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Methadone maintenance therapy versus no opioid replacement therapy for opioid dependence (Review)

Mattick RP, Breen C, Kimber J, Davoli M, Breen R

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## TABLE OF CONTENTS

<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>HEADER</td>
<td>1</td>
</tr>
<tr>
<td>ABSTRACT</td>
<td>1</td>
</tr>
<tr>
<td>PLAIN LANGUAGE SUMMARY</td>
<td>2</td>
</tr>
<tr>
<td>BACKGROUND</td>
<td>2</td>
</tr>
<tr>
<td>OBJECTIVES</td>
<td>3</td>
</tr>
<tr>
<td>METHODS</td>
<td>3</td>
</tr>
<tr>
<td>RESULTS</td>
<td>4</td>
</tr>
<tr>
<td>DISCUSSION</td>
<td>5</td>
</tr>
<tr>
<td>AUTHORS’ CONCLUSIONS</td>
<td>6</td>
</tr>
<tr>
<td>ACKNOWLEDGEMENTS</td>
<td>6</td>
</tr>
<tr>
<td>REFERENCES</td>
<td>7</td>
</tr>
<tr>
<td>CHARACTERISTICS OF STUDIES</td>
<td>7</td>
</tr>
<tr>
<td>DATA AND ANALYSES</td>
<td>13</td>
</tr>
<tr>
<td>Analysis 1.1. Comparison 1 Methadone maintenance treatment vs no methadone maintenance treatment, Outcome 1 Retention in treatment.</td>
<td>13</td>
</tr>
<tr>
<td>Analysis 1.2. Comparison 1 Methadone maintenance treatment vs no methadone maintenance treatment, Outcome 2 Morphine positive urines.</td>
<td>14</td>
</tr>
<tr>
<td>Analysis 1.3. Comparison 1 Methadone maintenance treatment vs no methadone maintenance treatment, Outcome 3 Self reported heroin use.</td>
<td>15</td>
</tr>
<tr>
<td>Analysis 1.4. Comparison 1 Methadone maintenance treatment vs no methadone maintenance treatment, Outcome 4 Criminal activity.</td>
<td>16</td>
</tr>
<tr>
<td>Analysis 1.5. Comparison 1 Methadone maintenance treatment vs no methadone maintenance treatment, Outcome 5 Mortality.</td>
<td>17</td>
</tr>
<tr>
<td>WHAT’S NEW</td>
<td>17</td>
</tr>
<tr>
<td>HISTORY</td>
<td>17</td>
</tr>
<tr>
<td>CONTRIBUTIONS OF AUTHORS</td>
<td>18</td>
</tr>
<tr>
<td>DECLARATIONS OF INTEREST</td>
<td>18</td>
</tr>
<tr>
<td>SOURCES OF SUPPORT</td>
<td>18</td>
</tr>
<tr>
<td>INDEX TERMS</td>
<td>18</td>
</tr>
</tbody>
</table>
Methadone maintenance therapy versus no opioid replacement therapy for opioid dependence

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ABSTRACT

Background

Methadone maintenance was the first widely used form of opioid replacement therapy developed to treat heroin dependence, and it remains the best-researched treatment for this problem. Despite the widespread use of methadone in maintenance treatment for opioid dependence in many countries, it is a controversial treatment whose effectiveness has been disputed.

Objectives

To evaluate the effects of methadone maintenance treatment (MMT) compared with treatments that did not involve opioid replacement therapy (i.e., detoxification, offer of drug-free rehabilitation, placebo medication, wait-list controls) for opioid dependence.

Search strategy

We searched all the following databases up to 2001: Cochrane Drugs and Alcohol Review Group Register, the Cochrane Controlled Trials Register, MEDLINE, EMBASE, Current Contents, Psychlit, CORK [www.state.vt.us/adap/cork], Alcohol and Drug Council of Australia (ADCA) [www.adca.org.au], Australian Drug Foundation (ADF-VIC) [www.adf.org.au], Centre for Education and Information on Drugs and Alcohol (CEIDA) [www.ceida.net.au], Australian Bibliographic Network (ABN), and Library of Congress databases, available NIDA monographs and the College on Problems of Drug Dependence Inc. proceedings, the reference lists of all identified studies and published reviews; authors of identified RCTs were asked about other published or unpublished relevant RCTs.

Selection criteria

All randomised controlled clinical trials of methadone maintenance therapy compared with either placebo maintenance or other non-pharmacological therapy for the treatment of opioid dependence.

Data collection and analysis

Reviewers evaluated the papers separately and independently, rating methodological quality of concealment of allocation, data were extracted independently for meta-analysis and double-entered.
Main results

Six studies met the criteria for inclusion in this review, all were randomised clinical trials, two were double-blind. There were a total number of 954 participants. The method of concealment of allocation was inadequate in one study, not clearly described in four studies, but adequate in a sixth study. Based on the meta-analysis, methadone appeared statistically significantly more effective than non-pharmacological approaches in retaining patient in treatment (3 RCTs, RR=3.05; 95%CI: 1.75-5.35) and in the suppression of heroin use (3 RCTs, RR=0.32; 95%CI: 0.23-0.44), but not statistically in criminal activity (3 RCTs, RR=0.39; 95%CI: 0.12-1.25).

