Comparing the Effects of Methadone, Buprenorphine, and Opium Tincture Maintenance Therapy on Sexual Function

Ali Kheradmand1, Ahad Fazeli2, Azadeh Mazaheri Meybodi1

Abstract

Background: Opioid use disorder is a major concern to public health, and opioid maintenance treatment on methadone or buprenorphine is a widely used approach. On the other hand, in recent years, there has been more regards for the use of opium tincture for detoxification and maintenance treatment of opioid dependence in certain parts of the world. The purpose of our research was to compare sexual impairments of methadone maintenance treatment (MMT), buprenorphine maintenance treatment (BMT), and opium tincture maintenance treatment (OMT) in patients.

Methods: The study sample consisted of opium-addicted men candidates for maintenance treatment in an addiction quitting clinic in Tehran, Iran, from November 2017 to February 2018. Participants (n = 84) were randomly assigned to three groups (of the equal number), receiving either methadone tablet, buprenorphine sublingual tablet, and opium tincture. The average score for sexual function was evaluated using the Arizona Sexual Experiences Scale (ASEX) at the beginning and after 3 months after treatment.

Findings: Although there was no significant different in ASEX scores between the groups at the beginning and end of the study (P > 0.05), but the difference was significant in each group in comparing by themselves.

Conclusion: These results showed that sexual dysfunction became better after opioid substitution therapies, and no differences were observed on sexual dysfunction between the three groups.

Keywords: Methadone; Buprenorphine; Opium tincture; Sexual dysfunctions; Opioid substitution therapies

Citation: Kheradmand A, Fazeli A, Mazaheri Meybodi A. Comparing the Effects of Methadone, Buprenorphine, and Opium Tincture Maintenance Therapy on Sexual Function. Addict Health 2019; 11(2): 120-8.

Received: 30.11.2018 Accepted: 05.02.2019

1- Taleghani Hospital Research Development Committee AND Department of Psychiatry, School of Medicine, Shahid Beheshti University of Medical Sciences, Tehran, Iran
2- Department of Psychiatry, School of Medicine, Shahid Beheshti University of Medical Sciences, Tehran, Iran
Correspondence to: Ali Kheradmand, Email: a.alikheradmand@sbmu.ac.ir

This is an open-access article distributed under the terms of the Creative Commons Attribution-NonCommercial 4.0 Unported License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.
Introduction

Opioids are one of the most common substance abuse materials for which patients seek treatment. Globally the burden is estimated to be 32.4 million with an overall prevalence of 0.7%. Methadone maintenance treatment (MMT) is identified as a potent substitute therapy for opioid addiction.

Sexual dysfunctions such as erectile dysfunction (ED), ejaculatory disturbances, and lack of desire for sexual relations are often reported in a considerable number of men patients on methadone maintenance. The fundamental process may be the reaction of hypothalamus and pituitary system to heroin and methadone, and also the lower performance of dopaminergic neurons of the mesolimbic area.

Methadone’s long-term stimulation of the μ-opioid receptors changes the function of the tubero-infundibular axis and dopaminergic influences on prolactin, which can affect sexual performance. High presence of circulating prolactin inhibits gonadotropin-releasing hormone (GnRH), which decreases sex hormones including testosterone. Decrease in testosterone measures may cause low sexual desire.

In conjunction with MMT, there were studies that showed that buprenorphine maintenance treatment (BMT) was an alternative effective treatment for opioid addiction. Buprenorphine is a mixed agonist-antagonist opioid, with a low intrinsic activity and a high affinity for the μ-opioid receptor, and no intrinsic activity but a high affinity for the κ-opioid receptor.

Although animal studies showed no significant changes in the basement membrane, seminiferous tubules, Sertoli cells, interstitial tissue, or sperm compared to a group that received methadone but in humans, some studies reported more frequent sexual and ED to methadone, while other ones found no significant differences between the two maintenance treatments.

Recently MMT and opium tincture maintenance treatment (OMT) for detoxification has achieved especial attention in some parts of the world. Opium tincture also called Laudanum is a clear, reddish-brown hydroalcoholic preparation of opium with a characteristic odor and bitter taste. Morphone is the active ingredient of opium tincture with the chemical formula of C_{17}H_{19}NO_{3}, and each milliliter of opium tincture contains 10 mg of morphine equivalent.

