

Injectable and implantable sustained release naltrexone in the treatment of opioid  
addiction

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## **Summary**

### *Background*

Sustained release technologies for administering the opioid antagonist naltrexone (SRX) have the potential to assist opioid-addicted patients in their efforts to maintain abstinence from heroin and other opioid agonists. Recently, reliable SRX formulations in intramuscular or implantable polymers that release naltrexone for 1-7 months have become available for clinical use and - research.

### *Methods*

This qualitative review of the literature provides an overview of the technologies currently available for sustained release naltrexone (SRX) and their effectiveness in reducing opioid use and other relevant outcomes.

### *Results*

The majority of studies indicate that SRX is effective in reducing heroin use, and the most frequently studied SRX formulations have acceptable adverse events profiles. Registry data indicate a protective effect of SRX on mortality and morbidity. In some studies, SRX also seems to affect other outcomes like concomitant substance use, vocational training attendance, needle use, and risk behaviour for blood-borne diseases like Hepatitis or HIV. There is a general need for more controlled studies, in particular comparing SRX with agonist maintenance treatment, combinations of SRX with behavioural interventions, and with at-risk groups like prison inmates or opioid addicted pregnant patients.

### *Conclusion*

The literature suggests that sustained release naltrexone is a feasible, safe and effective option for assisting abstinence efforts in opioid addiction.

## Introduction

Heroin is used by an estimated 0.4% of the world's population, but heroin-related problems account for nearly 60% of the treatment demand in Europe and Asia (1). The best candidate explanation for this lies in the comprehensive nature of heroin addiction: the sedative effects of the opioid agonist heroin greatly increases the risk of fatal or near-fatal overdose, while a high incidence of injecting use greatly increases the risk of introducing bacterial, viral or fungal agents due to non-sterile injecting practices. Regular heroin users also have an increased occurrence of mental health disorders, and often engage in the regular use of at least two other illicit drugs (2). In the United States of America, diversion and misuse of prescription opioids is an increasing problem (3). Environmental factors associated with illicit opioid use, such as engagement in criminal activities, poor living standards and 'less stable environments' (i.e. exposure to violence, accidents, injury and suicide) (4). All these factors contribute to increase the risk of death from regular illicit opioids to a rate of about 8.6 deaths per 1000 person-years (5). This risk is heightened following detoxification and discharge from a controlled environment, as opioid receptors are thought to readjust to function without exogenous opioid intake. For example, one study found risk of overdose death was 12 times that of the pre-admission risk following discharge from inpatient treatment like detoxification (6). Another study found mortality risk was up to 34 times elevated during the first two weeks following release from a prison setting (7). Recovery from heroin addiction often takes several years with at least occasional relapse and setbacks; it is thus often understood as a chronically relapsing disease (8). While most of our present knowledge on opioid addiction comes from experience with illicit heroin users, all types of opioid agonists

share the same basic neurophysiological pathways and thus the risk of dependence, tolerance, withdrawal, intoxication and abuse.

### **Present treatment alternatives**

Until recently, treatment options for heroin addiction were limited to three main alternatives: *Detoxification* followed by long-term residential treatment; *Opioid maintenance treatment* (OMT) and *Oral Naltrexone*.

*Detoxification* followed by long-term residential treatment has been found to result in some reduction in drug use for a large minority of patients, but suffers from problems with retention in treatment and risk of overdose upon discharge (9). *Opioid maintenance treatment* maintains or substitutes dependence on heroin via the supervised administration of opioid agonist medications including methadone, buprenorphine or medically dispensed heroin (10). While OMT is effective in reducing mortality, morbidity and drug-related criminal activity, chief concerns are dropout during the initial months of treatment and that only a minority of patients are able to achieve normal vocational and social functioning. For those who do achieve such integration, there is currently no validated alternative to life-long dependence on the opioid agonists administered daily in OMT.

### **Naltrexone - an opioid antagonist**

Naltrexone induces a competitive antagonism at all main types of opioid receptors, with some preference for the mu receptor. Although both naltrexone and naloxone were developed based on modifications of oximorphan, naltrexone's overall affinity for opioid receptors is higher and its half-life significantly longer than that of naloxone. Thus naloxone is better suited for acute purposes like reversing the effects

of opioid-induced sedation, while naltrexone is better for scenarios that require prolonged antagonism, e.g. assisting abstinence from opioid agonists following detoxification and/or reducing addiction-related craving. While a full review of these latter types of effects is beyond the scope of this article, the high prevalence of comorbid substance use problems makes them relevant to the overall therapeutic effect, especially for heroin users.

