Available at www.samhsa.gov)
Global abuse of opiates

Overview:

- Sixteen million (0.4%) of world’s population aged 15-64 abuse opiates
- Heroin abusers make up about 71% of opiate abusers
- Opiates account for 2/3 of all treatment demands in Asia and 60% of treatment demand in Europe

Regional Breakdown of Opiate Abusers

Asia: 54%
Europe: 25%
Americas: 14%
Africa: 6%
Oceania: 1%

Notes
Proportions of heroin abuse vary by region:
- Almost all of opiate consumers in Africa are reportedly abusing heroin.
- 2/3 of opiate abusers consume heroin in Asia. Use of opium is still widespread in a number of countries.
- Asia & Europe together account for 80% of the world’s heroin abusers.

Sources: UNODC, Annual Reports Questionnaire Data, Govt. reports, reports of regional bodies, UNODC estimates.
### Annual Prevalence of Opiate Abuse, 2003 - 2005

<table>
<thead>
<tr>
<th>Region</th>
<th>Abuse of opiates</th>
<th>of which abuse of heroin</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Number of abusers</td>
<td>in % of population age 15-64</td>
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<tr>
<td><strong>EUROPE</strong></td>
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<tr>
<td>West &amp; Central Europe</td>
<td>1,565,000</td>
<td>0.5</td>
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<tr>
<td>South-East Europe</td>
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<tr>
<td><strong>ASIA</strong></td>
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<td>Oceania</td>
<td>90,000</td>
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<tr>
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<tr>
<td><strong>GLOBAL</strong></td>
<td>15,840,000</td>
<td>0.4</td>
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</tbody>
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- Above global average
- Around global average
- Below global average

Sources: UNODC, Annual Reports Questionnaire data, various Govt. reports, reports of regional bodies, UNODC estimates.
Trends in Opiate Use

Notes

Opiate Abuse Levels

- Rising in Asia, mainly among countries close to Afghanistan, though falling in East & South East Asia (reflecting the strong declines of opium production in Myanmar & Lao PDR).

- Stable decline in West & Central Europe; rising in East Europe (Eastern Europe suffering from the supply push of Afghan opiates).

- Stable decline in Americas (Falling opium production levels in Latin America & South-East Asia, the two main traditional supply lines for the North American market, may have contributed to this.

-Oceania continues to remain below levels recorded in 2000 (major heroin shortage in 2001, prompted by the dismantling of some major heroin trafficking networks).

-Opiate use in Africa starts rising (upward trend is particularly noted in South Africa, where heroin used to account for less than 1% of treatment demand; by the first two quarters of 2005, this proportion had increased to 7%).

Sources: UNODC, Annual Reports Questionnaire Data, Government reports, UNODC Field Offices, UNODC’s Drug Abuse Information Network for Asia and the Pacific (DAINAP), EMCDDA, CICAD, HONLEA reports and local studies.
Change in Abuse of Heroin and Other Opiates
(2004, or latest year available)

Map & Changes in abuse of heroin and other opiates, 2004 (or latest year available)

Opioids

**Opiate (n)**

“An unlocked door in the prison of identity. It leads to the jail yard.”

Ambrose Bierce
*The Devil’s Dictionary* (1906)

**Notes**

Opium is the milky juice or dried exudate of the opium poppy. In the week after the flowers fall off, the pod, if cut, will excrete a tar-like substance. The tar is brown in colour, and has an unpleasant odour and bitter taste. The gum is drained, dried, boiled in water and filtered to produce opium paste. Two additional products can then be isolated (morphine and codeine).

Heroin (diacetylmorphine), a semi-synthetic substance, is the result of a chemical process that combines opium with two additional molecules.

Opium contains around 1–15% morphine, 1–2% codeine, and 75–80% substances which have little or no pharmacological activity (Victoria Police, 2001).

Opium poppies are grown in the Middle East, Asia, China, Afghanistan, and increasingly, the Americas.


Medicine in Quotations Online [www.acponline.org/cgi-bin/medquotes](http://www.acponline.org/cgi-bin/medquotes)

Opioid-related problems

- Most prominent problems are associated with heroin dependence.
- Not all users of heroin develop dependence. Between 1 in 4 to 1 in 3 regular users develop dependence.
- Development of heroin dependence usually requires regular use over months (or longer, when use is more irregular).
The revolving door

- Heroin dependence is a chronic, relapsing disorder. It is a dependency that is very difficult to resolve.
- Relapse is extremely common. It is part of the process of resolving the dependence – much like giving up tobacco.
- A principle health care objective is to get the patient into treatment, help keep them in treatment, and return them to treatment when relapse occurs.

Notes
Heroin dependence is a chronic, relapsing–remitting condition. Long-term follow-up of those entering treatment suggests:

- Only 10% of heroin users will become and remain abstinent in the first year after treatment
- Approximately 2%–3% of people who use heroin will achieve abstinence and remain abstinent in each subsequent year.

Polydrug use: Patterns and risks

- Polydrug use is the norm among drug users
- Most people who use illicit drugs use a variety of different drugs
- Heroin users also are heavy users of alcohol and benzodiazepines
- As CNS depressants, these combinations are especially dangerous and known to be significant contributors to overdose
- Patients should be advised against the use of these combinations and told of the risks involved
Detecting opioid dependence

**Look for a pattern (not an isolated event):**
- In which a patient frequently runs out of scripts for a prescribed opioid
- In which a patient is on a high and increases the dose of prescribed opioids
- In which a patient injects oral medications
- Of observed intoxication or being in withdrawal
- Which presents plausible conditions that warrant prescribed opioids, but with specific requests for medication type and amount
- In which the patient threatens or harasses staff for a fit-in appointment
- In which a patient alters, steals, or sells scripts
- In which a patient is addicted to alcohol or other drugs
Classification of Opioids

Full Agonists

- Non-synthetic
  - opium
  - papaverine
  - morphine
  - codeine

- Semi-synthetic
  - heroin
  - hydromorphone
  - oxycodone

- Synthetic
  - LAAM
  - fentanyl
  - Meperidine / pethidine
  - hydrocodone
  - methadone
  - pentazocine

Partial Agonists

- buprenorphine

Antagonist

- naltrexone

Notes

Pure opioid agonists: of 20 naturally occurring alkaloids, only morphine and codeine have analgesic properties.

Semisynthetics: e.g., heroin and “homebake” (morphine made out of codeine by a home-made method) are chemical derivatives of morphine.

Synthetics: e.g., methadone and dextropropoxyphene, share a common structure that enables interaction with opioid receptors. These entirely artificial drugs have been synthesised without commencing the process with a naturally occurring opioid.

Commonly used opioid-based preparations include:

- heroin/homebake
- morphine/morphine-based medications such as Pethidine
- codeine phosphate and codeine based preparations, e.g., cough mixtures, and preparations such as ‘Codral Forte’, ‘Panadeine Forte’, ‘Mersyndol Forte’
- methadone
- oxycodone-based medications such as ‘Endone’ and ‘Prolodone’
- dextropropoxyphene contained in medications such as ‘Digesic’, ‘Doloxene’ and dextromoramide (Palfium).

Sources:


Notes

Refer also to next slide.

Three main types of opioid receptors in the CNS and periphery have been identified – κ, δ and μ. It is also believed that there are several other subtypes whose characteristics are yet to be determined. There are also four groups of endogenous peptides (enkephalins, endorphins, dynorphins, and endomorphins) produced by peptidases that cleave inactive precursor peptides. Opioid peptides and their receptors are widely distributed through the CNS and non-neuronal tissues, such as the GI tract. Opioid receptors, acting via G-proteins, are inhibitory. They inhibit adenylate cyclase, open potassium channels, and block voltage-gated calcium channels, therefore reducing neurotransmitter release (5-HT, acetylcholine, glutamate and GABA) (Young et al., 2002, p. 80).


Picture: Society for Neuroscience
http://apu.sfn.org/content/publications/BrainBriefings/addition.html#fullsize

Notes:

“PET” stands for “Positron Emission Tomography.”
Pharmacodynamics

Opioids act on 3 main families of opioid receptors (µ, κ, and σ). Endogenous opioid peptides (e.g. enkephalins, endorphins, dynorphins, endomorphins) produced by peptidases also bind with opioid receptors. Both the opioid peptides and receptors are located in the brain, spinal cord, and periphery (including non-neuronal pathways in the GI tract), resulting in effects such as cough and respiratory suppression, reduced GI motility (hence nausea and vomiting), miosis, and urinary retention (from increased bladder and urethral tone). Effects exerted through the limbic system produce changes in emotions, such as the euphoric high.

Opioid receptors are inhibitory and act via G-proteins. They inhibit adenylate cyclase, open potassium channels and block voltage-gated calcium channels thereby inhibiting the release of neurotransmitters such as 5-HT, GABA, glutamate and acetylcholine (Young et al., 2002, p. 80).

The endogenous opioid system is activated by stress. It can modulate pain perception, mood and physiological systems (e.g., the respiratory or immune systems) (Young et al., 2002, p. 81).

All prescription opioids produce morphine-like effects but rather than removing pain, they alter perceptions of the pain so that it is more tolerable and less aversive. Although cognition is impaired, consciousness and coordination are generally intact at low doses. Opioids produce analgesia and euphoria, decrease muscle tone, slow movement of the digestive tract, may alter hormonal balance and have a role in regulating immune function. Inhibition of the respiratory system and potential for overdose occur due to the brainstem response to carbon dioxide.

Opioids are distinguished from sedative hypnotics through their powerful analgesic, anti-diarrhoeal, and cough suppressant properties.

Heroin

- Morphine is produced through heroin hydrolysis
  \[ \text{heroin} \rightarrow \text{monoacetylmorphine (MAM)} \rightarrow \text{morphine} \]
- Heroin and MAM are lipophilic, hence more rapid action
- Heroin excreted in urine as free and conjugated morphine
- Heroin metabolites are present in urine for approximately 48 hours following use

Notes

Pharmacokinetics

The variety of chemical structures in the opioid class result in important differences in their pharmacokinetics. Although most are metabolised by oxidation, morphine and buprenorphine are conjugated with glucuronic acid in the liver. As morphine is rapidly metabolised by the liver after oral administration, only a small amount reaches systemic circulation (Young et al., 2002, p. 81).

Heroin (diacetylmorphine) is hydrolysed firstly to monoacetylmorphine (MAM), then to morphine. As heroin and MAM are more lipophilic than morphine, they cross the adult blood-brain barrier more rapidly than morphine, resulting in feelings of euphoria.

Codeine is also converted to morphine via demethylation by the enzyme CYP2D6.

For the 8–10% of Caucasians and 2% of South-East Asians who do not have the enzyme CYP2D6, codeine will have no analgesic effect.