Opium tincture has gained growing popularity since its introduction in the national protocol of opioid addiction treatment in Iran in 2010, and is now widely used (64000 patients) as the second most common medication after methadone or maintenance treatment of opioid reliance.

Despite the increasing prevalence of opium tincture use as maintenance treatment in Iran, however, there is insufficient information about long-term use of opium tincture as a new option of maintenance treatment in opioid-addicted patients on sexual function.

A qualitative research showed a number of patients on MMT who faced sexual impairment had pulled back from intercourse with their partners leading to dissensions. Outcome of these dissensions were undesirable influences on the rehabilitation, early treatment termination, methadone dose cutting, and using under counter drugs for sexual power. Sexual impairments are not potentially fatal but can reduce the quality of life due to withdrawal from sexual relations.

Given that sexual impairment is a serious problem, the present study aimed to assess the sexual impairment in patients undergoing MMT, BMT, or OMT.

Methods

This randomized, open trial was conducted from November 2017 to February 2018 in an addiction quitting clinic, affiliated to Shahid Beheshti University of Medical Sciences, Tehran, Iran. This research was accepted by the Research and Ethics group of the School of Medicine at Shahid Beheshti University of Medical Sciences. All patients were enrolled in the study after obtaining written informed consent.

The study sample consisted of opium-addicted men candidates for maintenance treatment. After obtaining written informed consents, the subjects (n = 84) were randomly assigned to three groups (of the equal number), receiving either methadone, buprenorphine tablet, or opium tincture for 12 weeks. Regular follow-up was conducted every two weeks by a psychiatrist asking patients about their sexual problems.

Patients permitted were those who had the following criteria: 1. receiving maintenance

http://ahj.kmu.ac.ir, 04 April
treatment for the first time; 2. having no comorbidity that could affect sexual performance; 3. no alcohol consumption; 4. recent cessation of benzodiazepines; and 5. not using drugs or stimulants.

The criteria for exclusion were as follows: 1. leaving the MMT program; 2. the psychiatrist’s conclusion for ending the bupropion or opium tincture treatment for any rational; and 3. seizures.

The average score of sexual performance was evaluated using the Arizona Sexual Experiences Scale (ASEX) score. Methadone, bupropion, or opium tincture was administered by a nurse who was not involved in the rating of patients. The sample size was calculated as 28 subjects per group to detect 20% difference in ASEX score between groups.

The ASEX is a brief 5-item questionnaire designed to measure sexual functioning in the domains of sexual drive, arousal, penile erection/vaginal lubrication, satisfaction, and ability of orgasm. A 6-point scale from 1 (hypo-function) to 6 (hyper-function) was used, total score was between 5 and 30. Higher score was in favor of higher sexual dysfunction. The subjects were assessed for the past three months. The ASEX had a Cronbach’s alpha of 90%, a good reliability, and a correlation coefficient of 0.80.21

Before and after intervention, data were collected from all participants, using ASEX. Organization, supervision, data collection, data evaluation, and conclusion were performed by a psychiatric resident. For data analysis, t, chi-square, and ANCOVA tests were performed, using SPSS software (version 18, SPSS Inc., Chicago, IL, USA). P-value of less than 0.05 was statistically significant.

**Results**

Eighty-four men candidates for maintenance treatment participated in this study (each group had 28 patients). There was no remarkable difference between mean age of participants in MMT (37.05 ± 10.40 years), BMT (38.78 ± 9.40) and OMT (42.21 ± 9.60 years) group (P > 0.05).

Table 1 shows demographic features of patients including employment status, level of education, and marital status in three groups at baseline. There was no significant differences between the groups.

| Variable                      | Group            | P  |
|-------------------------------|------------------|----|
| Employment                    | MMT              |    |
| Employed                      | 10               | 15 | 9  | > 0.05 |
| Unemployed                    | 18               | 13 | 19 |
| Education (years)             | MMT              |    |
| <12                           | 13               | 15 | 8  | > 0.05 |
| 12-16                         | 11               | 13 | 16 |
| >16                           | 4                | 0  | 4  |
| Marital status                | MMT              |    |
| Single                        | 10               | 11 | 13 |
| Married                       | 18               | 17 | 15 |

MMT: Methadone maintenance treatment; BMT: Buprenorphine maintenance treatment; OMT: Opium tincture maintenance treatment

