Pain in methadone patients: Time to address undertreatment and suicide risk (ANRS-Methaville trial)

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Abstract

Background

Pain in opioid-dependent patients is common but data measuring the course of pain (and its correlates) using validated scales in patients initiating methadone treatment are sparse. We aimed to assess pain and its interference in daily life, associated correlates, and undertreatment before and during methadone treatment.

Methods

This is a secondary analysis using longitudinal data of a randomized trial comparing two methadone initiation models. We assessed the effect of methadone initiation and other correlates on pain intensity and interference (using the Brief Pain Inventory) at months 0, 6 and 12 using a mixed multinomial logistic regression model.

Results

The study group comprised 168 patients who had data for either pain intensity or interference for at least one visit. Moderate to severe pain was reported in 12.9% of patients at M0, 5.4% at M6 and 7.3% at M12. Substantial interference with daily functioning was reported in 36.0% at M0, 14.5% at M6 and 17.1% at M12. Of the 98 visits where patients reported moderate to severe pain or substantial interference, 55.1% reported no treatment for pain relief, non-opioid analgesics were reported by 34.7%, opioid analgesics by 3.1% and both opioid and non-opioid analgesics by 7.1%. Methadone was associated with decreased pain intensity at 6 months (OR = 0.29, p = 0.04) and 12 months (OR = 0.30, p = 0.05) of follow-up and tended to be associated with substantial pain interference. Suicide risk was associated with both pain intensity and pain interference.
Conclusions
Methadone in opioid-dependent patients can reduce pain. However, undertreatment of pain in methadone patients remains a major clinical concern. Patients with pain are at higher risk of suicide. Adequate screening and management of pain in this population is a priority and needs to be integrated into routine comprehensive care.

Introduction
Methadone is a long-acting opioid agonist prescribed to treat opioid dependence and pain as a second-line treatment or in opioid rotation [1,2]. Prevalence of pain among patients on Methadone Maintenance Treatment (MMT) is estimated at 29–80% [3–6].

Among MMT patients, pain is associated with older age [4,7], a high number of comorbidities [6–11] and psychiatric disorders [6,7,11,12] including severe depressive symptoms [8,10,12]. Prescription and non-prescription medication use [3,4,6,9]—including prescription opioids [6,8] and benzodiazepines [4]—have also been associated with pain. Furthermore, some studies have found that pain is associated with poorer treatment outcomes such as increased use of non-prescribed opioids [9,11,13], persistence of substance use after detoxification [13] and lower retention during maintenance treatment (methadone and buprenorphine) [14].

Although one would expect methadone to reduce pain among opioid-dependent patients, given that it is a powerful analgesic, to our knowledge, no study has assessed the course of pain (and its correlates) before and during MMT using a validated scale [15].

Clinicians treating opioid-dependent patients are often faced with the complex challenge of treating pain in patients with a history of opioid-dependence [16,17]. A recent study showed that despite the high prevalence of pain among people who inject drugs, clinicians may be reluctant to prescribe opioid-based analgesics to those with a history of drug use or addiction, especially in people already enrolled in MMT [17].

We used data from the ANRS Methaville trial to 1) assess the impact of MMT initiation and other correlates on two dimensions of pain—intensity and interference—using the BPI [18] in opioid-dependent patients, 2) estimate undertreatment in this population.

Materials and methods
Study design
The ANRS Methaville study is a pragmatic multi-site, open-label, randomized and controlled, non-inferiority trial, which compares methadone induction in France in specialized centers (hereafter “standard care”) with methadone induction by primary care physicians. Two previous articles describe the complete protocol [19] and the primary outcomes [20]. The study was approved by the Ethics Committee for the Protection of Patients in Paris, France.

Study group
From January 2009 to January 2010, 195 men and women were recruited in 10 sites and followed up for 12 months. Inclusion criteria were as follows: over 18 and under 70 years old, opioid-dependent in accordance with the DSM-IV criteria, and having an indication for methadone treatment (i.e. being methadone naive or not having had methadone treatment during the previous 30 days or switching from buprenorphine treatment). Exclusion criteria were as...
follows: having a triple (opioid, benzodiazepine and alcohol) dependence and not being reachable by phone for interview. All patients who agreed to participate in the study provided written informed consent. Subjects were identified only by number.

Data collection

Each participant was followed up for 12 months during which he/she had 4 programmed medical visits: at enrolment and at months 3, 6 and 12 (M0, M3, M6 and M12, respectively). At each visit, a medical questionnaire and a short self-administered questionnaire were completed. A phone interview (Computer-Assisted Telephone Interview: CATI) was conducted after each visit.