Naltrexone has long been known to cause a reduction in craving sensation for many types of addictive substances including alcohol (11) and amphetamine (12). There has also been reports of a similar effect on certain types of compulsive behaviours, such as bodily self-harm (13) and gambling addiction (14). The precise mechanism for craving reduction has not been determined, but the most likely is that naltrexone causes antagonism of opioid pathways to the nucleus accumbens, reducing the total amount of dopamine released. Naltrexone at very low doses (0.25 mg/day) seems to reduce the severity and/or longevity of opioid withdrawal during detoxification (15), possibly assisting a restoration of normal opioid receptor functioning (16) and attenuating noradrenergic withdrawal systems (17). In addition, opioid antagonists like naltrexone affects other biological systems like G-receptor second messenger systems (18), the immune system (19), and the HPA axis (20).

### **Compliance problems with oral naltrexone**

Studies of oral naltrexone tablets taken daily or bi-daily have generally failed to show superiority over placebo, mostly due to rapid dropout in the active naltrexone group. However, modestly improved results can be achieved when oral naltrexone is taken as part of a compliance-reinforcing scheme like contingency management (21). The lack

of clinical success with oral naltrexone were recognized in the first clinical studies of oral naltrexone (22,23). Consequently, research efforts were started in order to develop sustained release technologies that would decrease compliance problems by reducing the number of dropout opportunities. As part of development efforts for a sustained release formulation, two central SRX characteristics were formulated:

- 1) for blocking street heroin doses, the minimum plasma level of naltrexone was estimated to be about 1 ng/ml, although some of this blockade is also provided by the metabolite 6-beta naltrexol (24). And 2) A clinically useful SRX formulation was thus considered to release naltrexone at levels of 1ng/ml plasma or above for the duration of at least four weeks, with an acceptable rate of tissue-related adverse events. Following more than 30 years of development efforts, this goal has recently been achieved.

### **Sustained release naltrexone (SRX) formulations**

Currently two main types of sustained release technologies are used to release naltrexone: injectable intramuscular suspension and surgically implantable pellets. This section provides a summary of the data from the literature on the currently available SRX technologies, and their ability to block opioid agonists such as heroin or morphine. While there are other sustained release technologies available e.g. for buprenorphine (25), these have not been developed for naltrexone.

#### *Poly lactide suspension*

The naltrexone release of this class of SRX medications is based on the slow biodegrading of a 380 mg poly-lactide and naltrexone suspension providing therapeutic blood levels of naltrexone over a period of 28 days. An intramuscular

SRX suspension of this type was recently FDA-approved for prescription for opioid dependence in the US, after being approved for the treatment of alcohol dependence in 2006. The intramuscular suspension is administered via injection into the gluteus muscle, alternating sides every 4 weeks. A research-only formulation can be injected subcutaneously. With the latter formulation, a heroin challenge study was conducted where participants were administered a 380 mg dosage of subcutaneous and then received IV dosages of heroin at 0, 6.25, 12.5 or 25 mg of heroin in a double-blind design. The suspension provided satisfactory blockade of both self-rated and objective measures (e.g pupil diameter) of heroin for between four and five weeks (26). Recently, a similar experiment was conducted using the FDA-approved intramuscular suspension in reduced dosages of 75, 150 or 300 mg of naltrexone and using hydromorphone instead of heroin for the challenge tests; 3 mg of hydromorphone was blocked by the 300 mg SRX formulation for 28 days, whereas the lower SRX dosages blocked this challenge for a correspondingly shorter duration (27).

#### *Surgically implanted capsules*

The other main type of SRX technology consists of pellets with biodegradable solid polymer surgically inserted or implanted under the skin or fatty tissue with the use of local anaesthetic. The wound is then sealed with 1-3 sutures, with the wound inspected after about one week. The two formulations of surgically implanted naltrexone that have been used in the majority of controlled studies are an Australian type with release periods as long as 7 months when 30 pellets are inserted (28) and a Russian type with a release period of 2-3 months (29). Other manufacturers of naltrexone implants exist, but little research has been published on their reliability or production methods (see (30) for an exception to this). Case data support the view that

SRX implants releasing naltrexone at or above 1ng/ml blood will block normal dosages of laboratory-administered heroin as well as high dosages of illicit heroin (24,31,32).

### **Effect on opioid use**

The majority of RCTs on SRX have shown promising increases in heroin abstinence in the SRX group relative to controls, despite diversity in sample composition, study design, and cultural settings. Two studies have been conducted of 4-week intramuscular SRX suspensions: An eight-week double-blind study from the US of a selected sample divided into a high-dosage to low-dosage and placebo (33), and a 24-week double-blind trial of SRX vs placebo in a sample of Russian heroin users (34). Both studies found significant increases in the proportion of urine samples negative for heroin use. On implantable naltrexone, five RCTs will be reviewed here: Three RCTs utilized a six-month version of the Australian implant: One open-label study randomizing to treatment as usual in a Norwegian treatment setting (35) and a placebo-controlled, double-dummy design with oral naltrexone in Western Australia (36) both found significant decreases in heroin use. A Norwegian open-label study randomizing to methadone OMT or naltrexone implant in probationer settings experienced dropout problems, and found similar reductions in opioid use among the patients who remained (37).