Morphine: Immediate effects (1)

- Perception altered, possible delirium
- Analgesia, to some degree
- Impaired cognition, though consciousness may be preserved
- Autonomic nervous system affected
- Suppression of cough reflex
- GI system affected
- Hypothermia

Notes

Perception: euphoria, flushing, sense of tranquillity, peace or contentment.

Analgesia: pain is not removed but perception of pain altered so that the experience is no longer aversive.

Impaired cognition: consciousness and coordination intact at low doses.

Autonomic nervous system: reduced brainstem response to CO₂ inhibits respiratory system; low blood pressure.

Suppression of cough reflex, nausea and vomiting: opioids stimulate the chemoreceptor trigger zone in medulla.

Through activation of the µ receptors in the mesolimbic reward pathway, morphine increases the release of dopamine through inhibiting GABA interneurones. The release of dopamine is believed to contribute to the dependence-producing potential of opioids. The euphoric effects of opioids, especially when injected, can be highly reinforcing to vulnerable individuals. Effects such as euphoria, flushing and the abdominal ‘buzz’ (described by many as akin to orgasm) are specific to recreational experiences and are not generally seen when opioids are used in clinical situations.

All opioids exert a morphine-like effect, producing drowsiness, clouding of sensorium and perception, mood changes (usually euphoria or contentment), analgesia and respiratory depression. The CNS depressant effects can be reversed by the opioid antagonist naloxone.

At high doses, the muscle tone of the large trunk and intercostal muscles may increase (tightly), hence further impairing breathing.

Morphine: Immediate effects (2)

- Miosis
- Urinary retention
- Reduced GI motility
- Endocrine
- Non-cardiogenic pulmonary oedema
- Coma or death (from respiratory depression)
- Other
  - pruritis; flushed skin; dry mouth, skin, and eyes

Notes

**Miosis**: due to increasing parasympathetic tone in the pupil.

**Urinary retention**: increased urethral and bladder tone.

**Reduced GI motility**: opioid receptors are present in GI tract. Reduced GI motility can result in constipation. Opioids increase muscle tone, specifically affecting the Sphincter of Oddi (increasing the muscle tone).

**Endocrine**: changes sex hormones in women – decreased follicle-stimulating hormone (FSH) and lutenising hormone (LH); raised prolactin resulting in menstrual changes, reduced libido, galactorrhoea; reduced testosterone in men with reduced libido. Also increases ADH, decreases ACTH.

Tolerance to opioids develops rapidly, commencing with the first dose and involves:
  - down-regulation – reduced number of receptors
  - desensitisation – diminished response to receptor action.

Opioids: Long-term effects (1)

- Little evidence of long-term direct toxic effects on the CNS from opioid use
- Long-term health-related complications may result from:
  - dependence
  - poor general self-care
  - imprisonment
  - drug impurities or contaminants, BBV

Notes
There is little evidence of long-term direct toxic effects on the CNS from using opiates. (See next slide for chronic use complications).

Notes

There is little evidence of long-term direct toxic effects on the CNS from using opioids. However, the following complications may result from long-term chronic opioid use.

**Narcotic bowel syndrome**
- Characterised by bloating, vague abdominal discomfort
- Physical examination and investigations are negative though patients may have a dilated bowel (with no obstruction)
- Intervention – taper to discontinue the drug use.

**Medication induced headaches**
- This condition generally refers to patients who are not regular heroin users but who are receiving mixed opioid/non-opioid analgesics such as paracetamol with codeine for management of migraine. Patients may report increased headache frequency since commencing the use of opioid-based medications which stop on cessation of analgesia.

**Depression**
- Changing drug-use behaviours requires significant social change. It is not unusual for patients to experience depression or sadness in the face of significant change and take time to adjust to a different lifestyle. Ongoing assessment is important to ensure adequate support is provided and for detecting the possible emergence of any mental health problems.

Notes

This slide highlights areas of special consideration when assessing or treating an injecting drug user who uses opioids.

It is advisable to prescribe methadone for opioid-using pregnant patients. Cessation of opioids is not recommended because of risks to the fetus from withdrawal. Refer patients to an authorised methadone prescriber.

Urinalysis:

• may be valuable in confirming drug use history, although this is an expensive process and the results are not immediately available
• indicates evidence of recent use but does not identify dependence, nor does it indicate problem areas
• does little to assist in building rapport with patient.

Physical exam

Signs of opioid dependence:
- Needle marks on wrists, antecubital fossa, legs (inner thighs), feet, hands, neck
- Intoxication: pinpoint pupils, “nodding off,” drowsiness, sweating

Notes
Note that track marks are not always in the obvious locations and some injecting drug users will go to considerable lengths to use sites that are less obvious and less easy to detect on examination, e.g. soles of the feet.
The following slides depict complications from use, dependence, and overdose.
Notes

Extensive “track marks” – IV drug use.
Notes

A typical track mark due to IV heroin use.
The person is pointing to slight inflammation (red line) up the arm. This is thrombophlebitis – inflammation of the vein.
Plastic surgery can remove the track so it is less obvious.
Notes

Venous abscess – IV drug user.
Infections may also include septicaemia or a septic joint in IV drug users.
Hot, painful joints should be assumed to be septicaemia until proven otherwise.
Opioid withdrawal

**Signs**
- Yawning
- Lacrimation, mydriasis
- Diaphoresis
- Rhinorrhea, sneezing
- Tremor
- Piloerection
- Diarrhoea and vomiting

**Symptoms**
- Anorexia and nausea
- Abdominal pain or cramps
- Hot and cold flushes
- Joint and muscle pain or twitching
- Insomnia
- Drug cravings
- Restlessness / anxiety

**Notes**
People using opioids may experience a moderate to severe but not life-threatening withdrawal syndrome.

The onset and duration of withdrawal varies according to the half life of the drug used, e.g., withdrawal symptoms from heroin (usually manifest in a marked drive to obtain and use the drug) may commence 6–12 hours after the last dose, and may last for 5–7 days. With methadone, withdrawal may not commence for 2–3 days after most recent dose and last for up to 3 weeks.

Knowledge of the half-life of the drugs used (methadone vs. heroin) and the time for likely onset of withdrawal after the last dose can assist in predicting, identifying, and managing opioid withdrawal.

Signs and symptoms of opioid withdrawal may be mistaken for a bad dose of the ‘flu’.

Despite depictions of heroin withdrawal in popular culture, opioid withdrawal is rarely, and is unlikely to be, fatal.

Withdrawal (and the culture or lifestyle associated with use, or withdrawal from that lifestyle) may precipitate dysthymia or depression.

Notes
Dilated pupils – opiate withdrawal.
Pupils will be constricted when intoxicated.
Progress of the Acute Phase of Opioid Withdrawal Since Last Dose

Withdrawal from heroin
Onset: 6–24 hrs
Duration: 4–10 days

Withdrawal from methadone
Onset: 24–48 hrs, sometimes more
Duration: 10–20 days, sometimes more

Severity of signs and symptoms

Days


Predictors of withdrawal severity

- Main predictors
  - Greater regular dose
  - Rapidity with which drug is withdrawn
- Also consider
  - Type of opioid used, dose, pattern, and duration of use
  - Prior withdrawal experience, expectancy, settings for withdrawal
  - Physical condition (poor self-care, poor nutritional status, track marks)
  - Intense sadness (dysthymia, depression)
  - Constipation or "Narcotic Bowel Syndrome"
  - Impotence (males) or menstrual irregularities (females)

Notes

Signs on the slide are indicative of long-term use and may predict severity of withdrawal. Despite potential severity, opioid withdrawal does not present a risk for fatality, except in the neonate or when other significant medical conditions are present.
Opioid withdrawal scales

Withdrawal scales:

- guide treatment
- monitor progress of withdrawal (subjective and objective signs)
- do not diagnose withdrawal but describe severity
- guide ongoing assessment

*If the withdrawal pattern is unusual, or the patient is not responding, suspect other conditions.*

Tools
The Subjective Opioid Withdrawal Scale (SOWS) (See Handouts).
The Objective Opioid Withdrawal Scale (OOWS) (See Handouts).
Use of the SOWS during assessment enables patients to be involved in their own care, and can assist in reducing their anxiety.


Withdrawal management aims to:

- reverse neuroadaptation by managing tolerance and withdrawal
- promote the uptake of post-withdrawal treatment options

Withdrawal management may occur:

- as an outpatient
- in a residential / treatment setting

Opioid withdrawal treatment

Involves:
- reassurance and supportive care
- information
- hydration and nutrition
- medications to reduce severity of somatic complaints (analgesics, antiemetics, clonidine, benzodiazepines, antispasmodics)
- opioid pharmacotherapies (e.g., methadone, buprenorphine)

Notes
Reassurance, arranging supportive care, insuring adequate hydration and nutrition, and providing accurate information about withdrawal (what to expect) for patients and their caregivers can significantly reduce anxiety and assist in the effective management of the patient.

A range of medications can assist in reducing the severity of somatic complaints and increase the comfort of the patient.

Buprenorphine is increasingly used for withdrawal management, as it:
  - offers less intense withdrawal compared with methadone tapering
  - has fewer side-effects when compared to clonidine.

The main complications from opioid withdrawal are not life-threatening.


Notes:
Opioid detoxification by itself should not be considered “treatment” for heroin addiction.
Opioid withdrawal complications

- Anxiety and agitation
- Low tolerance to discomfort and dysphoria
- Drug-seeking behaviour (requesting or seeking medication to reduce symptom severity)
- Muscle cramps
- Abdominal cramps
- Insomnia

Opioid withdrawal

- Non-life threatening
- Commences 6 – 24+ hours after last use
- Peaks at around 24 – 48 hours after use
- Resolves after 5 – 7 days

There is increasing recognition of the existence of a protracted phase of withdrawal lasting some weeks or months, characterised by reduced feelings of wellbeing, insomnia, dysthymia, and cravings.

Notes

Protracted phase – monitor for dysthymia/depression, which may need to be treated.

Note: hallucinations and seizures are not features of heroin withdrawal except in neonates. Assess for other conditions.
**Notes**

This slide depicts the various treatment options and pathways available for dependent patterns of opioid use and the role of harm reduction and relapse prevention strategies as valuable components of the model.

**DSM IV criteria for opioid dependence**

- Tolerance
- Withdrawal symptoms on cessation of drug use
- Increasing quantity or frequency of use
- Persistent desire for the drug or unsuccessful attempts to cut down
- Salience of drug use over other responsibilities (most of a patient’s time involves taking, recovering from, or obtaining drugs)
- Continued use despite evidence of psychological or social problems

**Notes**

About one in three heroin users develops dependence.

Dependence has grades of severity – it is not an “all or nothing” phenomenon.

The most salient feature of the dependence syndrome is loss of control over the use of a drug, with persistent use despite significant harms.

Physical dependence is not a requisite for drug dependence.