The results represented in table 2 show the mean scores of sexual dysfunction in the MMT, BMT, and OMT groups at the start. No remarkable difference was seen in baseline sexual function mean scores, based on ASEX questionnaire, between three groups at the start of the study.

| Domain                      | Group            | P  |
|-----------------------------|------------------|----|
| Desire/drive                | MMT              |    |
| 2.3 ± 1.1                   |                  |    |
| Arousal                     | MMT              |    |
| 2.5 ± 1.3                   |                  |    |
| Erection                    | MMT              |    |
| 2.4 ± 1.3                   |                  |    |
| Ability to reach orgasm     | MMT              |    |
| 2.9 ± 1.7                   |                  |    |
| Satisfaction with orgasm    | MMT              |    |
| 2.8 ± 1.3                   |                  |    |
| Total                       | MMT              |    |
| 13.1 ± 5.6                  |                  |    |

The amounts are presented as mean ± standard deviation (SD).

ASEX: Arizona sexual experiences scale; MMT: Methadone maintenance treatment; BMT: Buprenorphine maintenance treatment; OMT: Opium tincture maintenance treatment

*Based on the one-way ANOVA test.
The follow-up of sexual function assessment in patients on MMT, BMT, and OMT at week 12 of the study did not show any significant difference between groups (Table 3).

Although there was no significant different in ASEX scores at the beginning and end of the study between groups, but this difference was significant in each group in comparing by themselves (Table 4).

An analysis of marital status showed that there were no important differences in ASEX scores between married (Table 5) and single (Table 6) patients in all groups (P > 0.05).

Table 3. Comparison of the mean Arizona sexual experiences scale (ASEX) domain scores in study groups after 3 months of therapy

| Domain            | MMT       | BMT       | OMT       | P*  |
|-------------------|-----------|-----------|-----------|-----|
| Desire/drive      | 3.3 ± 1.1 | 3.6 ± 0.7 | 3.4 ± 1.1 | 0.60|
| Arousal           | 4.3 ± 1.1 | 4.2 ± 1.1 | 4.1 ± 1.0 | 0.75|
| Erection          | 3.3 ± 1.3 | 3.5 ± 1.2 | 3.6 ± 0.8 | 0.70|
| Ability to reach orgasm | 4.0 ± 1.1 | 4.2 ± 1.1 | 4.1 ± 0.9 | 0.81|
| Satisfaction with orgasm | 3.9 ± 1.5 | 3.3 ± 0.9 | 3.3 ± 1.2 | 0.12|
| Total             | 19.0 ± 5.1| 18.6 ± 4.1| 18.8 ± 4.0| 0.92|

The amounts are presented as mean ± standard deviation (SD).
ASEX: Arizona sexual experiences scale; MMT: Methadone maintenance treatment; BMT: Buprenorphine maintenance treatment; OMT: Opium tincture maintenance treatment
*Based on the one-way ANOVA test.

Table 4. Comparison of the mean Arizona sexual experiences scale (ASEX) total scores in the study groups after 3 months of therapy

| Group | ASEX score Before | After | P*  |
|-------|-------------------|-------|-----|
| MMT   | 13.1 ± 5.6        | 19.0 ± 5.1 | < 0.01 |
| BMT   | 13.2 ± 5.3        | 18.6 ± 4.1 | < 0.01 |
| OMT   | 11.0 ± 1.4        | 18.8 ± 4.0 | < 0.01 |

The amounts are presented as mean ± standard deviation (SD).
ASEX: Arizona sexual experiences scale; MMT: Methadone maintenance treatment; BMT: Buprenorphine maintenance treatment; OMT: Opium tincture maintenance treatment
*Based on the paired-samples t test.

Table 5. Comparison of the mean Arizona sexual experiences scale (ASEX) scores of married patients in study groups after 3 months of therapy

| Domain            | MMT       | BMT       | OMT       | P*  |
|-------------------|-----------|-----------|-----------|-----|
| Desire/drive      | 3.3 ± 1.1 | 3.4 ± 0.7 | 3.5 ± 1.1 | 0.66|
| Arousal           | 4.0 ± 1.1 | 4.0 ± 1.1 | 4.2 ± 1.0 | 0.84|
| Erection          | 3.3 ± 1.3 | 3.3 ± 1.2 | 3.5 ± 0.8 | 0.77|
| Ability to reach orgasm | 3.7 ± 1.1 | 3.9 ± 1.1 | 4.3 ± 0.9 | 0.14|
| Satisfaction with orgasm | 3.2 ± 1.5 | 3.3 ± 0.9 | 3.3 ± 1.2 | 0.89|
| Total             | 17.9 ± 5.1| 18.2 ± 4.1| 18.3 ± 4.0| 0.92|