Authors’ conclusions

Methadone is an effective maintenance therapy intervention for the treatment of heroin dependence as it retains patients in treatment and decreases heroin use better than treatments that do not utilise opioid replacement therapy. It does not show a statistically significant superior effect on criminal activity.

Plain Language Summary

Methadone maintenance treatment can keep people who are dependent on heroin in treatment programs and reduce their use of heroin

Methadone is the most widely used replacement for heroin in medically-supported maintenance or detoxification programs. Several non-drug detoxification and rehabilitation methods are also used to try and help people withdraw from heroin. However the review found that people have withdrawn from trials when they are assigned to a drug-free program. Consequently, there are no trials comparing methadone maintenance treatment with drug-free methods other than methadone placebo trials, or comparing methadone maintenance with methadone for detoxification only. These trials show that methadone can reduce the use of heroin in dependent people, and keep them in treatment programs.

Background

Currently, the major form of medical therapy for heroin dependence internationally involves orally administered methadone. Methadone is an analgesic medication developed to treat pain in the 1940s. It has been, and is still, prescribed widely for the management of pain in America, Australia and Europe.

It was in New York in the 1960s, during an increase in heroin use and heroin dependence, that researchers (Dole 1965; Dole Nyswander 1967) examined different prescribed opioids to manage heroin dependence, and reported that they found that methadone was most suitable to the task. They believed that long-term heroin use caused a permanent metabolic deficiency in the central nervous system and an associated physiological disease, which required regular administration of opiates to correct the metabolic deficiency (Dole 1969 & Nyswander, 1965). The disorder of opioid dependence has been represented in the International Classification of Disease of the World Health Organisation. It is a chronic or long-term and relapsing disorder, and some believe that it requires ongoing maintenance medication.

The aspects of methadone that have led to its use as a substitute drug for heroin include the number of pharmacological features of opioids. At the basis of methadone maintenance treatment (MMT) is the observation that opioid analgesics can be substituted for one another (Jaffe 1990). Methadone at adequate doses (of 20mg to more than 100 mg) prevents or reverses withdrawal symptoms (Ward 1992), and thus reduces the need to use illegal heroin (Jaffe 1990). Methadone remains effective for approximately 24 hours, requiring a single daily dose rather than the more frequent administration of three to four times daily which occurs with the shorter-acting heroin (Jaffe 1990). Methadone can “block” the euphoric effects of heroin, discouraging illicit use and thereby relieving the user of the need or desire to seek heroin (Dole 1969). This allows the opportunity to engage in normative activities, and “rehabilitation” if necessary. Methadone can cause death in overdosage, like other similar medications such as morphine, and for this reason it is a treatment which is dispensed under medical supervision and relatively strict rules. In summary, methadone is a long-acting opioid analgesic with well-understood pharmacological characteristics which make it suitable for stabilising opioid dependent patients in a maintenance treatment approach.
There is evidence that the quality of the therapeutic relationship with staff in methadone clinics plus the intensity of these ancillary services, combined with the dose of methadone prescribed will all act to enhance the outcome for methadone treatment (Ward 1992), although this is not the focus of this review.

Methadone maintenance treatment remains one of the best researched treatments for opioid dependence (Cooper 1983; Gerstein 1990; Hargreaves 1983; Mattick 1993; Ward 1992). It is the only treatment for opioid dependence which has been clearly demonstrated to reduce illicit opiate use more than either no-treatment (Dole 1969; Yancovitz 1991), drug-free treatment (Gunne 1981), placebo medication (Newman 1979; Strain 1993a; Strain 1993a), or detoxification (Vanichseni 1991) in clinical controlled trials. These trials have been conducted by different research groups, in markedly differing cultural settings, yet have converged to provide similar results.

**OBJECTIVES**

The present systematic review aimed to provide an evaluation of the effectiveness of methadone maintenance treatment on opioid dependence compared with treatments that did not include an opioid replacement therapy. The focus of the review is on retention in treatment, opioid use as measured by objective urine results and from self-report, as well as criminal activity and patient mortality.

**METHODOLOGY**

**Criteria for considering studies for this review**

**Types of studies**
The literature was reviewed for all clinical controlled trials of MMT against another treatment which does not use opioid replacement therapy.

**Types of participants**
Individuals who were opioid dependent were the target population for this review. No distinction was made between those using heroin and those who have been in methadone treatment prior to entering the research trial treatment. No restrictions were imposed in terms of studies of outpatients, inpatients, those with comorbid states, etc.

**Types of interventions**
Interventions were included if they used methadone maintenance therapy (MMT). The MMT interventions were included even where they also employed other treatments, such as behavioural therapies or outpatient rehabilitation. The control groups were treated with placebo medication, withdrawal or detoxification (with or without ancillary medication), drug-free rehabilitation treatment (such as therapeutic communities), and no treatment or wait-list controls.

**Types of outcome measures**
Outcome measures:
- Primary outcomes
  - 1) retention in treatment
  - 2) mortality
  - 3) proportion of urinalysis results positive for heroin (or morphine)
  - 4) self-reported heroin use
  - 5) criminal activity
- Secondary outcomes
  - 1) use of other drugs
  - 2) physical health
  - 3) psychological health

**Search methods for identification of studies**
The search strategy was developed in consultation with a drug and alcohol research information specialist without language restrictions.