Most dependent heroin users describe first using heroin in their late teens to early twenties, with regular use usually commencing several years later.

Heroin dependence is a chronic, relapsing–remitting condition. Long-term follow-up of those entering treatment suggests:

- 10% of heroin users will become and remain abstinent in the first year after treatment
- Approximately 2%–3% of people who use heroin will achieve and remain abstinent in each subsequent year.

**Some characteristics of dependent heroin use in Australia**

- Dependent heroin use is difficult to sustain for most people.
- Heroin is a short-acting drug: 2 to 4 injections a day is common.
- Illicit heroin has variable concentration and adulterants, and is expensive (costing $50 to $200 per day in 2001).
- Stigma associated with heroin use can deter people from seeking treatment or disclosing their drug use to family, friends, work colleagues, and health workers.

Polydrug use is common: Over half of dependent heroin users use cannabis regularly and approximately one third used benzodiazepines within last month.

ICD-10 criteria for opioid dependence

- A strong desire or sense of compulsion to take opioids
- Difficulties in controlling opioid-taking behaviour (onset, termination, or levels of use)
- A physiological withdrawal state
- Evidence of tolerance
- Progressive neglect of alternative pleasures or interests because of opioid use
- Use of opioids despite overtly harmful consequences

Additional information on diagnostic guidelines
A definite diagnosis of dependence should usually be made only if three or more of the following have been experienced or exhibited at some time during the previous year:
(a) a strong desire or sense of compulsion to take opioids;
(b) difficulties in controlling opioid-taking behaviour in terms of its onset, termination, or levels of use;
(c) a physiological withdrawal state when opioid use has ceased or been reduced, as evidenced by the characteristic withdrawal syndrome for opioid; or use of the same (or a closely related) substance with the intention of relieving or avoiding withdrawal symptoms;
(d) evidence of tolerance, such that increased doses of opioid are required in order to achieve effects originally produced by lower doses (clear examples of this are found in opioid-dependent individuals who may take daily doses sufficient to incapacitate or kill nontolerant users);
(e) progressive neglect of alternative pleasures or interests because of opioid use, increased amount of time necessary to obtain or take the substance or to recover from its effects;
(f) persisting with opioid use despite clear evidence of overtly harmful consequences, such as depressive mood states consequent to periods of heavy substance use, or drug-related impairment of cognitive functioning; efforts should be made to determine that the user was actually, or could be expected to be, aware of the nature and extent of the harm.

Narrowing of the personal repertoire of patterns of opioid use has also been described as a characteristic feature.

It is an essential characteristic of the dependence syndrome that either opioid-taking or a desire to take opioids should be present; the subjective awareness of compulsion to use drugs is most commonly seen during attempts to stop or control substance use. This diagnostic requirement would exclude, for instance, surgical patients given opioid drugs for the relief of pain, who may show signs of an opioid withdrawal state when drugs are not given but who have no desire to continue taking drugs.

(Source: ICD-10 World Health Organization. Available at http://azpsychiatry.info/icd/substance/dependence/opioid.htm)
### General principles of pharmacotherapies: Pharmacodynamics

- **Agonists**
  - directly activate opioid receptors (e.g., morphine, methadone)

- **Partial agonists**
  - unable to fully activate opioid receptors even with very large doses (e.g., buprenorphine)

- **Antagonists**
  - occupy but do not activate receptors, hence blocking agonist effects (e.g., naloxone)

## Notes

Both agonists and antagonists can demonstrate selectivity for specific receptor sub-types, e.g.:

- as a partial agonist, buprenorphine can block some of the reinforcing effects of full $\mu$ agonists
- as a mixed kappa agonist and $\mu$ antagonist, pentazocine produces dysphoria rather than euphoria.

Mixed agonist/antagonists may precipitate withdrawal if given to a person who is opioid dependent.

Pure antagonists, e.g., naloxone/naltrexone, have no analgesic or other reinforcing properties, hence they are suitable for maintenance treatment.

Maintenance pharmacotherapies

- Methadone
- Buprenorphine
- LAAM

Notes

Until recently, methadone was the most studied treatment modality for responding to opioid dependence. Patients prescribed methadone have been found to:

- reduce their use of drugs
- be more likely to be retained in treatment
- have improved health, social relationships and general functioning, including
  - reduced risk of transmission of BBV through reduced frequency of injecting
  - opportunity to withdraw from drug-using lifestyle
  - stable dosing, which enables stability and development of routine.
- have reduced chance of risky behaviours associated with opioid use and premature death
- have reduced participation in criminal activities.

LAMM has been withdrawn in Europe. LAAM was approved for clinical use in America (1994) and in Europe (1997).

However, LAAM was subsequently withdrawn in Europe after several cases of serious cardiac arrhythmias associated with QT prolongation, torsade de pointes, cardiac arrest, and deaths. In the U.S., the Food and Drug Administration did not completely suspend LAAM, but recommended using other opiate pharmacotherapy first. (Fegus Law)
Key outcomes of maintenance pharmacotherapy programs

- ↑Retention in treatment
- Facilitates reduction / cessation of opioid use
- Reduces risky behaviours associated with opioid use
- Enables opportunity to engage in harm reduction measures
- ↓Mortality and morbidity
- ↑Psychological, emotional, and physical wellbeing of patients
- ↓Social costs associated with illicit drug use
- ↓Crime

Notes

Patients are more likely to cut down or cease use as they age and mature. Retaining them in treatment is the first priority. GPs can greatly assist in initiating and supporting patients’ engagement in treatment.

Involvement in a comprehensive treatment program is crucial to insure the provision of patient supports required to adjust to new lifestyles and to take the opportunities afforded by maintenance therapies. The legal, social, health, and lifestyle changes that occurred when a patient was (or is still) using drugs may take considerable time to resolve. Hence, the support offered by members of the health team can significantly assist the patient and the treatment plans.

Harm reduction measures reduce mortality and morbidity rates associated with opioid use. Pharmacotherapies increase psychological, emotional, and physical wellbeing of patients.

Methadone: Clinical properties

The “Gold Standard” Treatment
- Synthetic opioid with a long half-life
- µ agonist with morphine-like properties and actions
- Action – CNS depressant
- Effects usually last about 24 hours
- Daily dosing (same time, daily) maintains constant blood levels and facilitates normal everyday activity
- Adequate dosage prevents opioid withdrawal (without intoxication)

Notes
Methadone is recognised as the “gold standard” of treatment for managing opioid dependence and has been found to be an effective public health and harm reduction measure. Its use is generally restricted to specific medical conditions, such as opioid dependence and the management of chronic pain.

In Australia, methadone is provided by public government services (public programs) and privately through trained GP prescribers.

Methadone is highly effective when taken orally. When used repeatedly, such as during maintenance for opioid dependence, its effects persist and the duration of its effect is extended.

Although a potent analgesic for chronic pain, the analgesic effect can lasts for 24 hours (variable) because of its variable half-life.

Methadone:
- is detectable in plasma for 30 minutes following ingestion and it might be detected in plasma for hours to days
- has a peak concentration after about 4 hours
- has a single dose half-life of 15–22 hours (high variability)
- has a maintenance dosing half-life of 22 hours and suppression of withdrawal for 24–36 hours
- stability varies with metabolic rate, which varies according to genetic makeup and environmental and disease-state factors (e.g., pregnancy increases methadone metabolism)
- oral form is only marginally less potent than IM form.

Buprenorphine

- Derived from the morphine alkaloid thebaine
- Partial opioid agonist at µ opioid receptors
- Antagonist at k opioid receptor
- Blocks opioid receptors, diminishes cravings, prevents opioid withdrawal

Notes

Blocks opioid receptors (which blocks the effects of extraneous opioids), thereby diminishing cravings for opioids and preventing opioid withdrawal.

Buprenorphine:

- is a partial opioid agonist at µ receptors. It can displace other opioids competing at the same receptor. It may therefore precipitate onset of withdrawal on initial administration. It has an euphoric effect but a less sedating effect than full opioid agonists
- binds strongly to the receptor and is not easily displaced
- also a kappa opioid receptor antagonist

Metabolism occurs through two pathways:

- conjugation with glucuronic acid
- N-de-alkylation

Metabolites are excreted in the biliary system, with enterohepatic cycling of buprenorphine and its metabolites, mainly in urine and faeces

Pain management: includes post-operative, terminal and chronic pain

Extended duration of action thought to relate to:

- a high affinity to µ receptors
- high lipophilicity (low levels are released from fat stores with chronic dosing)
- reabsorption after intestinal hydrolysis of conjugated metabolites

Sources: adapted from CDHA (Commonwealth Department of Health and Ageing) 2002, Illicit Drug Training for Pharmacists, CDHA, Canberra, pp. 89–90.
Buprenorphine vs. Methadone

### Buprenorphine Advantages
- Milder withdrawal
- Convenient (dose every 2/7)
- Relative ease of use, i.e., ready transmission from heroin withdrawal state or methadone
- Easier to taper than methadone
- Wider safety margin

### Buprenorphine Disadvantages
- SL route results in reduced bio-availability compared with IV preparations
- Difficult to reverse respiratory depression if it does occur
- Increased time required for supervised dosage (to get dissolution)
- Risk of abuse difficult to supervise
- Failure to reduce cravings in people with significant emotional trauma.

**Notes**
As a partial agonist, buprenorphine induces a lower level of dependence, hence withdrawal is significantly easier.

Buprenorphine is convenient – patients able to travel short distances (with weekend pickup) and takeaways, decrease chemist contact.

Less likely to be diverted or sold on the streets, as it may precipitate withdrawal in opioid-dependent people.

Safer in accidental overdose (e.g., children) because of poor oral absorption (therefore less respiratory depression).

Initially the pharmacists’ practice of crushing tablets was intended for those suspected of diversion. However, recent changes in practice suggest that crushing is increasingly common, and that both pharmacists and patients prefer the tablets crushed as this enables the patient to feel the tablet dissolving and reduces the time spent in the pharmacy. Few people report problems with crushed tablets. Others suggest that crushing decreases the bio-availability of buprenorphine. Further research is needed on this.