Pain intensity, interference with daily functioning and pain treatments. We assessed pain intensity and interference (i.e. the repercussions of pain on daily functioning) using the self-administered BPI-Short Form [21] at enrolment (before methadone initiation), at M6 and at the end of the study (M12).

Pain intensity was measured using an 11-point visual analogue scale from 0 (no pain) to 10 (worst possible pain). Patients used this scale to record pain “right now,” average pain in the previous 24 hours, and their highest and lowest pain levels during the previous 24 hours. A pain intensity score was calculated when patients completed all four related questionnaire items. It equaled the average of the four different scores. We used cut-off points employed by Serlin et al., 1995 [22]. A pain rating of 0 corresponded to “no pain”, >0 and <5 to “mild pain”, and ≥5 to “moderate to severe pain”.

Seven additional questions on a 10-point visual scale (0: does not interfere, 10: completely interferes) evaluated pain interference in the previous 24 hours with the following seven items: general activity, mood, walking ability, normal work, relations with other people, sleep, and enjoyment of life. An interference score was obtained by averaging the scores of the interference items when at least four of the seven items were completed, as suggested by Cleeland, 1994 [18]. We used statistical cut-off points (25th and 75th interquartiles) as, to our knowledge, no standard cut-off points exit for the interference score. A rating of 0 corresponded to “no interference”, >0 and <4 to “mild interference”, and ≥4 to “substantial interference”.

The BPI-short Form included a question about treatments taken against pain. Analgesics were classified as follows: non-opioid analgesics, opioid analgesics and combination of non-opioid and opioid analgesics.

Socio-demographic characteristics. At enrolment, information on socio-demographic characteristics including gender, age, educational level, employment status, having child(ren) or not, and housing situation (renter or owner of their personal housing, living with family, living in a hospital or clinic, in a social care institution or emergency accommodation, at a friend’s home, in a hotel, and finally, being homeless) was collected. At the M12 visit, employment status and housing situation were reassessed.

Medication use and addictive practices. After each follow-up visit, drug use in the previous month was assessed using a section of the Opiate Treatment Index (OTI) questionnaire focusing on drug use [23]. Individuals who reported using non-prescribed opioids (i.e., illicit opioids or opioid medication obtained without a prescription), cocaine and cannabis at least once in the previous month, were defined as non-prescribed opioid, cocaine and cannabis users, respectively.

At each follow-up visit, patients who reported drug injection at least once during the previous month were defined as active injectors. Patients reporting drug injection at least once during their lifetime were defined as having a history of drug injection.
We recorded anxiolytic/hypnotic agent use as positive when at least one of the following data sources indicated their use: medical questionnaire, self-report and urine test. Alcohol consumption was assessed at M0 and M12 during the CATI using the AUDIT questionnaire, with a cut-off point higher than 13 identifying alcohol dependence [24]. Binge drinking was defined as drinking 6 glasses or more of alcohol on one occasion at least once a month. At M0 and M6, nicotine dependence was screened for using the Fagerström test for nicotine dependence, a score ≥6 indicating high/very high dependence [25].

Daily prescribed methadone doses were recorded at M6 and M12. During each medical visit, physicians collected data on withdrawal symptoms using the Objective Opioid Withdrawal Scale (OOWS) which comprises a list of 13 withdrawal symptoms [26].

The self-administered questionnaire at M0, M6 and M12 included 2 screening tools. The first was the Adult ADHD (Attention Deficit Hyperactivity Disorder) Self-Report Scale-Version 1.1 (ASRS-V1.1). The scoring algorithm used was the summed score obtained after adding the scores (0–4) of the first 6 items of the adult ADHD scale [27]. We defined ADHD diagnosis using 14 as a cut-off point [28]. The second tool was a 20-item self-reported inventory, the Beck hopelessness scale (BHS), where a score of 9 or more indicates suicide risk [29].

During CATI, depressive symptoms were assessed at M0, M6 and M12 using the Center for Epidemiologic Studies Depression (CES-D) scale. The validated French cut-off points of 17 for men and 23 for women was employed [30].