We searched:
2. Cochrane Controlled Trials Register for trials of methadone maintenance therapy to 2001.
3. MEDLINE (1966-2001) (OVID) was searched using the Cochrane Collaboration sensitive search strategies used to identify randomised trials in conjunction with the following to identify studies comparing methadone maintenance therapy and no methadone maintenance therapy:
   
   #1 exp methadone
   
   #2 (placebo or withdrawal or detoxification or untreated or no treatment or drug free or wait list).ti, ab, rw,sh.
   
   #3 exp pain/ or pain.ti, ab, rw, sh.
   
   #4 (1 and 2) not 3

   EMBASE (1980-2001) was searched using the following (OVID):
   
   #1 exp methadone/ or exp methadone treatment/ ct (limit to clinical trials)
   
   #2 exp drug dependence or exp substance abuse or exp drug abuse
   
   #3 1 and 2
   
   #4 limit to human
As several drug and alcohol journals are not indexed on the main electronic databases, Current Contents, Psychlit (-2001), CORK [www.state.vt.us/adap/cork], Alcohol and Drug Council of Australia (ADCA) [www.adca.org.au], Australian Drug Foundation (ADF -VIC) [www.adf.org.au], Centre for Education and Information on Drugs and Alcohol (CEIDA) [www.ceida.net.au], Australian Bibliographic Network (ABN), and Library of Congress databases were searched for studies and book chapters comparing methadone maintenance with other treatment.

4. The references of all identified studies and published reviews were inspected for more trials.

**Data collection and analysis**

Each potentially relevant study located in the search was obtained and independently assessed for inclusion by two of three reviewers. Data extraction for each study was undertaken by the same two reviewers, again independently. A standardised checklist was used for data extraction. Disagreement was dealt with by the third reviewer, acting as a mediator. If unresolved disagreements on inclusion, study quality or extraction occurred they were referred to the editor.

It is generally not the case that these trials were blinded. As such, methodological quality was assessed by assessment of the randomisation procedure and the likelihood that randomisation was not biased:

A. Low risk of bias (allocation clearly independent of clinical staff);  
B. Moderate risk of bias (some doubt about the independence of the allocation procedure); and  
C. High risk of bias (inadequate separation of randomisation from clinical staff).

A standardised effect size was calculated for each study, based on the main outcome measure reported. Where possible (relative risks and 95% confidence intervals for dichotomous outcomes (retention) using a random effects model and standardised mean differences for continuous outcomes were presented. To assess for statistical heterogeneity a test of homogeneity was undertaken. A standardised checklist was used for data extraction. Disagreement was dealt with by the third reviewer, acting as a mediator. If unresolved disagreements on inclusion, study quality or extraction occurred they were referred to the editor.

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**RESULTS**

**Description of studies**

See: Characteristics of included studies; Characteristics of excluded studies.

The information provided in the tables present the characteristics of the excluded and the included studies.

One study was not included. A study by Bale (Bale 1980) was an attempt to compare methadone against therapeutic community treatment of detoxification. Because the study failed, no useful data could be obtained from it for this review.

In total, six studies were included in this review. The first study by Dole (Dole 1969) was a two group randomised trial where patients either received methadone or placebo on a wait-list. The second study (Gunne 1981) randomly allocated patients to receive methadone maintenance or to be allocated to a drug-free rehabilitation. None of the patients allocated to drug-free rehabilitation took up the offer, refusing treatment after they had learnt that they would not receive methadone. There were two placebo controlled trials (Newman 1979, Strain 1993a). Finally, there have been two randomised clinical trials, one assessing methadone maintenance against methadone detoxification and the other assessing methadone maintenance against a wait-list control (Vanichseni 1991, Yancovitz 1991).

All studies were assessed to determine whether they provided data on retention in treatment, codeable results from urine analysis, self-reported drug use (particularly heroin use), criminal activity, and mortality. After reviewing the studies, it was realised that it was not possible to include urine results for cocaine and benzodiazepines as these were not reported in an analysable form for most studies. Thus, it was not possible to analyse data on either cocaine or benzodiazepine positive urine from these studies. However, it was possible to code data on retention in treatment, morphine positive urine, self-reported heroin use, criminal activity, and mortality.

**Risk of bias in included studies**

Of the six studies included in this review, two were placebo-controlled trials (Newman 1979, Strain 1993a). Both of these studies were double-blind but neither of these two studies provided sufficient data to be confident about the concealment of allocation. The first study conducted by Dole (Dole 1969) seemed to have adequate concealment of allocation. The other three studies had
reasonable concealment, and particularly the Yancovitz (Yancovitz 1991) had good concealment of allocation. The sample sizes in these studies were sometimes small, in that two studies having sample sizes of 32 and 34 (Dole 1969; Gunne 1981), respectively. The other four studies had sample sizes ranging from 100 to 240 (Newman 1979; Vanichseni 1991) patients up to 247 to 301 patients (Strain 1993a; Yancovitz 1991). The dosages of methadone used in these studies appears to have been adequate. In the first study, (Dole 1969) the dose at release from prison was 35 milligrams but patients were entered into a community program where blockade doses of approximately 100 milligrams were standard. In the study by Gunne (Gunne 1981) the doses are not clearly stated. The placebo-controlled study by Newman (Newman 1979) have an average dose on 97 milligrams per day. An average of 74 milligrams per day was reported in the study from Thailand (Vanichseni 1991). Strain (Strain 1993a) used doses of methadone of 50 and 20 milligrams per day. Finally, the study by Yancovitz 1991 used a maintenance dose of approximately 80 milligrams per day. As such, the results from the studies appear to use moderate to high doses on average.