Two randomized studies have been conducted in Russia using a Russian naltrexone implant: A 10-week study of n=100 patients (n=50 in the SRX and placebo groups, respectively) who were both amphetamine and heroin dependent found significant reductions in heroin use (29). A larger study that followed n=306 opioid dependent

patients over 6 months in a three-group, double-dummy design found a significantly larger proportion of urine samples were opioid-negative in the active SRX group compared to both oral naltrexone and placebo (38).

The magnitude of the reduction in opioid use with SRX is typically about 50% at a group level when compared to oral naltrexone or usual-treatment controls, although there is considerable individual variation among patients. In summary, sustained release naltrexone seems to succeed in assisting patients in achieving abstinence from opioids. The consistency of this finding despite diversity in study designs, cultural setting, and SRX formulation reinforces the impression that SRXs' effect on heroin use is a clinically robust finding. There are few data regarding the effectiveness of SRX in the treatment of addiction to prescription opioids.

#### **SRX and heroin-related overdose**

Naltrexone's ability to compete against heroin for opioid receptors means it should provide protection against overdose and – death. The RCTs thus far completed have an insufficient number of participants to permit meaningful analyses of mortality rates. A series of registry cohort studies from Western Australia have used samples of several thousand patients; these studies suggest SRX reduces the number of deaths among heroin users compared to methadone users and oral naltrexone (39–41). The same open cohort was used for the SRX implant patients in two of these studies. Case reports have been published of patients 'breaking the naltrexone blockade' with large doses of opioids (e.g. (42)), as well as post-mortem cases (43) often do not account for potential confounding factors. Data from Norwegian SRX patients confirm that a minority of patients report 'breakthrough'-like experiences, but that the use of non-

opioid illicit drugs makes it difficult to verify which substance induced the experience (32). The concept of true receptor agonism or 'breakthrough' in the presence of naltrexone also appear inconsistent with case stories of naltrexone blocking large quantities of heroin (24,32).

An extension of this question is whether death from an overdose of heroin can occur in active SRX patients. Like any pharmacotherapy, naltrexone's binding at the receptor site is of a competitive type that it is technically possible to outperform using extreme quantities of normal-affinity opioids or high-affinity synthetic opioids like fentanyl. In clinical settings, obtaining and self-administering agonists of the right type or quantity would be very difficult; deaths in patients treated with a reliable SRX formulation are thus more likely to be caused by exposure to the many non-opioid mortality sources common in the heroin demographic.

#### **Retention in SRX for heroin users**

Ambivalence between remaining in treatment and recommencing heroin use means heroin users are often tempted to drop out from treatment. Thus retention in treatment is considered a highly important measure of the clinical feasibility of any treatment for heroin addiction, including OMT and SRX. For naltrexone treatment, the inability to retain patients in oral naltrexone regimens has strongly contributed to why oral naltrexone treatment has seen minimal adoption in clinical settings with heroin users (21). A central clinical advantage of sustained release – over oral naltrexone – is the reduction in dropout opportunities, e.g. one intramuscular injection every 28<sup>th</sup> day instead of a tablet every day. In one RCT (33), retention was 62% between the first and second 28-day intramuscular SRX administration. In the Russian study of

intramuscular SRX (28 days' naltrexone release), attrition at the end of six months' administration of intramuscular SRX administrations was about 50% (34). This is similar to retention between the first and second administration of six-month implantable SRX (44). For patients receiving the 10-week Russian implant, retention was 63% over 6 months among Russian heroin users (38) and 52% in the study of patients with both opioid and amphetamine dependence (29). Differences in study design and - setting, as well as differences in readministration frequencies and adverse event profile make it difficult to infer beyond that retention rates for SRX are within a clinically acceptable range and tend to be better than their comparison group. Thus in this respect SRX seems to confirm hopes that it would constitute an improvement over oral naltrexone (21).

#### **Integration with other behavioural interventions**

A study from the Johns Hopkins behavioural laboratory found that when entry into a voucher-based workplace system was contingent on acceptance of a monthly intramuscular SRX, compliance and retention was improved when patients could enter the workplace freely versus those who were simply prescribed SRX monthly: 74% of contingency patients accepted all six injections, whereas only 26% of prescription patients did the same (45). This is consistent with previous findings from contingency management with oral naltrexone (46). This suggests that the retention in SRX can be greatly improved when combined with behavioural interventions in order to maximise its clinical usefulness.

SRX administered as part of a planned release from prison is another area of considerable interest, in particular due to the increase in overdose mortality reported

in several studies (e.g. (7,47)). As heroin is less available in prison, inmates are more likely to maintain abstinence from heroin that greatly facilitates naltrexone induction (48). Several studies on oral naltrexone for opioid dependent inmate populations concluded with beneficial outcomes when naltrexone was integrated with psychosocial support to enhance external motivation, e.g. work-release programmes and parole including follow-up by criminal justice staff (49–52). Although treatment attrition was still high in these trials, those who stayed on oral naltrexone were less likely to relapse to heroin and less likely to engage in criminal activity than comparison groups not receiving naltrexone. A recent pilot study suggests intramuscular SRX is feasible in probationers with participants displaying reductions in opioid use (53). This is consistent with findings from a Norwegian OMT-SRX randomized study (37), where heroin abstinence rates were equivalent between the two groups six months post release. There is debate regarding the ethical aspects of mandating SRX for heroin users as part of sentencing or parole conditions (e.g. (54)).