The amounts are presented as mean ± standard deviation (SD).
ASEX: Arizona sexual experiences scale; MMT: Methadone maintenance treatment; BMT: Buprenorphine maintenance treatment; OMT: Opium tincture maintenance treatment
*Based on the one-way ANOVA test.
In a meta-analysis by Yee et al., sexual dysfunction was significantly higher in MMT group in comparison with BMT group.\textsuperscript{24} Our justification for the differences is the differences in characteristics of study population. Moreover, the difference in results may be due to socio-economic differences across the communities.\textsuperscript{25}

Some research shows the interference of methadone with the production of hormones of hypothalamus and pituitary regulatory which enhances serum prolactin, and reduces gonadotropin releasing hormone leading to reduced testosterone production.\textsuperscript{15} Reduction in testosterone levels cause tiredness, weakness, disturbances of the mood, reduction of libido, and sexual performance.\textsuperscript{36} The effect of methadone and anti-androgen drugs on sexual performance are similar.\textsuperscript{27} On the other side, buprenorphine, a partial opioid agonist of the μ receptor and an antagonist for the κ opioid receptor, causes the release of dopamine and does not inhibit the sex hormones to the same degree that the methadone does.\textsuperscript{28}

As our knowledge, our study is the first that investigated sexual adverse effects of opium tincture in patients undergoing maintenance treatment. Our results showed that sexual dysfunction in OMT group increased significantly during 3 months of study and the final total score of ASEX was higher than BMT group and lower than MMT group.

Morphine is the main substance of opium tincture that may be the important reason for sexual dysfunction in OMT group. Animal studies show that the misuse of morphine remarkably lessens testosterone levels.\textsuperscript{29} Long-term use of morphine affects luteinizing hormone (LH), testosterone, and body weight in rats, with no notable changes in follicle-stimulating hormone (FSH) and testicular weight.\textsuperscript{30} Cicero et al. stated a decrease in testicular and seminal vesicle weight.\textsuperscript{31} Morphine effects on the measures of hypothalamic monoamines has been shown in rats.\textsuperscript{30,32} Gabriel et al. showed the suppression of GnRH by morphine and testosterone.\textsuperscript{32}

Ahmadnia et al. showed the reduction of sex hormonal features and spermatogenesis, LH levels, and mature sperms of the target group.\textsuperscript{33}

In human studies also evidence showed that long-term use of morphine is associated with sexual dysfunction.\textsuperscript{34} Ajo et al. showed that sexual dysfunction is higher in men who received a significantly higher mean of opioid.\textsuperscript{35}

In subtest analysis, there was no remarkable difference in sexual dysfunction between three groups. Not only biological issues determine sexual desire but interpersonal (existence of partner) and social issues influence psychological part of the sexual desire.\textsuperscript{36} Orgasm means a quick release of sexual excitement during a sexual cycle starting with the rhythmic contractions of the pelvic muscles which is known as sexual pleasure.

To determine orgasm, medical doctors point out physiological changes but psychologists and psychiatrists point out spiritual and cognitive changes.\textsuperscript{37}

In the current study, we excluded all patients who had a history of psychiatric illness, and treated with medication during the study to exclude psychological effects on sexual function. Marital status has a remarkable role in the sexual functioning of patients during maintenance treatment. Some patients continued single without a sexual partner because of orgasmic complications in the study of Chekuri et al.\textsuperscript{38} Yee et al. found that men with no sexual partner on MMT had greater orgasmic complications than men