Data on retention in treatment, self-reported heroin use, criminal activity, mortality and morphine positive urine were provided in the studies.

**Effects of interventions**

1. **Selection of Studies/Participants/Interventions**

Six studies were included in the review. The participants (n=954) were from a range of geographic regions including USA, Sweden, Hong Kong. Thailand and they were largely typical of heroin dependent individuals, in terms of age and gender characteristics. In some studies, only males were included but where females were included the gender distribution was as one would expect with majority of the participants being male.

As shown in the table of included studies, the interventions generally lasted for significant time of several weeks up to two years, although one study only ran for 45 days.

2. **Quantitative Analysis**

Retention in treatment could be coded from three studies, and the results showed that methadone has a superior retention rate compared with control conditions. When compared with placebo medications and with the wait-list control there was an advantage in terms of retention for methadone over the control groups (3 studies, 505 patients; RR= 3.05, 95% CI 1.75-5.35). The relative risk on a random effect model was applied. The chi-square test for heterogeneity was significant (p=0.018). The examination of the graphical representation of the relative risks showed that one study (Newman 1979) did appear to be slightly different from the other two studies, with higher RR, even though all single RRs where in the same direction of a positive effect (Vanichseni 1991; Strain 1993a).

Turning to the data from morphine positive urines, only two studies (Vanichseni 1991; Yancovitz 1991) provided data which were usable because of the way in which the data were typically reported. Specifically, many studies did not provide dichotomous data as to whether patients had morphine positive urines at the follow-up. However, the results from the two studies providing data on the presence/absence of morphine in urine at the follow-up showed an advantage of methadone above the control conditions (2 studies, 409 patients, RR= -0.32, 95% CI -0.40 -0.23), in this case detoxification or wait-list control, in reducing heroin use as shown by a lack of heroin metabolites in urine.

The results from the objective data on morphine positive urine were also supported by self-report data from three studies. In particular, studies from the USA and from Sweden (Dole 1969; Yancovitz 1991; Gunne 1981) all concurred to show an advantage for methadone above control in reduction of heroin use as reported by the patients (3 studies, 230 patients RD=0.32, 95% CI:0.23-0.44). The test for heterogeneity was not significant, indicating the results can be interpreted confidently. The results show clearly an advantage for methadone above the control conditions which included no treatment, referral to drug-free treatment, or frequent contact on a waiting list.

The results for the criminal activity variable, available for three studies, were consistent with the reduction in heroin use, even though the advantage for methadone beyond control in reducing criminal activity was not statistically significant (3 studies, 363 patients RR=0.39, 95% CI:0.12-1.25). The test for heterogeneity was not significant.

Turning finally to the evidence concerning the ability of methadone to prevent deaths, available for three studies, the results showed a trend in favour on methadone that was not statistically significant (3 studies, 435 patients RR=0.39, 95% CI:0.06-4.23). Other measures (e.g., use of other drugs, physical health, and psychological health) are too infrequently and irregularly reported in the literature to be usefully integrated in the quantitative review.

**DISCUSSION**

The results of the meta-analysis indicate that methadone is able to retain patients in treatment better than the drug-free alternatives (placebo medication, offer of drug-free treatment, detoxification, or wait-list control), to suppress heroin use based on morphine (the heroin metabolite) found in urine samples, and patient self-report. There was a greater reduction in criminal activity and mortality among the MMT patients, but these differences were not statistically significant. There is evidence from other literature showing that mortality is decreased in patients who are in methadone treatment.

Interestingly, the results from these six randomised trials all showed statistically significant positive benefits from methadone treat-
ment, despite their small sample sizes. Additional support for the efficacy of methadone maintenance treatment comes from the results of many observational studies wherein some statistical form of control has addressed alternative explanations of apparent effectiveness. These large scale observational studies have generally supported the results from the randomised clinical trials in showing that methadone maintenance treatment reduces the use of heroin and decreases criminal activity (Ward 1992). As noted earlier there is a broader international literature showing advantages for methadone beyond other treatments in terms of reduction of death (Ward 1992), even though the randomised trial data do not show this result.

Another relevant outcome to be considered would be seroconversion for HIV, which is the object of a separate Cochrane review in progress. Methadone maintenance treatment has been shown to reduce HIV risk taking behaviour (specifically reduction in needle sharing) and thereby has achieved a reduction in the transmission of HIV. Consistent with this it has been shown that methadone maintenance treatment is protective of patients, reducing HIV infection in geographic locations where HIV had spread rapidly among injecting drug users who had not entered treatment. We have commented elsewhere on two large prospective cohort studies in the USA which found methadone maintenance treatment protected against HIV infection. This outcome could not be addressed here as there are no randomised trials of methadone that have included HIV status as a measure, the evidence coming from observational studies (Ward 1992).