#### **Concomitant substance use**

Several studies have examined whether SRX also reduces concomitant use of non-opioid illicit drugs. Naltrexone has been known to reduce craving for a number of addictive substances (see elsewhere in this issue), often resulting in a subsequent reduction in substance use. Of the available studies, RCTs with stricter inclusion criteria seem to confirm a change in non-opioid drug use (33,34); this effect does not reach significance in studies with less strict inclusion criteria (28,34,35). This indicates that SRX may have an effect on concomitant drug use in heroin users, but less dramatic than the effect seen on heroin consumption; the division along inclusion criteria may also indicate that a reduction in concomitant substance use is more likely

to occur in subgroups of heroin users that are pre-screened to reduce the incidence of potential confounders.

### **Somatic & mental health outcomes**

A registry cohort study in Australia followed cohorts of both SRX and methadone patients, and found their rate of mental health related hospitalization similarly reduced (55). In a similar study, SRX patients presented with fewer psychiatric hospital admissions after entering SRX (56). For somatic hospitalizations, overdose admissions were reduced to zero among SRX implant patients in a registry linkage study, and continued to be reduced compared to pre-admission levels for an additional six months following the expiry of naltrexone from the SRX implant (39).

### **Adverse events**

Moderate adverse effects such as nausea, vomiting, and muscle twitches are experienced by heroin users in both SRX and oral naltrexone treatment (22,57). The majority of adverse effects are described as mild to moderate (35), and are more likely to occur in active SRX groups than in placebo patients (29,33,34). As SRX releases naltrexone into the bloodstream gradually at concentrations typically in the 1-5 ng/ml range, the intensity of adverse effects is much reduced compared to oral naltrexone, where blood naltrexone levels can remain at 10-30 ng/ml for several hours every day following tablet intake. The blockade of endogenous opioids thought to result from treatment with SRX has not been reported to have consequences for the occurrence of mood disorders in any of the RCTs thus far published, even though the majority of them administered instruments to measure depression. While there have been reports of depression in users of oral naltrexone (58,59), subsequent investigations failed to

confirm any effects on mood (60,61). Clinicians should perhaps be more concerned that naltrexone blocks the effects of opioid-agonist based analgesics in an accident-prone population, although increasing the dosage or using other types of analgesics will often resolve the problem. It has also been suggested that naltrexone increases the sensitivity of the opioid receptor system, making patients more vulnerable than usual to heroin overdose once SRX is concluded (62). However, findings from toxicological examinations of heroin-related deaths comparing patients with or without prior naltrexone exposure do not support this hypothesis (63). In addition, a recent database study found a reduction in deaths among SRX patients during the first months following treatment when compared to oral naltrexone patients (41).

An important difference between SRX and oral naltrexone is the occurrence of site-related adverse events (64). For implantable SRX, these may appear as mild allergic itching or redness around the implantation site, infection of the skin, stitching or underlying tissue (65). These events are reported to occur in 2-5% of patients (e.g. (29,35)) and usually resolve with symptomatic treatment but in extreme cases may require removal of the implant. Some patients have cosmetic concerns with the fact that some implantable SRX formulations may take months or years to biodegrade completely (66). Similarly, recipients of SRX with intramuscular suspension can often experience some site pain, while a few percent experience more serious site reactions like induration and infection.

Hepatic health is sometimes a concern with heroin users, especially for patients recently infected with Hepatitis C. There is little evidence that SRX in ordinarily administered doses is hepatotoxic. Intramuscular SRX has been found to be well tolerated in alcohol dependent patients with hepatic impairment requiring no dose

adjustments (67,68). A pilot study of implantable SRX in heroin users found key hepatic indicators such as ALT to improve over the course of treatment (31), and the influence of SRX on indicators in other studies have generally been below levels of clinical significance. A clinical study of 50 SRX implant patients undergoing antiviral therapy for Hepatitis C found 62% were HCV negative following completion of HCV treatment and 6 months of SRX (69). Still, caution may be warranted in administering SRX to patients who present with severely reduced hepatic functioning, e.g. who qualify for an impairment classification corresponding to Child-Pugh grade C.

Pregnancy is a debated topic in SRX research, as with heroin users in general (70–72). SRX medication is now available for regular prescription in the US, and there is an interest among pregnant drug users despite a general lack of knowledge about SRX's effects on foetal health. While this lack of knowledge is unfortunate from a medical point of view, the risk of return to heroin use upon discontinuation of SRX may be considered an even worse outcome. Historically, the solution most often adopted has been to continue the pharmacotherapies for pregnant heroin users and initiating short- and long-term studies on adverse effects following delivery of the child (73). Only one case has been reported following this approach, with no adverse effects detected in mother or child (74).