**Statistical analysis**

We used a mixed multinomial logistic regression model to test the following factors for association with pain intensity (mild versus no pain, and moderate to severe versus no pain), and interference with daily functioning (mild versus no interference and substantial versus no interference) at M0, M6 and M12: 1) Follow-up characteristics: follow-up time (i.e. time on methadone: M6 versus M0 and M12 versus M0), methadone initiation (specialized centers versus primary care physician); 2) Socio-demographic characteristics: gender, age, educational level (< high school certificate versus ≥ high school certificate), having child(ren), employment status, stable housing (i.e. renter or owner of their personal housing versus other), receiving food assistance; 3) Addictive practices and medication use: methadone dose, anxiolytic/hypnotic agents, cocaine, non-prescribed opioids, and cannabis consumption, current drug injection, alcohol dependence, binge drinking, nicotine dependence; 4) History of drug use: switching from buprenorphine to methadone, history of drug overdose, history of drug injection, time since first regular drug use; 5) Health conditions: number of withdrawal symptoms, ADHD, current major depressive episode and suicide risk.

Variables associated with pain intensity and pain interference severity with a p-value lower than 0.20 in the univariable analysis for at least one category of the outcome were considered eligible to enter multivariable models. A stepwise procedure was used to identify the best model by removing variables one at a time based on a p-value of >0.05 for at least one of the outcome’s categories. When coefficient estimates for different categories of pain and interference were similar in the multivariable analysis, the categories were collapsed together to obtain a single estimate.

All analyses were performed using Intercooled Stata 12 (StataCorpLP, College Station, TX) software packages. The GLAMM Stata procedure was used to create a multinomial distribution.
Results

Description of the study group characteristics

The study group included 168 individuals with a score for pain intensity or interference at least once over the follow-up visits (M0, M6 or M12). No statistical difference was found between included (n = 168) and excluded (n = 27) patients regarding the following sociodemographic variables: age, gender, employment, housing, having children and educational level.

Among the 168 patients of the study group, 14% were female, aged 33.2 years on average (median = 32, interquartile range = [32.1–34.4]) and 31% had high school certificates. At M0, 52% were employed, 32% had unstable housing and 39% had a child(ren). Use of anxiolytic/hypnotic agents was reported by 24% of patients, cocaine by 29%, non-prescribed opioids by 72% and daily cannabis by 83%. Thirty percent had injected drugs in the previous month while 14% were alcohol dependent. Half the study group was switching from buprenorphine maintenance at trial inclusion, and 47% had a history of injection.

Pain intensity

At M0, M6 and M12, respectively, 21.6%, 36.9% and 31.3% patients reported no pain. Moderate to severe pain was reported by 12.9% of patients at M0, 5.4% at M6 and 7.3% at M12. Average score of pain intensity varied from 2.4 (SD = 2.0) at M0 to 1.3 (SD = 1.5) at M6 and 1.5 (SD = 1.8) at M12.

In univariable analysis, mild and moderate to severe pain (versus no pain) were negatively associated with MMT at 6 and 12 months (Table 1).

Results of the multivariable multinomial regression for mild pain and moderate to severe pain (versus no pain) were similar. For this reason, the multivariate analysis of the dependent variable pain versus no pain is presented in Table 1. This analysis showed that after adjustment for history of injection drug use and withdrawal symptoms, MMT was negatively associated with pain at 6 months (OR [95%CI] = 0.29 [0.09–0.92]; p = 0.04) and after 12 months (OR [95%CI] = 0.30 [0.09–0.97]; p = 0.05), while suicide risk was positively associated with pain (OR[95%] = 5.90 [1.49–23.3], p = 0.01).

Interference with daily functioning

At M0, 24.5% of the study group reported no pain interference. This increased to 32.1% at M6 and 36.9% at M12. At M0, 36.0% of the patients reported substantial interference, 14.5% at M6 and 17.1% at M12. The average score of interference varied from 3.0 (SD = 2.6) at M0 to 1.7 (SD = 2.0) at M6 and 1.7 (SD = 2.0) at M12.

The results of the univariable analysis are presented in Table 2.

After adjusting for potential confounding factors (withdrawal symptoms and stable housing), being at suicide risk (OR [95%CI] = 3.78 [1.40–10.21]; p = 0.01) was positively associated with substantial interference (Table 2). MMT at M6 and M12 tended to be associated with a decrease in substantial interference (respectively OR [95%CI] = 0.36 [0.13–1.02]; p = 0.06 and OR [95%CI] = 0.39 [0.13–1.14]; p = 0.09).