**Notes:**
Dosing every two days may not be suitable for a proportion of patients.

The main disadvantages of buprenorphine are as follows:
(1) risk of abuse, as it is difficult to supervise
(2) failure to reduce cravings in some people with significant emotional “trauma”
Rationale for opioid agonist / partial agonist treatment

Advantages of opioid agonist / partial agonist medication over heroin

- Non-parenteral administration
- Known composition
- Gradual onset and offset
- Long-acting
- Far less reinforcing than heroin
- Medically supervised

Notes

Medications such as methadone, LAAM, and buprenorphine have several advantages over heroin. They can be administered by safer routes (oral or sublingual, rather than by injection); they are long-acting (so that dosing is daily or several times per week, rather than several times per day); they have known composition (so that dosing can be quantified and constant, and so that contaminants are eliminated and there is a known level of purity); their onset of action is gradual and their effects are mildly reinforcing (insuring compliance in taking the medication while decreasing abuse potential); and they are managed under medical supervision.
Opioid agonist treatment

- Most effective treatment for opioid dependence
- Controlled studies have shown that with long-term maintenance treatment using appropriate doses, there are significant:
  - Decreases in illicit opioid use
  - Decreases in other drug use

Notes
The prototypic opioid agonist maintenance medication is methadone. Controlled studies have shown that methadone, when delivered properly, can be a highly effective medication. Improvements among opioid-dependent patients treated with methadone are not limited to decreases in illicit opioid use. Methadone treatment can result in significant decreases in other drug use, and improvements in other areas (such as employment). Further information about methadone (and LAAM) will be provided in a later section.

References:
Ball J.C., Ross A. The Effectiveness of Methadone Maintenance Treatment. Springer-Verlag, New York, 1991; pages 166-168; 181-182.
Opioid agonist treatment (continued)

- Decreases in criminal activity
- Decreases in needle sharing and blood-borne virus transmission (including HIV)
- Improvements in pro-social activities
- Improvements in mental health
Injecting Drug Use and HIV/AIDS

Estimated number of deaths from AIDS up till now: 25 million

Estimated number of people with HIV infection in 2002/2003: 42 million

Estimated number of additional HIV infections till 2010: 45 million.

Notes
Injecting drug use is a major form of transmission of HIV/AIDS. Anyone engaging in injecting behaviour should be counselled about risks and illicit drug users advised to use safer routes of administration. Further information on harm reduction strategies (preventing HIV, etc.) is available in Volume D Topic 4 (Harm Reduction and HIV Risk Reduction Strategies) of the training materials.

By 2010, AIDS will have caused more deaths than any disease outbreak in history.

Injecting drug use is an important contributor to the spread of HIV.
91% of the world adult population (4 billion) is covered by the data. Information unavailable for 119 countries.

UN Reference Group on HIV/AIDS prevention and care among IDU
www.idurefgroup.org
WHO / UNODC / UNAIDS position paper: *Substitution Maintenance Therapy in the Management of Opioid Dependence and HIV/AIDS Prevention*

“Substitution maintenance treatment is an effective, safe and cost-effective modality for the management of opioid dependence. Repeated rigorous evaluation has demonstrated that such treatment is a valuable and critical component of the effective management of opioid dependence and the prevention of HIV among IDUs.”

Notes

The position paper *Substitution Maintenance Therapy in the Management of Opioid Dependence and HIV/AIDS Prevention* is available online at:

### Availability of Substitution Treatment

**85% + methadone is consumed in developed countries (2002)**

<table>
<thead>
<tr>
<th>Country</th>
<th>%</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>US</td>
<td>53%</td>
<td>8.7 tons</td>
</tr>
<tr>
<td>Spain</td>
<td>11%</td>
<td>1.8 tons</td>
</tr>
<tr>
<td>Germany</td>
<td>6%</td>
<td>916kg</td>
</tr>
<tr>
<td>Italy</td>
<td>5%</td>
<td>812kg</td>
</tr>
<tr>
<td>UK, Canada, Australia,</td>
<td>18%</td>
<td></td>
</tr>
<tr>
<td>Switzerland, France, Denmark,</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Belgium, etc.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Substitution treatment is also available in the following countries:**
- Argentina
- China
- Croatia
- India
- Indonesia
- Iran
- Kyrgyzstan
- Malaysia
- Moldova
- Nepal
- Singapore
- Thailand
- Ukraine

Most of the rest consumed by 8 other countries, mostly in Europe, and Australia

---

**Notes:**

Acknowledgments: Thanks to Gerry Stimson.
Estimated Opiate-Dependent Drug Users in Substitution Treatment per 100,000 Population

- Australia
- Spain
- United States
- Netherlands
- Italy
- UK
- Germany
- Denmark
- France
- Canada
- Sweden
- Thailand
- China
- India
- Nepal

0 50 100 150 200
Naltrexone

- Morphine antagonist, true blockade
- No direct psychoactive effect
- No withdrawal experienced upon cessation
- Reported to reduce cravings in some people

Notes
Because of the lack of psychoactive effect, naltrexone is not self-reinforcing.
Naltrexone: Mechanism of action

- Fully blocks μ receptors, preventing euphoria from opioid use; therefore
  
  “drug money spent = money wasted”

- Allows extinction of Pavlovian-conditioned response to opiate cues

- Prevents reinstatement of opioid dependence, but does not reinforce compliance
Naltrexone: Indications for use

- Prescribed for the management of opioid dependence by registered prescribers
- Primary role = relapse prevention
- Abstinence-based treatment option
- Non-dependence inducing
- Commenced at least 1 week after cessation of heroin use
- Optimally effective with motivated individuals who have higher levels of psychosocial functioning and family support
- Young addicts or short duration of dependence

Additional information on Naltrexone:

Because of the potential to induce severe withdrawal, naltrexone should only be commenced after the patient has been at least:
- 7 days heroin-free
- 10 days methadone-free.

Because Naltrexone does not have narcotic effect, it does not produce any withdrawal symptoms. Despite its potential advantage, it has little impact on the treatment of opioid addiction in the U.S. because of poor patient compliance.

It is important to mention that some researchers advise of the risk of heroin overdose due to the fact that patients may stop using naltrexone and relapse to heroin. However further research is needed to validate this concern.

(Source: U.S. Department of Health and Human Services. Substance Abuse and Mental Health Services Administration, Center for Substance Abuse Treatment. Medication-Assisted Treatment for Opioid Addiction in Opioid Treatment Programs: TIP 43. Available at www.samhsa.gov)
Questions?

Comments?
Thank you for your time!

End of Workshop 1
Workshop 2: Opiate Addiction Treatment with Methadone
At the end of this training, you will know:

1. The rationale for opiate agonist therapy
2. Medical withdrawal protocols using methadone
3. The basic purpose and background evidence to support the use of methadone for treating opiate dependence
4. The basic principles of maintenance treatment with methadone
5. Effective practices (evaluation, initial dose and management of dose; tapering procedures, etc.) in the implementation of methadone treatment
6. How to address concurrent use of other drugs and alcohol during methadone treatment
7. The contraindications and medical interactions with methadone
Methadone: Clinical properties

The “Gold Standard” Treatment
- Synthetic opioid with a long half-life
- μ agonist with morphine-like properties and actions
- Action – CNS depressant
- Effects usually last about 24 hours
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Methadone is recognised as the “gold standard” of treatment for managing opioid dependence and has been found to be an effective public health and harm reduction measure. Its use is generally restricted to specific medical conditions, such as opioid dependence and the management of chronic pain.

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Although a potent analgesic for chronic pain, the analgesic effect lasts for less than 24 hours because of its variable half-life.

Methadone:
- is detectable in plasma for 30 minutes following ingestion
- has a peak concentration after about 4 hours
- has a single dose half-life of 15–22 hours (high variability)
- has a maintenance dosing half-life of 22 hours and suppression of withdrawal for 24–36 hours
- stability varies with metabolic rate, which varies according to genetic makeup and environmental and disease-state factors (e.g. pregnancy increases methadone metabolism)
- oral form only marginally less potent than IM form.

Intrinsic Activity: Full Agonist, Partial Agonist and Antagonist

(Source: Drug and Alcohol Dependence 70 (Suppl.) Johnson et al. Buprenorphine: How to use it right. 59-77, 2003)
Methadone pharmacokinetics

- Good oral bioavailability
- Peak plasma concentration after 2-4 hrs
- 96% plasma protein bound
- Mean half-life around 24 hrs
- Steady state after 3-10 days

**Metabolism**
- Cytochrome P450 mediated
  - CYP3A4 main
  - also CYP2D6, CYP1A2, CYP2C9 and CYP2C19
- genetic variability
  → risk of drug interactions
Pharmacodynamics

- full opioid agonist
  - Main action on mu receptors
    - inhibit adenyl cyclase = ↓ cAMP
    - ↑ potassium channel opening
    - ↓ calcium channel opening
  - also inhibit serotonin reuptake
  - also non-competitive antagonist NMDA receptor
Safety overview

- Safe medication (acute and chronic dosing)
- Primary side effects: like other mu agonist opioids (e.g., nausea, constipation), but may be less severe
- No evidence of significant disruption in cognitive or psychomotor performance with methadone maintenance
- No evidence of organ damage with chronic dosing

Notes
1. Buprenorphine is a highly safe medication for use in patients with opioid dependence.
2. Note that it is also safe if inadvertently taken by a person who is not physically dependent on opioids (such as a child). In such a case, it is most likely the person would swallow the tablet and experience virtually no opioid agonist effect because of the poor oral bioavailability. Even if the person sucked on the tablet, there is a low likelihood that they would experience serious adverse effects. This is because buprenorphine is a partial opioid agonist, and there is a ceiling in the maximal effects produced.
3. Clinical trials with buprenorphine have found no significant organ damage associated with chronic dosing. However, buprenorphine may be associated with increases in liver function tests, and this may be especially true for patients with a history of hepatitis prior to the onset of buprenorphine treatment. Increases in liver function tests appear to be mild, and it is important to keep in mind that other factors commonly found in opioid-dependent patients (such as hepatitis and alcohol abuse) can lead to elevations in liver function tests.