### **Conclusions**

Since a Cochrane review in 2008 (75) concluded there were too few studies to conduct any meaningful assessment of sustained release naltrexone (SRX) in the opioid addicted, the amount of research published on SRX has accumulated to the point where this conclusion seems gradually less valid. SRX is showing promising,

consistent effects in supporting opioid users' efforts to achieve abstinence across different clinical study design and - treatment settings. The SRX formulations that have been the subject of the majority of research articles appear to have a satisfactory rate of consistency in naltrexone release and an acceptable adverse effects profile. The literature on SRX for opioid addiction still requires more studies in order to confirm initial findings on effects. There is a particular need for more knowledge on SRX compared with current standard treatments, the impact on poly-drug dependence, the use of SRX during pregnancy, and the combination of SRX with other interventions in order to maximise the impact on recovery.

**Declaration of interests**

The authors have completed the Unified Competing Interest form at [www.icmje.org/coi\\_disclosure.pdf](http://www.icmje.org/coi_disclosure.pdf) (available on request) and declare: no support from any organisation for the submitted work; GKH had entered into a contractual arrangement (via the University of Western Australia) with Go Medical Industries (who manufactures the Australian naltrexone implant) to conduct a number of research studies in the previous 3 years; GKH had co-authored with Dr. George O'Neil (Director, Go Medical Industries) on a number of previous publications.

## References

1. United Nations Office on Drugs and Crime. World Drug Report 2011. New York; 2011.
2. Ross J, Teesson M, Darke S, Lynskey M, Ali R, Ritter A, Cooke R. The characteristics of heroin users entering treatment: findings from the Australian treatment outcome study (ATOS). *Drug Alcohol Rev.* 2005/11/22 ed. 2005;24(5):411–8.
3. CDC grand rounds: prescription drug overdoses - a U.S. epidemic. *MMWR. Morbidity and mortality weekly report.* 2012 Jan 13;61(1):10–3.
4. Warner-Smith M, Darke S, Day C. Morbidity associated with non-fatal heroin overdose. *Addiction.* 2002 Aug;97(8):963–7.
5. Hulse GK, English DR, Milne E, Holman CDJ. The quantification of mortality resulting from the regular use of illicit opiates. *Addiction.* 1999 Feb;94(2):221–9.
6. Ravndal E, Amundsen EJ. Mortality among drug users after discharge from inpatient treatment: an 8-year prospective study. *Drug and alcohol dependence.* 2010 May 1;108(1-2):65–9.
7. Seaman SR, Brettle RP, Gore SM. Mortality from overdose among injecting drug users recently released from prison: database linkage study. 1998;316(7129):426.
8. McLellan AT, Lewis DC, O'Brien CP, Kleber HD. Drug dependence, a chronic medical illness: implications for treatment, insurance, and outcomes evaluation. *JAMA* □: the journal of the American Medical Association. 2000 Oct 4;284(13):1689–95.
9. Gossop M, Stewart D, Browne N, Marsden J. Factors associated with abstinence, lapse or relapse to heroin use after residential treatment: protective effect of coping responses. *Addiction.* 2002;97(10):1259–67.
10. Verthein U, Bonorden-Kleij K, Degkwitz P, Dilg C, Köhler WK, Passie T, Soyka M, Tanger S, Vogel M, Haasen C. Long-term effects of heroin-assisted treatment in Germany. *Addiction (Abingdon, England).* 2008 Jun;103(6):960–6; discussion 967–8.
11. Kranzler HR, Wesson DR, Billot L. Naltrexone depot for treatment of alcohol dependence: a multicenter, randomized, placebo-controlled clinical trial. *Alcohol Clin Exp Res.* 2004/07/15 ed. 2004;28(7):1051–9.
12. Jayaram-Lindström N, Hammarberg A, Beck O, Franck J. Naltrexone for the treatment of amphetamine dependence: a randomized, placebo-controlled trial. *The American journal of psychiatry.* 2008 Nov;165(11):1442–8.
13. Agarwal LJ, Berger CE, Gill L. Naltrexone for severe self-harm behavior: a case report. *The American journal of psychiatry.* 2011 Apr;168(4):437–8.
14. Grant JE, Kim SW, Hartman BK. A double-blind, placebo-controlled study of the opiate antagonist naltrexone in the treatment of pathological gambling urges. *The Journal of clinical psychiatry.* 2008 May;69(5):783–9.
15. Mannelli P, Patkar AA, Peindl K, Gorelick DA, Wu L-T, Gottheil E. Very low dose naltrexone addition in opioid detoxification: a randomized, controlled trial. *Addiction biology.* 2009 Apr;14(2):204–13.
16. Nestler EJ, Aghajanian GK. Molecular and cellular basis of addiction. *Science.* 1997;278(5335):58.