\begin{table}
\centering
\caption{Comparison of the mean Arizona sexual experiences scale (ASEX) scores of single patients in study groups after 3 months of therapy} \label{tab:6}
\begin{tabular}{|l|c|c|c|c|}
\hline
\textbf{Domain} & \textbf{MMT} & \textbf{BMT} & \textbf{OMT} & \textbf{P}^* \\
\hline
Desire/drive & 3.4 ± 1.1 & 3.4 ± 0.7 & 3.6 ± 1.1 & 0.86 \\
Arousal & 4.4 ± 1.1 & 4.5 ± 1.1 & 4.5 ± 1.0 & 0.93 \\
Erection & 3.5 ± 1.3 & 3.8 ± 1.2 & 3.8 ± 0.8 & 0.80 \\
Ability to reach orgasm & 4.0 ± 1.1 & 4.7 ± 1.1 & 4.4 ± 0.9 & 0.76 \\
Satisfaction with orgasm & 3.3 ± 1.5 & 3.6 ± 0.9 & 4.6 ± 1.2 & 0.08 \\
Total & 19.0 ± 5.1 & 19.8 ± 4.1 & 20.4 ± 4.0 & 0.82 \\
\hline
\end{tabular}
\footnotesize{The amounts are presented as mean ± standard deviation (SD). ASEX: Arizona sexual experiences scale; MMT: Methadone maintenance treatment; BMT: Buprenorphine maintenance treatment; OMT: Opium tincture maintenance treatment.} *Based on the one-way ANOVA test.
\end{table}
with no sexual partner on BMT. In a study by Ramdurg et al. sexual partner had a significant effect on orgasm problems in men underwent BMT.

Low serum testosterone, due to opioid effects on the hypothalamic-pituitary-gonadal axis, may explain libidinal depression. However, as psychological factors are common causes of depression of sex drive, and because psychiatric comorbidity is so prevalent in the substance-dependent population, mental and emotional health should be investigated in addition to hormonal assays. Conditions of potential importance include mood disorders, psychosis, situational stressors, gender identity issues, and age-related psychological issues. Medications other than opioid substitution treatment should also be reviewed, as these are also common causes of a depressed sex drive. Common offenders include anti-hypertensive and psychotropic agents.

Other etiologies should be ruled out; given the associations in the literature for the dose of methadone and serum testosterone, reasonable therapeutic approaches may include replacement (parenteral or transdermal) of irregular low testosterone or a reduction in daily methadone dose. In an open-label study, methadone-maintained men with depressed testosterone levels responded to transdermal testosterone in terms of serum testosterone levels, sexual function, and measures of well-being. Bromocriptine may be a therapeutic alternative, as well. Bromocriptine may act via reestablishment of central nervous system (CNS) levels of dopamine and normalization of dopaminergic regulation of prolactin production.

ED usually has an organic or iatrogenic origin. Non communicative diseases such as diabetes mellitus, chronic liver disease, renal failure, chronic pulmonary disease, cardiovascular disease, or malignancy can cause ED. Surgery, trauma, and congenital and anatomic anomalies in genitourinary can also lead to ED. Medications commonly associated with ED include antihypertensive, psychotropic agents, and anticholinergic drugs. Smoking has a strong correlation with ED. Every 10 pack-year of smoking enhances the relative danger for ED by 1.31. Mental and spiritual health problems may have a remarkable role. Symptoms of depression have strongly been correlated with ED; as 90% of men with severe depression showed ED in one research. Anxiety disorders has also been reported to have a correlation with ED.

In our study, there was no significant difference in subtest scores between groups. This finding can have two reasons. First, no remarkable difference was found between the number of single and married individuals in the investigated groups. Second, patients in the group of single patients may not have the reality of having a sexual partner due to social and religious issues.

Two main restrictions of our study was first, response bias related to sex privacy among patients, and the difficult feeling to talk about them with the researcher.

Second, LH and FSH were not studied in this research due to financial limitations. Previous studies show patients on MMT and OMT have reduced testosterone measures in comparison with patients on BMT; this is why sexual dysfunction is more in patients on MMT.

In brief, for rehabilitation of sexual performance is important to enhance the quality of life. Although a few studies exist assessing safety and efficacy of OMT in treating opioid use disorder with promising results, especially for detoxification, results of this study showed that use of opium tincture is associated with acceptable sexual dysfunction in compare with buprenorphine and methadone in opioid-dependent patients. Thus, future clinical trials are required to provide adequate evidence about the risks and success of OMT in opioid reliance treatment, particularly in long-term maintenance therapy.

Conclusion

These results showed that sexual dysfunction became better after opioid substitution therapies, and no differences had been observed on sexual dysfunction between three groups.

Conflict of Interests

The Authors have no conflict of interest.