Pain treatment

At M0, among the 55 patients reporting moderate to severe pain with substantial interference, 30 declared receiving no analgesic treatment, 13 opioid analgesics and 12 non-opioid analgesics. At M6, 20 patients reported moderate to severe pain with substantial interference, among them 11 declared receiving no analgesic treatment and 9 non-opioid analgesics. At M12, 23
patients reported moderate to severe pain with substantial interference, among them 11 declared receiving no analgesic treatment and 12 non-opioid analgesics.

Discussion

There are 3 main results of this study: 1) pain decreases with methadone treatment; 2) pain is considerably undertreated in MMT patients; 3) suicide risk is a major correlate of pain intensity and pain interference with daily functioning.
Table 2. Factors associated with mild and substantial interference during methadone treatment: Univariable and multivariable logistic regressions (ANRS-Methaville trial; n = 168 patients/168 included, 389 visits/403 visits).

| Follow-up       | Univariable analysis | Multivariable analysis |          |          |          |          |          |          |          |          |          |          |
|-----------------|----------------------|------------------------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|
|                 | No. of visits        | No. of patients        | Mild versus no interference | Substantial versus no interference | Mild versus no interference | Substantial versus no interference |          |          |          |          |          |          |
|                 |                      |                       | OR [IC95%] | p        | OR [IC95%] | p        | OR [IC95%] | p        | OR [IC95%] | p        | OR [IC95%] | p        |
| M0              | 147 (37.8)           | 147 (87.5)             | 1        | 1        | 1        | 1        |          |          |          |          |          |          |
| M6              | 131 (33.7)           | 131 (78.0)             | 0.71 [0.33; 1.54] | 0.39    | 0.21 [0.09; 0.50] | < 0.001 | 0.63 [0.25; 1.61] | 0.34    | 0.36 [0.13; 1.02] | 0.06    |          |          |
| M12             | 111 (28.5)           | 111 (66.1)             | 0.49 [0.21; 1.10] | 0.08    | 0.20 [0.08; 0.48] | < 0.001 | 0.54 [0.21; 1.42] | 0.21    | 0.39 [0.13; 1.14] | 0.09    |          |          |
| Socio-demographic characteristic |          |                       |          |          |          |          |          |          |          |          |          |          |
| Having child (ren)<sup>a</sup> |          |                       |          |          |          |          |          |          |          |          |          |          |
| No              | 233 (60.5)           | 102 (61.5)             | 1        | 1        |          |          |          |          |          |          |          |          |
| Yes             | 152 (39.5)           | 64 (38.5)              | 0.48 [0.18; 1.30] | 0.15    | 0.30 [0.11; 0.86] | 0.03    |          |          |          |          |          |          |
| Stable housing<sup>b</sup> |          |                       |          |          |          |          |          |          |          |          |          |          |
| No              | 149 (38.4)           | 72 (43.1)              | 1        | 1        | 1        | 1        |          |          |          |          |          |          |
| Yes             | 239 (61.6)           | 116 (69.5)             | 0.34 [0.14; 0.82] | 0.02    | 0.37 [0.14; 0.93] | 0.03    | 0.38 [0.15; 0.98] | 0.05    | 0.41 [0.15; 1.11] | 0.08    |          |          |
| Medication use and addictive practices |          |                       |          |          |          |          |          |          |          |          |          |          |
| Binge drinking<sup>c</sup> |          |                       |          |          |          |          |          |          |          |          |          |          |
| No              | 103 (44.4)           | 79 (52.7)              | 1        | 1        |          |          |          |          |          |          |          |          |
| Yes             | 129 (55.6)           | 95 (63.3)              | 2.66 [1.08; 6.57] | 0.03    | 2.16 [0.84; 5.55] | 0.11    |          |          |          |          |          |          |
| Non-prescribed opioid consumption<sup>d</sup> |          |                       |          |          |          |          |          |          |          |          |          |          |
| No              | 187 (51.9)           | 103 (61.7)             | 1        | 1        | 1        | 1        |          |          |          |          |          |          |
| Yes             | 173 (48.1)           | 118 (70.7)             | 2.05 [0.91; 4.60] | 0.08    | 3.45 [1.45; 8.20] | 0.01    |          |          |          |          |          |          |
| Health indicators |          |                       |          |          |          |          |          |          |          |          |          |          |
| No. of withdrawal symptoms<sup>e</sup> |          |                       |          |          |          |          |          |          |          |          |          |          |
| No              | 227 (68.0)           | 119 (76.3)             | 1        | 1        |          |          |          |          |          |          |          |          |
| Yes             | 107 (32.0)           | 72 (46.2)              | 1.53 [0.66; 3.57] | 0.32    | 3.53 [1.45; 8.58] | 0.01    |          |          |          |          |          |          |
| Current suicide risk<sup>f</sup> |          |                       |          |          |          |          |          |          |          |          |          |          |
| No              | 271 (71.3)           | 133 (80.6)             | 1        | 1        | 1        | 1        |          |          |          |          |          |          |