17. Mannelli P, Gottheil E, Peoples JF, Oropeza VC, Van Bockstaele EJ. Chronic very low dose naltrexone administration attenuates opioid withdrawal expression. *Biological psychiatry*. 2004 Aug 15;56(4):261–8.
18. Childers SR. Opioid receptor-coupled second messenger systems. *Life sciences*. 1991 Jan;48(21):1991–2003.
19. Gekker G, Lokensgard JR, Peterson PK. Naltrexone potentiates anti-HIV-1 activity of antiretroviral drugs in CD4(+) lymphocyte cultures. *Drug Alcohol Depend*. 2001;64(3):257.
20. Volavka J, Cho D, Mallya A, Bauman J. Naloxone increases ACTH and cortisol levels in man. *The New England journal of medicine*. 1979 May 3;300(18):1056–7.
21. Minozzi S, Amato L, Vecchi S, Davoli M, Kirchmayer U, Verster A. Oral naltrexone maintenance treatment for opioid dependence. *Cochrane database of systematic reviews* (Online). 2011 Jan;(4):CD001333.
22. Martin WR, Jasinski DR, Mansky PA. Naltrexone, an antagonist for the treatment of heroin dependence. *Effects in man*. 1973;28(6):784.
23. Resnick RB, Volavka J, Freedman AM, Thomas M. Studies of EN-1639A (naltrexone): a new narcotic antagonist. *Am J Psychiatry*. 1974/06/01 ed. 1974;131(6):646–50.
24. Chiang CN, Hollister LE, Gillespie HK, Foltz RL. Clinical evaluation of a naltrexone sustained-release preparation. *Drug Alcohol Depend*. 1985;16(1):1.
25. Ling W, Casadonte P, Bigelow G, Kampman KM, Patkar A, Bailey GL, Rosenthal RN, Beebe KL. Buprenorphine implants for treatment of opioid dependence: a randomized controlled trial. *JAMA* □: the journal of the American Medical Association. 2010 Oct 13;304(14):1576–83.
26. Sullivan MA, Vosburg SK, Comer SD. Depot naltrexone: antagonism of the reinforcing, subjective, and physiological effects of heroin. *Psychopharmacology (Berl)*. 2006;189(1):37–46.
27. Bigelow GE, Preston KL, Schmittner J, Dong Q, Gastfriend DR. Opioid challenge evaluation of blockade by extended-release naltrexone in opioid-abusing adults: Dose-effects and time-course. *Drug and alcohol dependence*. 2012 Jul 1;123(1-3):57–65.
28. Hulse GK, Arnold-Reed DE, O’Neil G, Chan CT, Hansson R, O’Neil P. Blood naltrexone and 6-beta-naltrexol levels following naltrexone implant: comparing two naltrexone implants. 2004;
29. Tiihonen J, Krupitsky E, Verbitskaya E, Blokhina E, Mamontova O, Föhr J, Tuomola P, Kuoppasalmi K, Kiviniemi V, Zwartau E. Naltrexone implant for the treatment of polydrug dependence: a randomized controlled trial. *The American journal of psychiatry*. 2012 May 1;169(5):531–6.
30. Olsen L, Christophersen AS, Frogopsahl G, Waal H, Morland J. Plasma concentrations during naltrexone implant treatment of opiate-dependent patients. 2004;58 (2):219.
31. Waal H, Frogopsahl G, Olsen L, Christophersen AS, Morland J. Naltrexone implants -- duration, tolerability and clinical usefulness. A pilot study. *Eur Addict Res*. 2006;12(3):138–44.
32. Kunøe N, Lobmaier P, Vederhus JK, Hjerkin B, Gossop M, Hegstad S, Kristensen Ø, Waal H. Challenges to antagonist blockade during sustained-release naltrexone treatment. *Addiction* (Abingdon, England). 2010 Sep;105(9):1633–9.