References:
Methadone: Advantages of treatment

- Suppresses opioid withdrawal
- Pure – no “cutting agents” present
- Oral administration (syrup or tablet forms used)
- Once-daily doses enable lifestyle changes
- Slow reduction and withdrawal can be negotiated with minimal discomfort
- Minimal reinforcing properties, relative to heroin
- Counselling and support assists long-term lifestyle changes
- Legal and affordable – reduced participation in crime
- Few long-term side effects

Notes
There are a few long-term health effects from use of methadone. Those known include:

- weight gain, possibly influenced by fluid retention and dietary changes
- reduced production of saliva – may contribute to dental problems
- endocrine changes – may result in impotence, low libido, disrupted menstrual cycle
- may be harmful in presence of underlying disease, e.g., kidney or liver problems
- some effects disappear when dose is adjusted.
Methadone: Disadvantages of treatment

- Initial discomfort to be expected during stabilisation phase
- Opioid dependence is maintained
- Slow withdrawal (preferably) negotiated and undertaken over a period of months
- Protracted withdrawal symptoms
- Can overdose, particularly with polydrug use
- Daily travel and time commitment
- Variable duration of action
- Diversion

Notes
Commitsing oneself to methadone maintenance therapy (MMT) can be off-putting to many and interferes with work activities or travel arrangements.
Maximising treatment adherence

- Address psychosocial issues as first priority
  - emotional stability
  - "chaotic" drug use
  - accommodation
  - income

- Opioid agonist pharmacotherapy can:
  - address psychosocial instability
  - increase opportunities to directly observe the administration of various HIV therapies
Assessment objectives

- Clarify nature and severity of problems
- Establish a therapeutic relationship
- Formulate problems into a treatment plan
Core assessment issues

- What does the patient want?
- Is the patient dependent?
- What is their level of tolerance?
- Is the patient using/dependent on other drugs?
- What is their motivation for change?
- What social supports exist?
- Are there other co-existing medical and psychiatric conditions?
Drug use history

- **Primary drug**
  - Average daily use (quantity / duration)
  - Time last used
  - Route of administration
  - Age commenced, periods of abstinence
  - Severity of dependence
  - Previous treatment(s)

- **Other drugs**
  - Current and previous
  - Dependence
Medical and psychiatric

- HIV/HCV
- Hepatitis B
- TB
- Pregnancy
- Other major medical conditions
  - Liver
  - Cardiac
- Major psychiatric conditions
  - Depression, suicide, psychosis
- Opioid-related overdose
Psychosocial

- Relationship with family
- Relationship with partner
- Education and employment
- Criminal justice
- Living circumstances
- Sources of income
Examination

- Mental state
  - Mood
  - Affect
  - Cognition
- Injection sites
- Signs of intoxication / withdrawal
- Stigmata of liver disease
- Nutritional state
Induction stabilisation phase (1)

- Dose adequacy and drug interactions
  - Signs of intoxication / withdrawal
  - Frequency of drug use
  - Frequency of sharing

- Case coordination and management
  - Psychological
  - Social
  - Medical
  - Health / welfare system interaction
Induction stabilisation phase (2)

- Risk Assessment
  - Drug use practises
    - polydrug
    - OD
    - sharing
  - Sexual practises
Safe initial dose

- 20 - 30mg methadone is generally safe
- Deaths have occurred with higher starting doses or polydrug use
- It may be safer to start opioid-dependent polydrug users as inpatients
### Methadone: Initial Effects and Side-Effects

- Relief from physical pain
- Feeling of wellbeing
- Constricted pupils
- Vasodilation
- Lowered sex drive
- Nausea and vomiting
- Loss of appetite
- Sweating
- Fluid retention
- Endocrine changes (loss of libido, menstrual changes)

| • Intense constipation |
| • Lowered temperature |
| • Bradycardia |
| • Hypotension |
| • Palpitations |
| • Shallow respirations |
| • Poor circulation |
| • Itching and skin rashes |
| • Recurrent dental problems |

*Polydrug use may cause overdose.*

### Notes

Effects may vary according to the individual, level of neuroadaptation, dosage, frequency taken, etc.
Opioid withdrawal scales

- guide treatment
- monitor progress (subjective and objective signs)
- do not diagnose withdrawal but describe severity
- guide ongoing assessment

If the withdrawal pattern is unusual, or the patient is not responding, suspect other conditions.

Notes
Tools
The Subjective Opioid Withdrawal Scale (SOWS; see Handouts).
The Objective Opioid Withdrawal Scale (OOWS; see Handouts).
Use of the SOWS during assessment enables patients to be involved in their own care and can assist in reducing their anxiety.

Sources: deCrespigny, C., Talmot, J, Modystack, K., Cusack, L. & Watkinson, J. 2003, Alcohol, Tobacco and Other Drugs Guidelines for Nurses and Midwives: Clinical Guidelines, Flinders University and Drug and Alcohol Services Council (DASC), Adelaide.
# Opiate withdrawal scale

<table>
<thead>
<tr>
<th>Resting Pulse Rate: _______ beats/minute</th>
</tr>
</thead>
<tbody>
<tr>
<td>Measured after patient is sitting or lying for one minute</td>
</tr>
<tr>
<td>0 pulse rate 80 or below</td>
</tr>
<tr>
<td>1 pulse rate 83-100</td>
</tr>
<tr>
<td>2 pulse rate 101-120</td>
</tr>
<tr>
<td>4 pulse rate greater than 120</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sweating: over past ½ hour not accounted for by room temperature or patient activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 no report of chills or flushing</td>
</tr>
<tr>
<td>1 report of chills or flushing</td>
</tr>
<tr>
<td>2 flushed or observable moistness on face</td>
</tr>
<tr>
<td>3 beads of sweat on brow or face</td>
</tr>
<tr>
<td>4 sweat streaming off face</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Restlessness Observation during assessment</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 able to sit still</td>
</tr>
<tr>
<td>1 reports difficulty sitting still but is able to do so</td>
</tr>
<tr>
<td>3 frequent shifting or extraneous movements of legs/arms</td>
</tr>
<tr>
<td>5 unable to sit still for more than a few seconds</td>
</tr>
</tbody>
</table>
## Opiate withdrawal scale

### Pupil Size
- 0 pupils pinned or normal size for room light
- 1 pupils possibly 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
*If patient was having pain previously, only the additional component attributed to opiates withdrawal is scored*
- 0 not present
- 1 mild diffuse discomfort
- 2 patient reports severe diffuse aching of joints/muscles
- 4 patient is rubbing joints or muscles and is unable to sit still because of discomfort

### Runny nose or tearing
*Not accounted for by cold symptoms or allergies*
- 0 not present
- 1 nasal stuffiness or unusually moist eyes
- 2 nose running or tearing
- 4 nose constantly running or tears streaming down cheeks

---

Continued
Opiate withdrawal scale

**GI Upset:** over last ½ hr
- 0 no GI symptoms
- 1 stomach cramps
- 2 nausea or loose stool
- 3 vomiting or diarrhoea
- 4 multiple episodes of diarrhoea or vomiting

**Tremor** *observation of outstretched hands*
- 0 no tremor
- 1 tremor can be felt but not observed
- 2 slight tremor observable
- 4 gross tremor or muscle twitching

**Yawning** *Observation during assessment*
- 0 no yawning
- 1 yawning once or twice during assessment
- 2 yawning three or more times during assessment
- 4 yawning several times/minute
Opiate withdrawal scale

Anxiety or Irritability
- 0 none
- 1 patient reports increasing irritability or anxiousness
- 2 patient obviously irritable or 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 ______
The total score is the sum of all 11 items
Initials of persons
Completing assessment ___________________
Methadone: Inappropriate dosing

Dose too low – Withdrawal
- “Flu-like” symptoms
- Runny nose, sneezing
- Abdominal cramps, diarrhoea
- Tremor, muscle spasm, aches, and cramping
- Yawning, “teary” eyes
- Hot and cold sweats
- Irritability, anxiety, aggression
- Aching bones
- Craving

Dose too high – Intoxicated
- Drowsy, “nodding off”
- Nausea, vomiting
- Shallow breathing
- “Pinned” (pinpoint) pupils
- Drop in body temperature
- Slow pulse, low BP, palpitations
- Dizziness
Stabilisation (1)

Rate of Dose Increase

- Increase 0-10mg methadone per 1-3 days during the first week according to physical assessment and SOWS score
- Maximum increase of 20-25mg over 1st week
- Subsequent dose increases should not exceed 10mg per week

Continued
Stabilisation (2)

Rate of Dose Increase

- gradual increase essential due to long half-life
- Best outcomes from maintenance doses > 60mg
- Lethal dose 20mg for children, as low as 50mg for opioid-naïve adults
- Repeated doses of 30mg can be fatal in adults

Notes:
Methadone dosage is individualised. 80 mg is the average, but some patients may require lower or higher doses.
Relationship between Methadone Dose and Heroin Use

(Adapted from Ball and Ross, 1991)
Stabilisation (3)

Frequency of Appointments

- First 5 - 7 days - see every 1-2 days
- Write prescription till next appointment only
- Always see the patient before increasing the dose
- Continue the assessment process, build the therapeutic relationship
Other treatment issues

- Promote compassionate opioid analgesia
  - Health care worker education especially at hospital
  - Role of maintenance treatment in analgesia
- Encourage good vein care
  - To maintain venous access
  - Important later, if applicable, in the clinical course of HIV infection
Ongoing management issues (1)

- Monitoring HIV progression
  - Co-infection
  - Cognitive state
- Mental health
  - Depression
  - Suicide ideation
  - ASPD
  - PTSD
- Pain management
- Drug substitution
Ongoing management issues (2)

- Risk exposure
  - dose
  - compliance with program rules
- Cost of medication
- Staff attitudes
Characteristics of effective programs

- Longer duration (2-4 years)
- Higher doses; > 60mg methadone
- Accessible prescriber and dispenser
- Integrated services
- Quality of therapeutic relationship
Drug interactions - metabolism

- Methadone
  - Metabolism Cytochrome P450 mediated
    - CYP3A4 main
    - also CYP2D6, CYP1A2, CYP2C9 and CYP2C19,
      - genetic variability
  - CYP3A4 breaks down 50% of drugs
    - Methadone mixed inhibitor
      - may increase other drug levels, e.g., Nifidepine, etc.
Questions?

Comments?
Thank you for your time!

End of Workshop 2
Workshop 3: Opiate Addiction Treatment with Buprenorphine
Training objectives

At the end of this training you will:

1. Understand medical withdrawal protocols using buprenorphine
2. Know the basic purpose and background evidence to support the use of buprenorphine for treating opiate dependence
3. Know the basic principles of maintenance treatment with buprenorphine
4. Know effective practices (evaluation, initial dose and management of dose; tapering procedures, etc.) in the implementation of buprenorphine treatment
5. Understand how to address concurrent use of other drugs and alcohol during buprenorphine treatment
6. Know contraindications and medication interactions with buprenorphine
Overview
Overview

- Buprenorphine is a thebaine derivative (classified in the law as a narcotic)
- High potency
- Produces sufficient agonist effects to be detected by the patient
- Available as a parenteral analgesic (typically 0.3 - 0.6 mg im or iv every 6 or more hours)
- Available as sublingual tablets (2-8 mg)
- Long duration of action when used for the treatment of opioid dependence contrasts with its relatively short analgesic effects

Notes

Buprenorphine is a thebaine derivative. This is important, because it leads to buprenorphine’s legal classification as an opioid. It has high potency. Buprenorphine has been available for years in the United States as a parenteral analgesic. Typical analgesic doses are 0.3-0.6 mg i.m. or i.v. every 6 (or more) hours.

Sublingual tablets of buprenorphine with naloxone are also available to reduce the potential for abuse (source: U.S. Department of Health and Human Services. Substance Abuse and Mental Health Services Administration, Center for Substance Abuse Treatment. *Clinical Guidelines for the Use of Buprenorphine in the Treatment of Opioid Addiction: TIP 40*. Available at www.samhsa.gov)
Affinity and dissociation

Buprenorphine has:

- high affinity for mu opioid receptor –
  - competes with other opioids and blocks their effects
- slow dissociation from mu opioid receptor –
  - prolonged therapeutic effect for opioid dependence treatment (contrasts to its relatively short analgesic effects)

Notes

1. Buprenorphine has high affinity for the mu opioid receptor. This means that it is hard for other opioids with lower affinity to displace buprenorphine from the mu receptor (so it blocks their effects).