Acknowledgements

We appreciate the support of Taleghani Hospital Research Development Committee to carry out this research project.
References

1. United Nations Office on Drugs and Crime (UNODC). World Drug Report 2016 [Online]. [cited 2016 May]; Available from: URL: https://www.unodc.org/doc/wdr2016/WORLD_DRUG_REPORT_2016_web.pdf

2. Chen W, Li X, Li X, Ling L, Xia Y, Chen J, et al. Erectile dysfunction among male heroin addicts receiving methadone maintenance treatment in Guangdong, China. J Addict Med 2012; 6(3): 212-8.

3. Hanbury R, Cohen M, Stimmel B. Adequacy of sexual performance in men maintained on methadone. Am J Drug Alcohol Abuse 1977; 4(1): 13-20.

4. Xia Y, Zhang D, Li X, Chen W, He Q, Jahn HJ, et al. Sexual dysfunction during methadone maintenance treatment and its influence on patient's life and treatment: A qualitative study in South China. Psychol Health Med 2013; 18(3): 321-9.

5. Ramdurg S, Ambekar A, Lal R. Sexual dysfunction among male patients receiving buprenorphine and naltrexone maintenance therapy for opioid dependence. J Sex Med 2012; 9(12): 3198-204.

6. Trajanovska AS, Vujovic V, Ignjatova L, Janicevic-Ivanovska D, Cibisev A. Sexual dysfunction as a side effect of hyperprolactinemia in methadone maintenance therapy. Med Arch 2013; 67(1): 48-50.

7. Sarkhel S. Kaplan and Sadock's Synopsis of Psychiatry: Behavioral Sciences/Clinical Psychiatry, 10th edition. Indian J Psychiatry 2009; 51(4): 331.

8. Maremmani I. The principles and practice of methadone treatment. Pisa, Italy: Pacini Editore Medicina; 2012. p. 69.

9. Brotto LA, Chik HM, Ryder AG, Gorzalka BB, Seal BN. Acculturation and sexual function in asian women. Arch Sex Behav 2005; 34(6): 613-26.

10. Walsh SL, Eissenberg T. The clinical pharmacology of buprenorphine: extrapolating from the laboratory to the clinic. Drug Alcohol Depend 2003; 70(2 Suppl): S13-S27.

11. Johnson RE, Chutuape MA, Strain EC, Walsh SL, Stitzer ML, Bigelow GE. A comparison of levomethadyl acetate, buprenorphine, and methadone for opioid dependence. N Engl J Med 2000; 343(18): 1290-7.

12. Heidari Z, Mahmoudzadeh-Sagheb H, Kohan F. A quantitative and qualitative study of rat testis following administration of methadone and buprenorphine. Int J High Risk Behav Addict 2012; 1(1): 14-7.

13. Quaglio G, Lugoboni F, Pattaro C, Melara B, Mezzelani P, Des Jarlais DC. Erectile dysfunction in male heroin users, receiving methadone and buprenorphine maintenance treatment. Drug Alcohol Depend 2008; 94(1-3): 12-8.

14. Yee A, Danaee M, Loh HS, Sulaiman AH, Ng CG. Sexual dysfunction in heroin dependents: a comparison between methadone and buprenorphine maintenance treatment. PLoS One 2016; 11(1): e0147852.

15. Hallinan R, Byrne A, Agho K, McMahon CG, Tynan P, Attia J. Hypogonadism in men receiving methadone and buprenorphine maintenance treatment. Int J Androl 2009; 32(2): 131-9.

16. Maguet O, Majeed M. Implementing harm reduction for heroin users in Afghanistan, the worldwide opium supplier. Int J Drug Policy 2010; 21(2): 119-21.

17. Alam Mehrjerdi Z, Zarghami M. Maintenance therapy with opium tincture for injecting drug users; implications for prevention from viral infections. Hepat Mon 2013; 13(4): e8334.

18. Jittiwutikarn J, Ali R, White JM, Bochner F, Somnogyi AA, Foster DJ. Comparison of tincture of opium and methadone to control opioid withdrawal in a Thai treatment centre. Br J Clin Pharmacol 2004; 58(5): 536-41.

19. Momtazi S, Noroozi A, Rawson RA. An Overview of iran drug treatment and harm reduction programs. In: el-Guebaly N, Carra G, Galanter M, editors. Textbook of addiction treatment: International perspectives. Milano, Italy: Springer Milan; 2015. p. 543-54.

20. Brown R, Balousek S, Mundt M, Fleming M. Methadone maintenance and male sexual dysfunction. J Addict Dis 2005; 24(2): 91-106.