33. Comer SD, Sullivan MA, Yu E, Rothenberg JL, Kleber HD, Kampman K, Dackis C, O'Brien CP. Injectable, sustained-release naltrexone for the treatment of opioid dependence: a randomized, placebo-controlled trial. *Arch Gen Psychiatry*. 2006/02/08 ed. 2006;63(2):210–8.
34. Krupitsky E, Nunes EV, Ling W, Illeperuma A, Gastfriend DR, Silverman BL. Injectable extended-release naltrexone for opioid dependence: a double-blind, placebo-controlled, multicentre randomised trial. *Lancet*. 2011 Apr 30;377(9776):1506–13.
35. Kunøe N, Lobmaier P, Vederhus JK, Hjerkin B, Hegstad S, Gossop M, Kristensen Ø, Waal H. Naltrexone implants after in-patient treatment for opioid dependence: randomised controlled trial. *The British journal of psychiatry* □: the journal of mental science. 2009 Jun;194(6):541–6.
36. Hulse GK, Morris N, Arnold-Reed D, Tait RJ. Improving clinical outcomes in treating heroin dependence: randomized, controlled trial of oral or implant naltrexone. *Arch Gen Psychiatry*. 2009/10/07 ed. 2009;66(10):1108–15.
37. Lobmaier PP, Kunøe N, Gossop M, Katevoll T, Waal H. Naltrexone implants compared to methadone: outcomes six months after prison release. *European addiction research*. 2010 Jan 26;16(3):139–45.
38. Krupitsky E, Zvartau E, Blokhina E, Verbitskaya E, Wahlgren V, Tsoy-Podosenin M, Bushara N, Burakov A, Masalov D, Romanova T, Tyurina A, Palatkin V, Slavina T, Pecoraro A, Woody GE. Randomized Trial of Long-Acting Sustained-Release Naltrexone Implant vs Oral Naltrexone or Placebo for Preventing Relapse to Opioid Dependence. *Archives of general psychiatry*. 2012 Sep 1;69(9):973–81.
39. Hulse GK, Tait RJ, Comer SD, Sullivan MA, Jacobs IG, Arnold-Reed D. Reducing hospital presentations for opioid overdose in patients treated with sustained release naltrexone implants. *Drug Alcohol Depend*. 2005;79(3):351–7.
40. Tait RJ, Ngo HT, Hulse GK. Mortality in heroin users 3 years after naltrexone implant or methadone maintenance treatment. *J Subst Abuse Treat*. 2007;
41. Kelty E, Hulse G. Examination of mortality rates in a retrospective cohort of patients treated with oral or implant naltrexone for problematic opiate use. *Addiction (Abingdon, England)*. 2012 May 5;
42. Krupitsky EM, Burakov AM, Tsoy MV, Egorova VY, Slavina TY, Grinenko AY, Zvartau EE, Woody GE. Overcoming opioid blockade from depot naltrexone (Prodetoxon). *Addiction*. 2007;102(7):1164–5.
43. Gibson AE, Degenhardt LJ, Hall WD. Opioid overdose deaths can occur in patients with naltrexone implants. *Med J Aust*. 2007;186(3):152–3.
44. Kunøe N, Lobmaier P, Vederhus JK, Hjerkin B, Hegstad S, Gossop M, Kristensen Ø, Waal H. Retention in naltrexone implant treatment for opioid dependence. *Drug and alcohol dependence*. 2010 Sep 1;111(1-2):166–9.
45. DeFulio A, Everly JJ, Leoutsakos J-MS, Umbricht A, Fingerhood M, Bigelow GE, Silverman K. Employment-based reinforcement of adherence to an FDA approved extended release formulation of naltrexone in opioid-dependent adults: a randomized controlled trial. *Drug and alcohol dependence*. 2012 Jan 1;120(1-3):48–54.
46. Stitzer ML, Vandrey R. Contingency management: utility in the treatment of drug abuse disorders. *Clinical pharmacology and therapeutics*. *American Society of Clinical Pharmacology and Therapeutics*; 2008 Apr 27;83(4):644–7.

47. Farrell M, Marsden J. Acute risk of drug-related death among newly released prisoners in England and Wales. *Addiction* (Abingdon, England). 2008 Feb;103(2):251–5.
48. Shewan D, Gemmell M, Davies JB. Behavioural change amongst drug injectors in Scottish prisons. *Social science & medicine* (1982). 1994 Dec;39(11):1585–6.
49. Brahen LS, Henderson RK, Capone T, Kordal N. Naltrexone treatment in a jail work-release program. 1984;45(9 Pt 2):49.
50. Chan KY. The Singapore naltrexone community-based project for heroin addicts compared with drugfree community-based program: the first cohort. 1996;3(2):87.
51. Cornish JW, Metzger D, Woody GE, Wilson D, McLellan AT, Vandergrift B, O'Brien CP. Naltrexone pharmacotherapy for opioid dependent federal probationers. *J Subst Abuse Treat*. 1998/01/23 ed. 1997;14(6):529–34.
52. Coviello DM, Cornish JW, Lynch KG, Alterman AI, O'Brien CP. A randomized trial of oral naltrexone for treating opioid-dependent offenders. *The American journal on addictions / American Academy of Psychiatrists in Alcoholism and Addictions*. 2010;19(5):422–32.
53. Coviello DM, Cornish JW, Lynch KG, Boney TY, Clark CA, Lee JD, Friedman PD, Nunes EV, Kinlock TW, Gordon MS, Schwartz RP, Nuwayser ES, O'Brien CP. A multisite pilot study of extended-release injectable naltrexone treatment for previously opioid-dependent parolees and probationers. *Substance abuse*: official publication of the Association for Medical Education and Research in Substance Abuse. Routledge; 2012 Jan;33(1):48–59.
54. Hall W, Capps B, Carter A. The use of depot naltrexone under legal coercion: the case for caution. *Addiction* (Abingdon, England). 2008 Dec;103(12):1922–4.
55. Ngo HTT, Tait RJ, Hulse GK. Hospital psychiatric comorbidity and its role in heroin dependence treatment outcomes using naltrexone implant or methadone maintenance. *Journal of psychopharmacology* (Oxford, England). 2011 Jun 1;25(6):774–82.
56. Ngo HT, Tait RJ, Arnold-Reed DE, Hulse GK. Mental health outcomes following naltrexone implant treatment for heroin-dependence. *Prog Neuropsychopharmacol Biol Psychiatry*. 2007;31(3):605–12.
57. Oncken C, Van Kirk J, Kranzler HR. Adverse effects of oral naltrexone: analysis of data from two clinical trials. *Psychopharmacology* (Berl). 2001;154(4):397.
58. Crowley TJ, Wagner JE, Zerbe G, Macdonald M. Naltrexone-induced dysphoria in former opioid addicts. 1985;142(9):1081.
59. Miotto K, McCann MJ, Rawson RA, Frosch D, Ling W. Overdose, suicide attempts and death among a cohort of naltrexone-treated opioid addicts. *Drug Alcohol Depend*. 1997;45(1-2):131.
60. Miotto K, McCann M, Basch J, Rawson R, Ling W. Naltrexone and dysphoria: fact or myth?[see comment]. [Review] [60 refs]. 2002;
61. Dean AJ, Saunders JB, Jones RT, Young RM, Connor JP, Lawford BR. Does naltrexone treatment lead to depression? Findings from a randomized controlled trial in subjects with opioid dependence. *J Psychiatry Neurosci*. 2006;31(1):38–45.
62. Yoburn BC, Sierra V, Lutfy K. Chronic opioid antagonist treatment: assessment of receptor upregulation. 1989;170(3):193.