It is notable that the doses of methadone used in the randomised clinical trials are probably slightly higher than are being used currently in routine clinical practice in some parts of the world. This relative under dosing in clinical practice may lead to a reduction in the effectiveness of methadone, as the response to methadone treatment is dose-dependent. In addition, it is important to recognise that methadone treatment in these trials was often provided with substantial ancillary services. These ancillary services have included counselling, psycho-social services, medical services and often psychiatric care. The quality of the therapeutic relationship with staff in methadone clinics plus the intensity of these ancillary services, combined with the dose of methadone prescribed will all act to enhance the outcome for methadone treatment. The extent that clinical programs move away from such an approach might be expected to impact on the effectiveness of methadone.

This does not imply that methadone maintenance treatment will become ineffective. Even allowing for some reduction in effectiveness when methadone is not provided in the fashion that it has been in the clinical trials, it is still likely to be effective. The effects of methadone may be modest, if they are judged by unrealistic expectations of patients can easily achieve enduring abstinence from opioid drugs. Methadone nonetheless attracts and retains more patients than alternative treatments, and it does produce better outcomes amongst those who complete treatment. Methadone maintenance appears to provide better outcomes than simple detoxification programs, where the evidence suggests that short-term detoxification has no enduring effect on drug use (Mattick 1996).

**Authors’ Conclusions**

**Implications for practice**

The implications of the results of the meta-analytic review conducted and reported herein for clinical practice are that methadone maintenance treatment is an effective intervention for the management of heroin dependence. Methadone retains patients in treatment and reduces heroin use. Methadone should be supported as a maintenance treatment for heroin dependence.

**Implications for research**

Overall there are a relatively limited number of randomised clinical trials on the efficacy of methadone treatment compared to placebo. It does not seem feasible at this stage to conduct further randomised trials of methadone treatment. However, evidence on reduction of criminal activity and mortality from clinical trials is lacking calling for an additional systematic review of observational studies. Moreover, monitoring of the outcome of standard methadone treatment in clinical practice may be important as a research activity to demonstrate its ongoing effectiveness, or to determine whether its effectiveness is being compromised through the reduction of ancillary services or reduction in adequate dose levels. A number of measures (e.g., of other drug use, physical health, and psychological health) were too infrequently and irregularly reported in the literature to be usefully integrated in the quantitative review, but future research might address these important areas.

**Acknowledgements**

We acknowledge the assistance of the Cochrane Review Group Coordinating Centre, Rome.
References to studies included in this review

Dole 1969 [published data only]

Gunne 1981 [published data only]

Newman 1979 [published data only]

Strain 1993a [published data only]

Vanichseni 1991 [published data only]

Additional references

Cooper 1983

Dole 1965

Dole Nyswander 1967

Gerstein 1990

Hargreaves 1983

Jaffe 1990

Mattick 1993

Mattick 1996

Ward 1992

* Indicates the major publication for the study

References to studies excluded from this review

Bale 1980 [published data only]
### Characteristics of included studies  
**[ordered by study ID]**

#### Dole 1969

| Methods | Two group, open, randomised controlled trial.  
| Randomisation: release dates of treatment applicants were selected by lottery. Applicants who were not selected and demonstrated motivation for treatment became untreated controls.  
| Follow-up for 50 weeks.|
| Participants | Geographic region: USA  
| n = 32 males  
| mean age = 30 years  
| 15% European descent, 10% African-American, 7% Hispanic  
| Participants were inmates eligible for release over a four month period from New York City Correctional Institute for Men.  
| Eligibility criteria: heroin dependence for 5 or more years, record of 5 or more previous convictions, not committed to custody of Addiction Services Agency|
| Interventions | Control: wait-list  
| Treatment: 10 day methadone maintenance pre-release.  
| Initial dose 10 mg, increasing to 35 mg at release.  
| Continued methadone maintenance in outpatient clinic after release|
| Outcomes | Urinalysis (weekly for heroin, amphetamines, cocaine, barbituates and alcohol)  
| Employment / education  
| Reincarceration|
| Notes | |
| Risk of bias | Item  
| Authors’ judgement | Description  
| Allocation concealment? | No  
| C - Inadequate |

#### Gunne 1981

| Methods | Two group randomised clinical trial.  
| Randomisation: after eligibility established subjects were randomly allocated to methadone maintenance or to drug-free treatment|
| Participants | Geographic region: Sweden  
| Study setting: psychiatric research centre  
| n = 34, 23.5% female  
| Eligibility criteria: 20-24 years, history of at least 4 years IV heroin use, withdrawal signs and positive urine on admission, a minimum of
### Gunne 1981 (Continued)

| Interventions | Control: no treatment, could not apply for the methadone program for two years  
<table>
<thead>
<tr>
<th></th>
<th>Treatment: methadone maintenance treatment</th>
</tr>
</thead>
</table>
| Outcomes      | Illicit drug use / Urinalysis (3 x week)  
|               | Criminality  
|               | Vocational adjustment  
|               | Health  
|               | Mortality |
| Notes         | 2 controls obtained methadone from private practitioners and were excluded |