21. Pezeshki MZ, Bayrami R. Reliability and construct validity of Arizona Sexual Experiences Scale (ASEX) among pregnant women referred to Tabriz urban health centers, 2004. Proceedings of 2nd National Congress on Family and Sexual Problems; 2005 Oct 25-26; Tehran, Iran.

22. Hassanipour Azgomi S, Arab Zozani M, Maghsoudi A, Mokhtari AM, Ghadiri Rad R. Comparing sexual dysfunction in maintenance therapy with Methadone and Buprenorphine in married male. International Journal of Epidemiologic Research 2016; 3(3): 232-8.

23. Tafreshian S, Javadi M, Fakhraei F, Fatemi SS. Sexual dysfunction in male patients receiving methadone and buprenorphine maintenance treatment in Iran. Heroin Addiction and Related Clinical Problems 2014; 16(3): 49-54.

24. Yee A, Loh HS, Hisham Hashim HM, Ng CG. The prevalence of sexual dysfunction among male patients on methadone and buprenorphine treatments: A meta-analysis study. J Sex Med 2014; 11(1): 22-32.

25. Galea S, Vlahov D. Social determinants and the
health of drug users: Socioeconomic status, homelessness, and incarceration. Public Health Rep 2002; 117(Suppl 1): S135-S145.

26. Bawor M, Bami H, Dennis BB, Plater C, Worster A, Varenbut M, et al. Testosterone suppression in opioid users: A systematic review and meta-analysis. Drug Alcohol Depend 2015; 149: 1-9.

27. Gladue BA, Clemens LG. Flutamide inhibits testosterone-induced masculine sexual behavior in male and female rats. Endocrinology 1980; 106(6): 1917-22.

28. Nimroozi R, Mastropietro MJ, Omidia H. Opioid abuse and deterrence: Buprenorphine products. J Develop Drugs 2014; 3: 1.

29. Cicero TJ, Wilcox CE, Bell RD, Meyer ER. Acute reductions in serum testosterone levels by narcotics in the male rat: stereospecificity, blockade by naloxone and tolerance. J Pharmacol Exp Ther 1976; 198(2): 340-6.

30. Yilmaz B, Konar V, Kutlu S, Sandal S, Canpolat S, Gezen MR, et al. Influence of chronic morphine exposure on serum LH, FSH, testosterone levels, and body and testicular weights in the developing male rat. Arch Androl 1999; 43(3): 189-96.

31. Cicero TJ, Adams ML, Giordano A, Miller BT, O'Connor L, Nock B. Influence of morphine exposure during adolescence on the sexual maturation of male rats and the development of their offspring. J Pharmacol Exp Ther 1991; 256(3): 1086-93.

32. Gabriel SM, Clark JT, Kalra PS, Kalra SP, Simpkins JW. Chronic morphine and testosterone treatment: effects on norepinephrine and serotonin metabolism and gonadotropin secretion in male rats. Brain Res 1988; 447(1): 200-3.

33. Ahmadnia H, Akhavan RA, Hoseyni M, Sharifi N, Khajedalooee M, Akhavan RA. Short-period influence of chronic morphine exposure on serum levels of sexual hormones and spermatogenesis in rats. Nephrourol Mon 2016; 8(4): e38052.

34. Ajo R, Segura A, Inda MD, Margarit C, Ballesta P, Martinez E, et al. Erectile dysfunction in patients with chronic pain treated with opioids. Med Clin (Barc) 2017; 149(2): 49-54.

35. Ajo R, Segura A, Inda MM, Planelles B, Martinez L, Ferrandez G, et al. Opioids increase sexual dysfunction in patients with non-cancer pain. J Sex Med 2016; 13(9): 1377-86.

36. Levine SB. The nature of sexual desire: A clinician's perspective. Arch Sex Behav 2003; 32(3): 279-85.

37. Rowland D, McMahon CG, Abdo C, Chen J, Jannini E, Waldinger MD, et al. Disorders of orgasm and ejaculation in men. J Sex Med 2010; 7(4 Pt 2): 1668-86.

38. Chekuri V, Gerber D, Brodie A, Krishnadas R. Premature ejaculation and other sexual dysfunctions in opiate dependent men receiving methadone substitution treatment. Addict Behav 2012; 37(1): 124-6.

39. Deglon JJ, Martin JL, Imer RL. Methadone patients' sexual dysfunctions: Clinical and treatment issues. Heroin Add & Rel Clin Probl 2004; 6(3): 17-26.