2. Buprenorphine’s slow dissociation from the mu receptor results in a prolonged therapeutic effect. Considerable evidence suggests buprenorphine can be given three times per week (rather than daily), and there is some evidence suggesting buprenorphine can be given even less frequently (e.g., two times per week).

3. Buprenorphine’s long duration of action when used as a medication for the treatment of opioid dependence contrasts with its relatively short analgesic effects.
Abuse potential

- Buprenorphine is abusable (epidemiological, human laboratory studies show)
- Diversion and illicit use of analgesic form (by injection)
- Relatively low abuse potential compared to other opioids

**Notes:**
Abuse potential of buprenorphine is similar to heroin in laboratory studies, but it is safer in overdose.
Notes:
Avoid referring to buprenorphine as an agonist/antagonist. It is confusing. It is sufficient to use the term partial agonist.
Positive effect does not necessarily correspond to the addictive potential. Buprenorphine is as addictive as methadone, if not more so, but it is safer in overdose.
Buprenorphine: Clinical pharmacology

- Partial agonist
  - high safety profile / ceiling effect
- Tight receptor binding at mu receptor
  - long duration of action
  - slow onset mild abstinence
- Antagonist at k receptor
Subjects’ Rating of Drugs’ Good Effect

![Graph showing subjects' rating of drugs' good effect]

- **Buprenorphine (mg):**
  - Peak Score:
    - 0.5 mg: 3.75
    - 2 mg: 15
    - 8 mg: 60
- **Methadone (mg):**
  - 3.75 mg: 0
  - 15 mg: 50
  - 60 mg: 100
Buprenorphine’s Effect on Respiration

Breaths/minute vs. Buprenorphine (mg)

Effect of Buprenorphine (mg) on Breaths/minute
Intensity of Abstinence Symptoms

Days after drug withdrawal

Himmelsbach scores

Buprenorphine
Morphine
Metabolism and excretion

- High percentage of buprenorphine bound to plasma protein
- Metabolised in liver by cytochrome P450 3A4 enzyme system into norbuprenorphine and other metabolites

References:
Patient selection: Issues for consultation (1)

Several factors may indicate a patient is less likely to be an appropriate candidate, including:

- Patients taking high doses of benzodiazepines, alcohol, or other central nervous system depressants
- Significant psychiatric co-morbidity
- Multiple previous opioid addiction treatment episodes with frequent relapse during those episodes (may also indicate a perfect candidate)
- Nonresponse or poor response to buprenorphine treatment in the past
Currently buprenorphine is not approved for use during pregnancy.

However, studies conducted to date suggest that buprenorphine may be an excellent option for pregnant women.

Randomized trials are underway to determine the safety and effectiveness of using buprenorphine during pregnancy.

Notes:

There is limited clinical experience with buprenorphine maintenance in pregnant women who are addicted to opioids. There is a need for further studies to determine if it is safe to use buprenorphine during pregnancy.
Patients with these conditions must be evaluated by a physician for appropriateness prior to buprenorphine:
- Seizures
- HIV and STDs
- Hepatitis and impaired hepatic function
- Use of alcohol, sedative-hypnotics, and stimulants
- Other drugs

Patient selection:
Issues for consideration (3)
Overview: Goal of induction

To find the dose of buprenorphine at which the patient:

- discontinues or markedly reduces use of other opioids
- experiences no cravings
- has no opioid withdrawal symptoms
- has minimal / no side effects
Buprenorphine induction:
For short-acting opioids (1)

Patients dependent on short-acting opioids (e.g., heroin, oxycodone): Day 1

Instruct patients to abstain from any opioid use for 12-24 hours (so they are in mild withdrawal at time of first buprenorphine dose) – may be easiest to schedule appointment early in day (decrease risk of opioid use prior to office visit)
Buprenorphine induction: For short-acting opioids (2)

Patients dependent on short-acting opioids (continued)

If patient is not in opioid withdrawal at time of arrival in office, then assess time of last use and consider either having them return another day, waiting in the office until evidence of withdrawal is seen, or leaving office and returning later in the day (with strict instructions to not take opioids while away from the office)
Buprenorphine induction:  
For short-acting opioids (3)

Patients dependent on **short-acting opioids (continued)**

- First dose: 2-4 mg sublingual buprenorphine
- Monitor in office for up to 2 hours after first dose
- Relief of opioid withdrawal symptoms should begin within 30-45 minutes after the first dose
Patients dependent on **short-acting opioids**

- If opioid withdrawal appears shortly after the first dose, it suggests that the buprenorphine may have precipitated a withdrawal syndrome.
- Clinical experience suggests the period of greatest severity of buprenorphine-related precipitated withdrawal occurs in the first few hours (1-4) after a dose, with a decreasing (but still present) set of withdrawal symptoms over subsequent hours.
Buprenorphine induction: For short-acting opioids (5)

Patients dependent on short-acting opioids (continued)

- If a patient has precipitated withdrawal consider:
  - giving another dose of buprenorphine, attempting to provide enough agonist effect from buprenorphine to suppress the withdrawal, or stopping the induction, provide symptomatic treatments for the withdrawal symptoms, and have patient return the next day
- Can re-dose if needed (every 2-4 hours, if opioid withdrawal subsides and then reappears)
- Maximum first-day dose of 8/2 mg buprenorphine / naloxone
Combination of buprenorphine plus naloxone

- Combination tablet containing buprenorphine with naloxone – if taken under tongue, predominant buprenorphine effect
- If opioid-dependent person dissolves and injects buprenorphine / naloxone tablet – predominant naloxone effect (and precipitated withdrawal)
Induction: Patient Physically Dependent on Short-acting Opioids, Day 1

Patient dependent on short-acting opioids?
Yes
Withdrawal symptoms present 12-24 hrs after last use of opioids?
Yes
Give buprenorphine/naloxone 4/1 mg, observe
No
Withdrawal symptoms continue or return?
No
Withdrawal symptoms relieved?
Yes
Daily dose established.
No
Repeat dose up to maximum 8/2 mg for first day
Yes
Withdrawal symptoms relieved?
No
Manage withdrawal symptomatically
Yes
Daily dose established.
No
Withdrawal symptoms return?
Daily dose established.
Yes
Return next day for continued induction.
Stop; Reevaluate suitability for induction

Daily dose established.
Buprenorphine induction: For long-acting opioids (1)

Patients dependent on long-acting opioids

- Experience suggests patients should have dose decreases until they are down to <30 mg/d of methadone
- Begin induction at least 24-36 hours after last dose of methadone
- Patient should be in mild withdrawal from methadone
- Give no further methadone once buprenorphine induction is started
Buprenorphine induction: For long-acting opioids (2)

- Use similar procedure as that described for short-acting opioids (i.e., first dose of 4/1 mg of buprenorphine/naloxone)
- Expect total first day dose of 8/2 mg sublingual buprenorphine/naloxone
- Continue adjusting dose by 2-4 mg increments until an initial target dose of 12-24 mg is achieved for the second day
- Continued dose increases are indicated after the second day to a maximum daily dose of 32/8 mg
Induction: Patient Physically Dependent on Long-acting Opioids, Day 1

Patient dependent on long-acting opioids?

If methadone, taper to <40 mg per day

24 hrs after last dose, give buprenorphine 4/1 mg

Withdrawal symptoms present?

Give buprenorphine 4/1 mg

Withdrawal symptoms continue?

Repeat dose up to maximum 12/3 mg/24 hrs

Withdrawal symptoms relieved?

Yes

No

Daily dose established

Yes

No

Manage withdrawal symptomatically

Daily dose established

GO TO INDUCTION FOR PATIENT PHYSICALLY DEPENDENT ON SHORT-ACTING OPIOIDS
Buprenorphine induction:
For short- or long-acting opioids

Patients dependent on short- or long-acting opioids

- After the first day of buprenorphine induction for patients who are dependent on either short-acting or long-acting opioids, the procedures are essentially the same.
- On Day 2, have the patient return, if possible, for assessment and Day 2 dosing.
- Assess if patient has used opioids since their last visit, and adjust dose according to the patient’s experiences after first-day dosing.
Patient returns to office on 8/2-12/3 mg

- Withdrawal symptoms present since last dose?
  - Yes: Increase buprenorphine/naloxone dose to 12/3-16/4 mg
    - Withdrawal symptoms continue?
      - Yes: Administer 4/1 mg doses up to maximum 24/6 mg (total) for second day
    - No: Withdrawal symptoms relieved?
      - Yes: Daily dose established.
      - No: Manage withdrawal symptomatically

- No: Maintain patient on 8/2-12/3 mg per day.

- Withdrawal symptoms return?
  - Yes: Daily dose established.
  - No: Return next day for continued induction; start with day 2 total dose and increase by 2/0.5-4/1 mg increments. Maximum daily dose: 32/8 mg

Notes:
Increase the dose if the patient is continuing using heroin or other illicit opioids.
Buprenorphine stabilisation / maintenance (1)

- The patient should receive a daily dose until stabilised
- Once stabilised, the patient can be shifted to alternate day dosing (e.g., every other day, MWF, or every third day, MTh)
- Increase dose on dosing day by amount not received on other days (e.g., if on 8 mg/d, switch to 16/16/24 mg MWF)
Buprenorphine stabilisation / maintenance (2)

- Stabilise on daily sublingual dose
- Expect average daily dose to be somewhere between 8/2 and 32/8 mg of buprenorphine / naloxone
- Higher daily doses more tolerable if tablets are taken sequentially rather than all at once
Maintenance treatment using buprenorphine

- Buprenorphine more effective than placebo
- Buprenorphine equally effective as methadone. However, methadone has better retention rates and probably less heroin use also.
- More research needed on if buprenorphine can be as effective as higher doses of methadone (e.g., 80-100 mg or more per day), and therefore may not be the treatment of choice for some patients with higher levels of physical dependence.
- Individuals with better levels of psychosocial functioning and support are optimal candidates for buprenorphine.

Notes
1. In general, these studies have shown buprenorphine and methadone are equivalent on primary outcome measures (treatment retention, rates of positive urine samples for illicit opioids). However, methadone has better retention and probably less heroin use also. There is even less heroin use at higher doses of methadone.
Comparison of buprenorphine maintenance vs. withdrawal:

Shows both the efficacy of maintenance treatment, and the poor outcomes associated with withdrawal (even when provided within the context of a relatively rich set of psychosocial treatments including hospitalisation and cognitive behavioral therapy)
Stabilisation / Maintenance

Induction phase completed?