63. Arnold-Reed DE, Hulse GK, Hansson RC, Murray SD, O'Neil G, Basso MR, Holman CD. Blood morphine levels in naltrexone-exposed compared to non-naltrexone-exposed fatal heroin overdoses. 2003;8(3):343.
64. Research Center for Drug Evaluation and Research. Postmarket Drug Safety Information for Patients and Providers - Information for Healthcare Professionals: Naltrexone Injection Site Reactions [naltrexone for extended-release injectable suspension (marketed as Vivitrol)]. Center for Drug Evaluation and Research, 2008. Accessed Sep 2012 on <http://www.fda.gov/Drugs/DrugSafety/PostmarketDrugSafetyInformationforPatientsandProviders/ucm126446.htm>
65. Hulse GK, Stalenberg V, McCallum D, Smit W, O'Neil G, Morris N, Tait RJ. Histological changes over time around the site of sustained release naltrexone-poly(DL-lactide) implants in humans. 2005;108(1):43–55.
66. Hulse GK, Low VH, Stalenberg V, Morris N, Thompson RI, Tait RJ, Phan CT, Ngo HT, Arnold-Reed DE. Biodegradability of naltrexone-poly(DL) lactide implants in vivo assessed under ultrasound in humans. *Addict Biol.* 2007;
67. Turncliff RZ, Dunbar JL, Dong Q, Silverman BL, Ehrich EW, Dilzer SC, Lasseter KC. Pharmacokinetics of long-acting naltrexone in subjects with mild to moderate hepatic impairment. *Journal of clinical pharmacology.* 2005 Dec 1;45(11):1259–67.
68. Lucey MR, Silverman BL, Illeperuma A, O'Brien CP. Hepatic safety of once-monthly injectable extended-release naltrexone administered to actively drinking alcoholics. *Alcoholism, clinical and experimental research.* 2008 Mar;32(3):498–504.
69. Jeffrey GP, MacQuillan G, Chua F, Galhenage S, Bull J, Young E, Hulse GK, O'Neil G. Hepatitis C virus eradication in intravenous drug users maintained with subcutaneous naltrexone implants. *Hepatology (Baltimore, Md.).* 2007 Jan;45(1):111–7.
70. Hulse GK, O'Neill G. A possible role for implantable naltrexone in the management of the high-risk pregnant heroin user. 2002;42(1):93.
71. Farid WO, Dunlop SA, Tait RJ, Hulse GK. The effects of maternally administered methadone, buprenorphine and naltrexone on offspring: review of human and animal data. *Current neuropharmacology.* 2008 Jul;6(2):125–50.
72. Jones HE, Chisolm MS, Jansson LM, Terplan M. Naltrexone in the treatment of opioid-dependent pregnant women: the case for a considered and measured approach to research. *Addiction (Abingdon, England).* 2012 May 4;
73. Unger A, Metz V, Fischer G. Opioid dependent and pregnant: what are the best options for mothers and neonates? *Obstetrics and gynecology international.* 2012 Jan;2012:195954.
74. Hulse GK, O'Neill G, Pereira C, Brewer C. Obstetric and neonatal outcomes associated with maternal naltrexone exposure. 2001;41(4):424.
75. Lobmaier P, Kornor H, Kunoe N, Bjorndal A. Sustained-release naltrexone for opioid dependence. *Cochrane Database Syst Rev.* 2008;(2):CD006140.