#### Risk of bias

<table>
<thead>
<tr>
<th>Item</th>
<th>Authors' judgement</th>
<th>Description</th>
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<tr>
<td>Allocation concealment?</td>
<td>Unclear</td>
<td>B - Unclear</td>
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### Newman 1979

| Methods | Double blind randomised clinical trial  
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<td>Randomisation: subjects randomly allocated on discharge from hospital after 2 week stabilisation on 60mg methadone to detoxification or continued maintenance</td>
</tr>
</tbody>
</table>
| Participants | Geographic region: Hong Kong  
|              | n = 100 males  
|              | Study setting: Hospital and outpatient clinic  
|              | mean age = 38 years  
|              | Eligibility criteria: male, 22-58 years, history of heroin dependence for at least 4 years and at least one previous treatment, current heroin dependence by three consecutive positive urine samples, voluntary application for admission (criminal justice referrals excluded), resident with fixed address, absence of past or present major psychiatric or medical illness |
| Interventions | Treatment: methadone maintenance - flexible dose (average 97 mg / day)  
|               | Control: detoxification from 60mg methadone at 1mg/day for 60 days, placebo thereafter |
| Outcomes      | Illicit drug use / Urinalysis (daily collection, analysed 2 x week for morphine only)  
|               | Retention  
|               | Criminal activity  
|               | Mortality |
| Notes         | |

#### Risk of bias

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<th>Item</th>
<th>Authors' judgement</th>
<th>Description</th>
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Strain 1993a

Methods
Three group, double-blind, placebo controlled randomised controlled trial. Patients were stratified by race and sex and randomly assigned to a fixed dose schedule at admission. Treatment group assignment, stabilisation dose and dosing schedules were blind to patient and clinic staff with patient contact.

Participants
Geographic region: USA
Study setting: methadone treatment research clinic
n = 247
mean age = 34
70% male
50% black
84% unmarried
62% unemployed
Eligibility criteria: 18-50 years, history of IV opioid dependence, no chronic medical illness, absence of major mental illness, negative pregnancy test and at least three months since last treatment at the clinic

Interventions
Initial treatment of active methadone for at least 5 weeks.
15 weeks of stable dosing at 50, 20 or 0 mg per day
Gradual tapering for those receiving active methadone from weeks 21-26
Individual counselling and group therapy (weekly).

Outcomes
Retention
Treatment compliance
Illicit drug use / Urinalysis (collected 3 x weekly, one sample selected at random for analysis for opioids, cocaine and benzodiazepines)

Notes
A subsample of 0mg patients (n=44) received an 8 week induction, reaching 0mg at 9 weeks. Data for patients in alternate 0mg treatment groups are collapsed

Risk of bias

<table>
<thead>
<tr>
<th>Item</th>
<th>Authors’ judgement</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allocation concealment?</td>
<td>Unclear</td>
<td>B - Unclear</td>
</tr>
</tbody>
</table>

Vanichseni 1991

Methods
Two group, open label, randomised clinical trial, with participants who applied for 45 day methadone detoxification and had at least six prior treatment episodes were randomly assigned to methadone maintenance or detoxification.

Participants
Geographic region: Thailand
Study setting: narcotics clinic

Risk of bias
Vanichseni 1991  

(Continued)

| n = 240 males |
| Eligibility criteria: heroin injectors applying for 45-day detoxification, at least 6 prior treatment episodes at the clinic |

Interventions

| Treatment: methadone maintenance (flexible dose, average 74mg) |
| Control: standard 45 day methadone detoxification |

Outcomes

| Retention |
| Illicit drug use |
| Urinalysis (2 x week for opiates) |

Notes

Risk of bias

| Item |
| Authors’ judgement |
| Description |

| Allocation concealment? |
| Unclear |
| B - Unclear |

Yancovitz 1991

Methods

Two group randomised clinical trial, with opioid dependent participants on waiting-lists for comprehensive methadone maintenance programs who were randomised to either the interim methadone program or wait list with frequent contact

Participants

| Geographic region: USA |
| n = 301 |
| Study setting: interim methadone clinic |
| 79.4% male |
| 10% White |
| 35% Black |
| 55% Hispanic |
| Eligibility criteria: wait list for comprehensive methadone maintenance program |

Interventions

| Control: wait-list with frequent contact |
| Treatment: “interim” methadone maintenance; standard physical exam on admission, flexible dosing 5 days a week, pick up on weekends from another site, minimal counselling, referral to community agencies |

Outcomes

| Urinalysis (2 x weekly for heroin and cocaine) |
| Entry into conventional treatment |

Notes

For the first 3 months of the study there were three experimental groups; interim methadone, wait-list with frequent contact and bi-weekly urinalysis, and the wait-list with no contact. Recruitment slowed which resulted in the protocol being changed two experimental groups; interim methadone and wait-list with frequent contact. The wait-list then only lasted one month at which time the participants were switched to a methadone program. Data from the initial discontinued minimal contact group is not include in the analysis.
Risk of bias

<table>
<thead>
<tr>
<th>Item</th>
<th>Authors’ judgement</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allocation concealment?</td>
<td>Yes</td>
<td>A - Adequate</td>
</tr>
</tbody>
</table>

Characteristics of excluded studies  [ordered by study ID]

<table>
<thead>
<tr>
<th>Study</th>
<th>Reason for exclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bale 1980</td>
<td>The authors planned to conduct a randomised controlled trial comparing methadone maintenance, therapeutic communities and detoxification programs. Ethical and practical problems prevented random assignment and the study therefore does not meet inclusion criteria for this review</td>
</tr>
</tbody>
</table>
### DATA AND ANALYSES