Yes

Continued illicit opioid use?

No

Withdrawal symptoms present?

No

Compulsion to use, cravings present?

No

Daily dose established

Yes

Daily dose established

Continue adjusting dose up to 32/8 mg per day

No

Continued illicit opioid use despite maximum dose?

No

Maintain on buprenorphine/naloxone dose, increase intensity of non-pharmacological treatments, consider if methadone transfer indicated
Withdrawal using buprenorphine (1)

**Withdrawal <=7 days**
- Buprenorphine is effective in suppressing opioid withdrawal symptoms
- Long-term efficacy is not known, and is likely limited
- Studies of other withdrawal modalities have shown that such brief withdrawal periods are unlikely to result in long-term abstinence

**Withdrawal <=7 days**
- Reports show buprenorphine suppresses opioid withdrawal signs and symptoms (better than clonidine)

**Withdrawal <=7 days**
- Using sublingual tablets:
  - First day: 8/2-12/3 mg sl
  - Second day: 8/2-12/3 mg sl
  - Third (last) day: 6/1.5 mg sl
Withdrawal over >30 day (long-term)

- Not a well-studied topic
- Literature on opioid withdrawal can provide guidance; suggests longer, gradual withdrawals more effective than shorter withdrawals

Although there are few studies of buprenorphine for such time periods, buprenorphine has been shown more effective than clonidine over this time period.
Withdrawal using buprenorphine (3)

Regardless of the buprenorphine withdrawal duration:

Consider use of ancillary medications to assist with symptoms of opioid withdrawal (e.g., medications for arthralgias, nausea, insomnia)
Overview of safety and side effects

- Highly safe medication (under both acute and chronic dosing circumstances)
- Also safe if inadvertently swallowed by someone not dependent on opioids (because of poor oral bioavailability and the ceiling on maximal effects)
- Primary side effects: like other mu agonist opioids such as methadone (e.g., nausea, constipation)
- Anecdotal reports indicate that symptoms may be less severe

Notes

1. Buprenorphine is a highly safe medication for use in patients with opioid dependence.
2. Note that it is also safe if inadvertently taken by a person who is not physically dependent on opioids (such as a child). In such a case, it is most likely the person would swallow the tablet and experience virtually no opioid agonist effect because of the poor oral bioavailability. Even if the person sucked on the tablet, there is a low likelihood that they would experience serious adverse effects. This is because buprenorphine is a partial opioid agonist, and there is a ceiling in the maximal effects produced.
3. Clinical trials with buprenorphine have found no significant organ damage associated with chronic dosing. However, buprenorphine may be associated with increases in liver function tests, and this may be especially true for patients with a history of hepatitis prior to the onset of buprenorphine treatment. Increases in liver function tests appear to be mild, and it is important to keep in mind that other factors commonly found in opioid dependent patients (such as hepatitis and alcohol abuse) can lead to elevations in liver function tests. Subsequent slides address the effects of buprenorphine on liver function tests.

References:


Precipitated withdrawal (1)

- The likelihood for buprenorphine-precipitated withdrawal is low for short-acting opioids.

- Buprenorphine-precipitated withdrawal seen in controlled studies has been mild in intensity and of short duration.

Notes
1. The potential for buprenorphine-precipitated withdrawal has been covered elsewhere in the Basic Pharmacology section, and will not be reviewed in detail here.
2. While it is possible for buprenorphine to precipitate withdrawal during buprenorphine induction, and this possibility has received significant attention and review in this curriculum, it is important to keep this potential in perspective. The likelihood for buprenorphine-precipitated withdrawal is low, and even when it does occur, it is mild in intensity and short in duration. The clinician should be aware of the potential, but not allow the potential to deter from the use of buprenorphine.
Risk factors that increase the possibility of buprenorphine-related precipitated withdrawal are:

- higher levels of physical dependence
- a short time interval between last use of an opioid and first dose of buprenorphine
- higher first doses of buprenorphine
Overdose with buprenorphine

- Low risk of clinically significant problems.
- No reports of respiratory depression in clinical trials comparing buprenorphine to methadone.
- Buprenorphine’s ceiling effect means it is less likely to produce clinically significant respiratory depression. However, overdose in which buprenorphine is combined with other CNS depressants may be fatal (reviewed later in this section).

Notes
1. The risk of developing clinically significant problems from a buprenorphine overdose is low. Unlike full agonist opioids (such as methadone and heroin), the maximal opioid agonist effect produced by buprenorphine – a partial agonist – is relatively low. The maximal effects of buprenorphine appear to occur in the 8-16 mg dose range for sublingual solution (in non-dependent opioid abusers). This is equal to 16-32 mg of sublingual tablets. This means that higher doses are unlikely to produce greater effects (and may actually produce fewer effects, based on pre-clinical evidence).
2. This ceiling on the effects produced means buprenorphine is less likely to produce clinically significant respiratory depression. However, overdose in situations where buprenorphine is combined with other CNS depressants may be fatal, as reviewed later in this section.

Reference:
Drug interactions with buprenorphine

1. Benzodiazepines and other sedating drugs
2. Medications metabolised by cytochrome P450 3A4
3. Opioid antagonists
4. Opioid agonists
Benzodiazepines and other sedating drugs (1)

- Reports of deaths when buprenorphine injected along with injected benzodiazepines. Also deaths from non-injection use of both drugs.
  - Reported from France, where buprenorphine without naloxone tablets are available (appears patients dissolve and inject tablets)
  - Probably possible for this to occur with other sedatives
  - Mechanism leading to death in these cases is not known
  - Not clear if any patients have died from use of sublingual buprenorphine combined with oral benzodiazepine. Most deaths appear to have been related to injection of the combination of dissolved buprenorphine tablets with benzodiazepine

Notes

1. It is not clear, based upon the French experience with buprenorphine-related deaths, if any patients have died from use of sublingual buprenorphine combined with oral benzodiazepine. It appears likely that most deaths have been related to injection of the combination of dissolved buprenorphine tablets with benzodiazepine. However, there have been cases of death from non-injection buprenorphine and benzodiazepines.

2. Note that the combination product (buprenorphine with naloxone) is designed to decrease the likelihood that people will dissolve and inject buprenorphine.

3. The mechanism leading to death in these cases is not known.

References:


Benzodiazepines and other sedating drugs (2)

Note that the combination product (buprenorphine with naloxone, Suboxone®) is designed to decrease the likelihood that people will dissolve and inject buprenorphine, so the risk of misuse of buprenorphine with benzodiazepines should be decreased with the availability of buprenorphine / naloxone.
Four possible groups that might attempt to divert and abuse buprenorphine / naloxone parenterally:

1. Persons physically dependent on illicit opioids
2. Persons on prescribed opioids (e.g., methadone)
3. Persons maintained on buprenorphine / naloxone
4. Persons abusing, but not physically dependent on opioids

Notes
1. Note that there are four possible groups that might attempt to abuse buprenorphine/naloxone.
2. For persons physically dependent on an illicit agonist opioid (like heroin), injection of buprenorphine/naloxone will precipitate withdrawal (or, if the dose is very low – e.g., 1/0.25 mg – it will produce placebo-like effects).
3. For persons physically dependent on a prescribed opioid (like methadone or LAAM), injection of buprenorphine/naloxone will precipitate withdrawal (or, again, if the dose is very low, it will produce placebo like effects).
4. For persons maintained on sublingual buprenorphine/naloxone, injection of buprenorphine/naloxone could produce opioid-agonist-like effects (with no precipitated withdrawal from the naloxone, since high doses of naloxone are needed to precipitate withdrawal in buprenorphine-maintained persons). Note that this is a population that will have access and may be very likely to dissolve and inject buprenorphine/naloxone tablets, since they will have a ready supply of them.
5. For persons not physically dependent on opioids, naloxone will not precipitate withdrawal and it is likely the buprenorphine will produce opioid agonist effects.
Notes
This slide shows results from a study in which persons with a history of opioid abuse, but who were not actively dependent upon opioids, received different doses of sublingual buprenorphine solution. The y-axis shows the percentage of identifications of the buprenorphine as placebo, opiate, or something else. As can be seen, as the dose of sublingual buprenorphine increases, the percent of identifications as opiate-like increases (and the proportion of identifications as something else -- placebo or other, decreases). This illustrates buprenorphine’s identification as an opioid-agonist-like drug by persons with a history of opioid abuse.
Maintenance treatment using buprenorphine

Following slides briefly review representative studies:

- Comparison of different doses of sublingual buprenorphine
- Buprenorphine-methadone flexible dose comparison
- Buprenorphine, methadone, LAAM comparison

Notes
The next several slides review four of the studies that have examined the efficacy of buprenorphine under different experimental conditions.
Different Doses of Buprenorphine: Opiate Use

(Ling et al., 1998)

Notes

• Dose effects were seen across a number of outcome measures -- for example, the 1 mg group had significantly poorer treatment retention for the 16 weeks (40%) compared to the 8 mg group (52%) and the 16 mg group (61%).

• Similarly, a significantly lower percentage of patients in the 1 mg group achieved 13 consecutive opioid-negative urine samples (18.5%) compared to the 8 mg group (32.9%).
Notes

Treatment retention was significantly better for the LAAM, buprenorphine, and high-dose methadone groups, compared to the low-dose methadone group. (It was also significantly better for the high-dose methadone group, compared to the LAAM group as well.)

Note that the rescue procedure started during week 6 of treatment, and the sharp drop off in treatment retention for the low-dose methadone group represents, in part, the substantial number of participants in this group who were switched to high-dose methadone.
Notes

This figure shows treatment retention, which was significantly better for the maintenance (buprenorphine) vs. control (withdrawal followed by placebo) group. All placebo patients who dropped out did so following relapse to drug use (as determined by urine testing). In the maintenance group, one patient dropped out of treatment, and four were discharged due to relapse in their drug use.

Urine results showed that 74.8% of samples were negative for drugs in the buprenorphine maintenance group over the course of the year.
Buprenorphine Maintenance / Withdrawal: Mortality

<table>
<thead>
<tr>
<th></th>
<th>Detox/Placebo</th>
<th>Buprenorphine</th>
<th>Cox regression</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dead</td>
<td>4/20 (20%)</td>
<td>0/20 (0%)</td>
<td>$\chi^2=5.9; p=0.015$</td>
</tr>
</tbody>
</table>

(Kakko et al., 2003)

Notes

While not the primary goal of the study, the study noted that four of the patients who underwent a withdrawal (which was inpatient, and lasted six days) had died after one year -- compared to none of the patients in the buprenorphine maintenance group.
Questions?

Comments?
Thank you for your time!