40. Shinderman MS, Maxwell S. Sexual dysfunction associated with methadone maintenance: Treatment with bromocryptine. Heroin Add & Rel Clin Probl 2000; 2(1): 9-14.

41. Kandeel FR, Kousa VK, Swerdlow RS. Male sexual function and its disorders: physiology, pathophysiology, clinical investigation, and treatment. Endocr Rev 2001; 22(3): 342-88.

42. Feldman HA, Goldstein I, Hatzichristou DG, Krane RJ, McKinlay JB. Impotence and its medical and psychosocial correlates: results of the Massachusetts Male Aging Study. J Urol 1994; 151(1): 54-61.

43. Rosen MP, Greenfield AJ, Walker TG, Grant P, Dubrow J, Bettmann MA, et al. Cigarette smoking: An independent risk factor for atherosclerosis in the hypogastric-cavernous arterial bed of men with arteriogenic impotence. J Urol 1991; 145(4): 759-63.

44. Araujo AB, Johannes CB, Feldman HA, Derby CA, McKinlay JB. Relation between psychosocial risk factors and incident erectile dysfunction: prospective results from the Massachusetts Male Aging Study. Am J Epidemiol 2000; 152(6): 533-41.

45. Sbrocco T, Weisberg RB, Barlow DH, Carter MM. The conceptual relationship between panic disorder and male erectile dysfunction. J Sex Marital Ther 1997; 23(3): 212-20.
مقایسه تأثیرات درمان‌های نگهدارنده با متادون، بوپرنورفین و تنتر اپیوم بر عملکرد جنسی

علی خردمند، ۱  احمد فاضلی، ۲ آزاده مظاهری میبدی، ۱

چکیده

مقدمه: درمان‌های نگهدارنده با متادون و بوپرنورفین، از جمله روش‌های رابط در درمان نگهدارنده مصرف مواد مخدر می‌باشند. هدف از انجام پژوهش حاضر، بررسی تأثیرات درمان نگهدارنده با متادون، بوپرنورفین و تنتر اپیوم بر عملکرد جنسی و مقایسه تأثیرات این سه درمان با هم در عملکرد جنسی بود.

روش‌ها: ۸۴ بیمار مرد که به صورت تصادفی از مراکز درمان سوء مصرف مواد شهر تهران در سال ۱۳۹۶-۱۳۹۷ انتخاب شده بودند، به سه گروه (هر گروه ۲۸ نفر) تقسیم شدند. جهت یک گروه درمان نگهدارنده با متادون، گروه دیگر بوپرنورفین و گروه سوم تنتر اپیوم شروع گردید. برای این بیماران، پرسشنامه تجربیات جنسی (Arizona Sexual Experiences Scale) در ابتدای درمان و سه ماه بعد برای تمام بیماران تکمیل شد.

یافته‌ها: نتایج نشان‌داد که درمان‌های ارائه شده به‌طور کلی عملکرد جنسی بالاتری نداشتند و تفاوت معنی‌داری بین سه گروه در نمرات مقیاس ASEX وجود نداشت. اما تفاوت معنی‌داری در نمرات سه ماه بعد از درمان هر گروه مشاهده شد.

نتیجه‌گیری: درمان‌های درمان نگهدارنده در بیماران مشخص مصرف مواد مخدر، همگی عملکرد جنسی را بهبود می‌بخشند. ولی نوع روش درمانی تأثیری در عملکرد جنسی بیماران ندارد.

واژگان کلیدی: متادون، بوپرنورفین، تنتر اپیوم، اختلال عملکرد جنسی، درمان‌های جایگزین مواد مخدر

ارجاعات: خردمند علی، فاضلی احمد، مظاهری میبدی آزاده. مقایسه تأثیرات درمان‌های نگهدارنده با متادون، بوپرنورفین و تنتر اپیوم بر عملکرد جنسی. مجله اعتیاد و سلامت ۱۳۹۸؛ ۱۱ (۲): ۱۲۸-۱۳۸.

تاریخ دریافت: ۱۳۹۷/۹/۹

تاريخ پذیرش: ۱۳۹۷/۱۱/۱۶

Email: a.alikheradmand@sbmu.ac.ir

 DOI: http://dx.doi.org/10.22122/ahj.v11i2.232
Published by Vesnu Publication

http://ahj.kmu.ac.ir, 04 April