#### Comparison 1. Methadone maintenance treatment vs no methadone maintenance treatment

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Retention in treatment</td>
<td>3</td>
<td>505</td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>3.05 [1.75, 5.35]</td>
</tr>
<tr>
<td>2 Morphine positive urines</td>
<td>2</td>
<td>409</td>
<td>Risk Difference (M-H, Random, 95% CI)</td>
<td>-0.32 [-0.40, -0.23]</td>
</tr>
<tr>
<td>3 Self reported heroin use</td>
<td>3</td>
<td>230</td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>0.32 [0.23, 0.44]</td>
</tr>
<tr>
<td>4 Criminal activity</td>
<td>3</td>
<td>363</td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>0.39 [0.12, 1.25]</td>
</tr>
<tr>
<td>5 Mortality</td>
<td>3</td>
<td>435</td>
<td>Risk Ratio (M-H, Random, 95% CI)</td>
<td>0.49 [0.06, 4.23]</td>
</tr>
</tbody>
</table>

#### Analysis 1.1. Comparison 1 Methadone maintenance treatment vs no methadone maintenance treatment, Outcome 1 Retention in treatment.

**Review:** Methadone maintenance therapy versus no opioid replacement therapy for opioid dependence

**Comparison:** 1 Methadone maintenance treatment vs no methadone maintenance treatment

**Outcome:** 1 Retention in treatment

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Methadone MT n/N</th>
<th>Control n/N</th>
<th>Risk Ratio M-H Random 95% CI</th>
<th>Weight</th>
<th>Risk Ratio M-H Random 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Newman 1979</td>
<td>38/50</td>
<td>5/50</td>
<td>22.6 % 7.60 [3.26, 17.71]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Strain 1993a</td>
<td>44/84</td>
<td>17/81</td>
<td>35.2 % 2.50 [1.56, 3.99]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vanichseni 1991</td>
<td>91/120</td>
<td>41/120</td>
<td>42.2 % 2.22 [1.70, 2.90]</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Total (95% CI)</strong></td>
<td><strong>254</strong></td>
<td><strong>251</strong></td>
<td><strong>100.0 %</strong> 3.05 [1.75, 5.35]</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Total events: 173 (Methadone MT), 63 (Control)

Heterogeneity: $\tau^2 = 0.17$; $\chi^2 = 8.01$, df = 2 ($P = 0.02$); $I^2 = 75$

Test for overall effect: $Z = 3.91$ ($P = 0.000092$)
### Analysis 1.2. Comparison 1 Methadone maintenance treatment vs no methadone maintenance treatment, Outcome 2 Morphine positive urines.

**Review:** Methadone maintenance therapy versus no opioid replacement therapy for opioid dependence

**Comparison:** 1 Methadone maintenance treatment vs no methadone maintenance treatment

**Outcome:** 2 Morphine positive urines

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>MMT</th>
<th>Control</th>
<th>Risk Difference</th>
<th>Risk Difference</th>
<th>Weight</th>
<th>Risk Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n/N</td>
<td>n/N</td>
<td>M-H, Random, 95% CI</td>
<td>M-H, Random, 95% CI</td>
<td></td>
<td>M-H, Random, 95% CI</td>
</tr>
<tr>
<td>Vanichseni 1991</td>
<td>70/120</td>
<td>109/120</td>
<td>66.2 %</td>
<td>-0.32 [ -0.43, -0.22 ]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yancovitz 1991</td>
<td>22/75</td>
<td>56/94</td>
<td>33.8 %</td>
<td>-0.30 [ -0.45, -0.16 ]</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Total (95% CI)</strong></td>
<td><strong>195</strong></td>
<td><strong>214</strong></td>
<td><strong>100.0 %</strong></td>
<td><strong>-0.32 [ -0.40, -0.23 ]</strong></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Total events: 92 (MMT), 165 (Control)

Heterogeneity: $\tau^2 = 0.0, \chi^2 = 0.06, df = 1$ ($P = 0.80$); $I^2 = 0.0$

Test for overall effect: $Z = 7.48$ ($P < 0.00001$)
**Analysis 1.3. Comparison 1 Methadone maintenance treatment vs no methadone maintenance treatment, Outcome 3 Self reported heroin use.**

Review: Methadone maintenance therapy versus no opioid replacement therapy for opioid dependence

Comparison: 1 Methadone maintenance treatment vs no methadone maintenance treatment

Outcome: 3 Self reported heroin use

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Methadone MT</th>
<th>Control</th>
<th>Risk Ratio</th>
<th>Weight</th>
<th>Risk Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n/N</td>
<td>n/N</td>
<td>M-H Random, 95% CI</td>
<td></td>
<td>M-H Random, 95% CI</td>
</tr>
<tr>
<td>Dale 1969</td>
<td>2/12</td>
<td>15/15</td>
<td>8.3 % 0.20 [0.06, 0.61]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gunne 1981</td>
<td>5/17</td>
<td>12/17</td>
<td>16.3 % 0.42 [0.19, 0.93]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yancovitz 1991</td>
<td>21/75</td>
<td>83/94</td>
<td>75.5 % 0.32 [0.22, 0.46]</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Total (95% CI)</strong></td>
<td><strong>104</strong></td>
<td><strong>126</strong></td>
<td><strong>100.0 % 0.32 [0.23, 0.44]</strong></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Total events: 28 (Methadone MT), 110 (Control)

Heterogeneity: $\tau^2 = 0.0$, $\text{Chi}^2 = 1.14$, df = 2 ($P = 0.57$); $I^2 = 0.0$

Test for overall effect: $Z = 6.96$ ($P < 0.00001$)
### Analysis 1.4. Comparison 1 Methadone maintenance treatment vs no methadone maintenance treatment, Outcome 4 Criminal activity.