End of Workshop 3
Workshop 4: Opiate Antagonist Treatment: Naloxone for Overdose, Naltrexone for Relapse Prevention
At the end of this training you will:

1. Understand the neurobiology-conditioning underpinning opiate relapse
2. Understand the rationale for the use of naloxone for opiate overdose
3. Know the protocol for the use of naltrexone for relapse prevention
4. Understand the challenges and limitations of naltrexone treatment
Naloxone for Opiate Overdose
Naloxone for opiate overdose

- Naloxone is a medication used to counter the effects of opioid overdose, for example heroin and morphine overdose.

- Specifically, naloxone is used in opioid overdoses for countering life-threatening depression of the central nervous system and respiratory system.

- It is marketed under trade names including Narcan, Nalone, and Narcanti.
Naloxone for opiate overdose

- The drug is derived from thebaine and has an extremely high affinity for \( \mu \)-opioid receptors in the central nervous system.
- Naloxone is a \( \mu \)-opioid receptor competitive antagonist, and its rapid blockade of those receptors often produces rapid onset of withdrawal symptoms.
Naloxone for opiate overdose

- Naloxone is injected, usually initially intravenously for fastest action
  
The drug acts after about two minutes, and its effects may last about 45 minutes.
Signs of opioid overdose

- Unconscious (does not respond verbally or by opening eyes when spoken to loudly and shaken gently)
- Constricted pupils
- Hypoventilation (respiration rate too slow or tidal volume too low)
- Cool moist skin
Opioid overdose: Steps to take (1)

If an opioid overdose is suspected:

- Oxygen, if available
- Naloxone – 0.4-0.8mg IV/IMI, (aliquots of 50mcg every 1-2 minutes may be used IV until arousal sufficient for airway maintenance and adequate ventilation). Dose may be repeated after 2 minutes if no response, to a maximum of 10mg
- Call ambulance
- Advise reception of emergency and location
Opioid overdose: Steps to take (2)

Assess the client:

If responsive

- Airway – open and clear
- Breathing – respiratory rate and volume
- Circulation – carotid pulse
If unresponsive, respiratory arrest, or hypoventilating

- **Call ambulance**
- Place in lateral coma position if breathing spontaneously
- Bag and mask, ventilate with oxygen for hypoventilation
- Naloxone 0.4-0.8mg IV (50mcg aliquots every 1-2 minutes) or IM if suspect opioid OD
Opioid overdose: Steps to take (4)

- If response is adequate
  - The patient will be fully conscious, oriented, alert, and responsive
- If response is inadequate or there is no response to naloxone
  - Continue oxygenation
  - Keep lateral
  - Monitor observations
  - Administer further naloxone
Opioid overdose: Steps to take (5)

- Advise client to go to the hospital for observation + naloxone infusion
- If refuses, advise no further drugs or alcohol that day
- Stay with a responsible person for > 2 hours
- Provide written information regarding above
- If client at risk (suicide / effects of drugs) consider detention order
Naloxone for opiate overdose

Naloxone has been distributed as part of emergency kits to heroin users, and this has been shown to reduce rates of fatal overdose. Projects of this type are underway in San Francisco and Chicago, and pilot projects started in Scotland in 2006.
Naltrexone for Relapse Prevention
Naltrexone for opiate relapse prevention (1)

- Naltrexone is an opioid antagonist treatment medication: It is a pure, potent mu antagonist that can be taken by mouth once daily or every other day, and has minimal side effects.
- It is neither reinforcing nor addicting and has no potential for abuse or diversion for unprescribed use.
Naltrexone, and its active metabolite 6-β-naltrexol, are competitive antagonists at μ- and κ-opioid receptors, and to a lesser extent at δ-opioid receptors.

This blockade of opioid receptors is the basis behind its action in the management of opioid dependence – it reversibly blocks or attenuates the effects of opioids.
Naltrexone for opiate relapse prevention (3)

- Naltrexone is not a narcotic
- It works by blocking the effects of narcotics, especially the “high” feeling that is produced by opiates
- It also may block the “high” feeling that is produced by alcohol
- It will not produce any narcotic-like effects or cause mental or physical dependence
Naltrexone for opiate relapse prevention (4)

- Naltrexone will cause withdrawal symptoms in people who are physically dependent on narcotics.
- Naltrexone treatment is started after an individual is no longer dependent on narcotics.
- It is important for an individual to be fully withdrawn from opiates.
- If naltrexone is taken by individuals who are incompletely detoxified from opiates, it can precipitate a rapid and unpleasant withdrawal syndrome.
The length of time between the last dose of opiate and the first dose of naltrexone is important.

The specific timetable depends on whether the opiate being used was a short-acting opiate (e.g., morphine or heroin) or a long-acting opiate (e.g., methadone) and how long the opiate was used (i.e., days, weeks months).

Before starting naltrexone it is important for the treating physician to have this information.
When opiate-dependent individuals desire to be inducted onto naltrexone, it is necessary to first detoxify them from opiates to avoid precipitated withdrawal.

It is not possible to use the two most effective withdrawal agents, methadone and buprenorphine, because of their agonist properties.

Therefore, detoxification methods that do not employ methadone and/or buprenorphine must be used.
Two commonly used agents are lofexidine and clonidine, both α-adrenergic agonists that relieve most opioid withdrawal symptoms without producing opioid intoxication or drug reward.

Opiate detoxification with these agents is less effective, since they do not relieve many opioid withdrawal symptoms. Therefore, adjunctive medicines often are necessary to treat insomnia, muscle pain, bone pain, and headache.
### Pre-naltrexone detoxification procedures (1)

- An appropriate protocol for clonidine is 0.1mg administered orally as a test dose.
- A dose of 0.2mg might be used initially for patients with severe signs of opioid withdrawal or for those patients weighing more than 200 pounds.
- The sublingual (under the tongue) route of administration also may be used.
- A similar procedure using lofexidine is appropriate; lofexidine produces significantly less hypotension than clonidine.

### Notes:

In some countries, clonidine is not longer registered (Europe). Buprenorphine can be successfully used for withdrawal management (gradually tapered) and then naltrexone started after 3-5 days for maintenance. This withdrawal procedure might be much more convenient than the use of clonidine, which has a significant effect on blood pressure.
Pre-naltrexone detoxification procedures (2)

- Clinicians should check the patient's blood pressure prior to clonidine administration, and clonidine should be withheld if systolic blood pressure is lower than 90 or diastolic blood pressure is below 60.

- These parameters can be relaxed to 80/50 in some cases if the patient continues to complain of withdrawal and is not experiencing symptoms of orthostatic hypotension (a sudden drop in blood pressure caused by standing).
Pre-naltrexone detoxification procedures (3)

- Clonidine (0.1 to 0.2mg orally) can then be given every 4 to 6 hours on an as-needed basis.
- Clonidine detoxification is best conducted in an inpatient setting, as vital signs and side effects can be monitored more closely in this environment.
- In cases of severe withdrawal, a standing dose (given at regular intervals rather than purely "as needed") of clonidine might be advantageous.
The daily clonidine requirement is established by tabulating the total amount administered in the first 24 hours, and dividing this into a three or four times per day dosing schedule.

Total clonidine should not exceed 1.2mg the first 24 hours and 2.0mg after that, with doses being held in accordance with parameters noted above.

The standing dose is then weaned over several days.

Clonidine must be tapered to avoid rebound hypertension.
Naltrexone for opiate relapse prevention

- For oral dosage form (tablets):
- For treating narcotic addiction:
  - Adults—25 milligrams (mg) (one-half tablet) for the first dose, then another 25 mg one hour later. After that, the dose is 350 mg a week. This weekly dose should be divided up according to one of the following schedules:
    - 50 mg (one tablet) every day; or
    - 50 mg a day during the week and 100 mg (two tablets) on Saturday; or
    - 100 mg every other day; or
    - 100 mg on Mondays and Wednesdays, and 150 mg (three tablets) on Fridays; or
    - 150 mg every three days
## Naltrexone for opiate relapse prevention (1)

<table>
<thead>
<tr>
<th>Side effects</th>
<th>Precautions</th>
</tr>
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<tbody>
<tr>
<td>• Acute opioid withdrawal precipitated (e.g., lethargy, aches, cramps, low energy)</td>
<td>• If naltrexone ceased and opioid use reinstated, reduced tolerance to opioids increases risk of overdose and death</td>
</tr>
<tr>
<td>• Depression, irritability</td>
<td>• Precipitates withdrawals in opioid-dependent patients</td>
</tr>
<tr>
<td>• Anxiety, nervousness</td>
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<tr>
<td>• Sleeping difficulties</td>
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<tr>
<td>• Skin rash</td>
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<tr>
<td>• Poor appetite</td>
<td></td>
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<tr>
<td>• Dizziness</td>
<td></td>
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<tr>
<td>• Nausea</td>
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</table>

## Notes

Depression induced by naltrexone responds rapidly with the use of SSRIs. Effects last ≤72 hours. During this period there is a gradual reduction in tolerance.

Patient non-compliance in part due to the absence of any agonist effects is a common problem. Therefore, a favourable treatment outcome requires a positive therapeutic relationship, careful monitoring of medication compliance, and effective behavioural interventions.

Effectiveness tends to be dependent on:
- situation, circumstances, support, commitment of patient
- inclusion as part of comprehensive treatment program (including counselling)
- Long-term treatment efficacy still under investigation
- While effective for some, inappropriate for others
Naltrexone - psychotherapy research

- Positive results when naltrexone is combined with cognitive behavioural therapy and treatment with the Matrix Model
- Contingency management also produces large increases in retention on naltrexone
- Family therapy also promotes successful treatment with naltrexone
- Using legal pressure (individuals sentenced to treatment by courts) to mandate people to take naltrexone can greatly increase retention on naltrexone and outcome success

Notes:
It is also important to mention the value of having a caring friend or significant other as a protective factor to increase retention.
Naltrexone for opiate relapse prevention

- Naltrexone can also be administered as a low-dose implant. These implants can remain effective for 30-60 days. They dissolve slowly and are usually put in under a local anaesthetic in the left iliac fossa.
- This implant procedure has not been shown scientifically to be successful in "curing" the patient of their addiction, although it does provide a better solution than oral naltrexone for medication compliance reasons.

Notes:
Implants have not been registered or tested to date.
Conclusion: Naltrexone for opiate addiction (1)

- Naltrexone, nonselective opioid antagonist
- Induction issues
- Retention
- Depot preparation
- Better outcomes with specific therapies or legal interventions
Conclusion: Naltrexone for opiate addiction (2)

- Treatment with opiate agonists (methadone) or partial agonists (buprenorphine) produces far better retention than does naltrexone.
- Use of these medications has gained far more acceptance by practitioners than has naltrexone treatment.
- Psychotherapy can substantially improve outcome with these medications as well.
Questions?

Comments?
Post-assessment

Please respond to the post-assessment questions in your workbook.

(Your responses are strictly confidential.)

10 minutes
Thank you for your time!

End of Workshop 4