**Review:** Methadone maintenance therapy versus no opioid replacement therapy for opioid dependence

**Comparison:** 1 Methadone maintenance treatment vs no methadone maintenance treatment

**Outcome:** 4 Criminal activity

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Treatment</th>
<th>Control</th>
<th>Risk Ratio</th>
<th>Weight</th>
<th>Risk Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n/N</td>
<td>n/N</td>
<td>M-H, Random, 95% CI</td>
<td></td>
<td>M-H, Random, 95% CI</td>
</tr>
<tr>
<td>Dole 1969</td>
<td>3/12</td>
<td>15/16</td>
<td>0.27 [0.10, 0.72]</td>
<td>65.9 %</td>
<td></td>
</tr>
<tr>
<td>Gunne 1981</td>
<td>0/17</td>
<td>2/17</td>
<td>0.20 [0.01, 3.88]</td>
<td>13.9 %</td>
<td></td>
</tr>
<tr>
<td>Yancovitz 1991</td>
<td>2/149</td>
<td>1/152</td>
<td>2.04 [0.19, 22.26]</td>
<td>20.2 %</td>
<td></td>
</tr>
</tbody>
</table>

**Total (95% CI)** | 178 | 185 | 100.0 % | 0.39 [0.12, 1.25] |

Total events: 5 (Treatment), 18 (Control)

Heterogeneity: Tau² = 0.29; Chi² = 2.54, df = 2 (P = 0.28); I² = 21%

Test for overall effect: Z = 1.59 (P = 0.11)
Analysis 1.5. Comparison 1 Methadone maintenance treatment vs no methadone maintenance treatment, Outcome 5 Mortality.

Review: Methadone maintenance therapy versus no opioid replacement therapy for opioid dependence

Comparison: 1 Methadone maintenance treatment vs no methadone maintenance treatment

Outcome: 5 Mortality

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Treatment</th>
<th>Control</th>
<th>Risk Ratio M-H, Random, 95% CI</th>
<th>Weight</th>
<th>Risk Ratio M-H, Random, 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gunne 1981</td>
<td>0/17</td>
<td>4/17</td>
<td>3.12%</td>
<td>0.11 [0.01, 1.92]</td>
<td></td>
</tr>
<tr>
<td>Newman 1979</td>
<td>3/50</td>
<td>1/50</td>
<td>39.6%</td>
<td>3.00 [0.32, 27.87]</td>
<td></td>
</tr>
<tr>
<td>Yancovitz 1991</td>
<td>0/149</td>
<td>2/152</td>
<td>29.2%</td>
<td>0.20 [0.01, 4.21]</td>
<td></td>
</tr>
<tr>
<td><strong>Total (95% CI)</strong></td>
<td><strong>216</strong></td>
<td><strong>219</strong></td>
<td><strong>100.0%</strong></td>
<td><strong>0.49 [0.06, 4.23]</strong></td>
<td></td>
</tr>
</tbody>
</table>

Total events: 3 (Treatment), 7 (Control)
Heterogeneity: Tau^2 = 1.77; Chi^2 = 3.89, df = 2 (P = 0.14); I^2 = 49%
Test for overall effect: Z = 0.65 (P = 0.52)

WHAT’S NEW

Last assessed as up-to-date: 23 February 2003.

<table>
<thead>
<tr>
<th>Date</th>
<th>Event</th>
<th>Description</th>
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<tbody>
<tr>
<td>26 March 2008</td>
<td>Amended</td>
<td>Converted to new review format.</td>
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</tbody>
</table>

HISTORY


Review first published: Issue 4, 2002
CONTRIBUTIONS OF AUTHORS

Contributions: Richard P Mattick, Jo Klimber and Courtney Breen reviewed the papers, with Courtney Breen and Richard P. Mattick coding data from the papers for meta-analysis.

Richard P. Mattick conceptualised the review and Courtney Breen conducted the initial literature searches.

Richard P. Mattick wrote the analysis sections and discussion. Marina Davoli was the contact editor of the review and contributed to the writing of the final version of the review.

Marica Ferri from the Rome Editorial Base provided comments and copyediting on the drafts of this review; Roberto D’Amico from the Cochrane Statistical Methods Group provided advice on statistical analysis issues.

DECLARATIONS OF INTEREST

None.

SOURCES OF SUPPORT

Internal sources

- National Drug and Alcohol Research Centre, University of New South Wales, Sydney, Australia.

External sources

- Commonwealth Department of Health and Aged Care, Canberra, Australia.

INDEX TERMS

Medical Subject Headings (MeSH)

Metabolic Detoxication, Drug; Methadone [*therapeutic use]; Narcotics [*therapeutic use]; Opioid-Related Disorders [*rehabilitation]; Randomized Controlled Trials as Topic
MeSH check words

Humans