(Continued)
At M0, 30 out of 55 patients reporting moderate to severe pain with substantial interference declared receiving no pain treatment, 11 out of 20 at M6 and 11 out of 23 at M12. This suggests that pain was inadequately treated, which is in line with other studies where pain of MMT patients was undertreated [31,32]. One explanation could be that clinicians may have deny additional analgesics to individuals already on methadone treatment [17]. Undertreatment of pain may have a negative effect on opioid dependence outcomes as many opioid maintenance treatment patients report frustration which encourages them to use illicit opioids for pain relief [17,33,34].

As shown by our results, methadone can reduce pain among opioid-dependent patients. However, its standard once-a-day administration for the treatment of opioid addiction often fails to provide sustained pain relief [35]. According to Blinderman et al., in order to treat pain, patients on MMT should receive additional methadone doses three to four times daily or every 6 to 8 hours [36]. A root cause of undertreated pain in MMT patients may be the tendency for clinicians to dichotomize MMT as either a treatment for pain or for addiction, rather than recognizing the comorbid nature of these two dimensions in individuals with a history of substance use [37]. Evidence-based guidelines have already been drawn up for pain management in opioid-dependent and MMT patients [38–40]. These guidelines should be put into clinical practice.

Suicide risk was significantly and independently associated with both pain intensity and interference in our study. Many factors may lead to suicide in pain patients. First, many experience concomitant depression [41]. Moreover, in such patients, depression may be misdiagnosed due to shared somatic symptoms between pain and depression [42]. This could explain the association between depression and pain intensity and interference found in unvariable analyses in our study (this was not confirmed in multivariable analysis, probably due to collinearity with suicide risk). Second, pain catastrophizing (exaggerated, negative focus on pain) can contribute to depression, pain intensity, disability [43] and mental defeat [44]. In fact, magnitude of depression and pain catastrophizing can predict the occurrence and degree of suicidal ideation [45]. Third, sleep disorders could mediate the relationship between pain and

### Table 2. (Continued)

|                        | Univariable analysis | Multivariable analysis |
|------------------------|----------------------|------------------------|
|                        | No. of visits | No. of patients | Mild versus no interference | OR [IC95%] | p | Substantial versus no interference | OR [IC95%] | p | Mild versus no interference | OR [IC95%] | p | Substantial versus no interference | OR [IC95%] | p |
| Yes                    | 109 (28.7)    | 71 (43.0)       | 2.37 [0.97; 5.81] | 0.06 | 3.81 [1.50; 9.72] | 0.01 | 2.21 [0.87; 5.67] | 0.10 | 3.78 [1.40; 10.21] | 0.01 |
| No                     | 286 (74.7)    | 142 (86.1)      | 1 | 1 | 1 | 1 | 1 | 1 |
| Yes                    | 97 (25.3)     | 66 (40.0)       | 1.60 [0.63; 4.08] | 0.32 | 4.07 [1.56; 10.64] | 0.004 | 1 | 1 |

* The percentages computed in this column do not add up to 100% as each refers to the proportion of individuals who had the characteristics at least once during follow-up. E.g.: 70.07% of patients reported they consumed non-prescribed opioids at least once during follow-up.

* c: At M0 and M12

* d: >6 glasses on the same occasion at least once a month

* e: During the previous 4 weeks

* f: CES-D score >17 for males and >23 for females

* g: During the previous 6 months

* h: Beck score > 9

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suicide attempt/ideation. Although not observed in the present study, the association between sleep disorders and pain and suicide risk was highlighted in a previous study using Methaville trial data [46]. Fourth, physical pain could lead to suicide attempt/ideation through its interaction with social pain (hopelessness, loss of work and changing family role, as well as isolation due to pain), as both types of pain may share some underlying neurological mechanisms [47]. Finally, through indicated and appropriate prescription of opioids, vulnerable patients may receive access to potentially lethal medications (i.e., opioids) [42].

Whether methadone can reduce suicide risk through pain relief is still unknown. A previous study using data from the Methaville trial showed that methadone and suicide risk were associated in univariable analysis but no longer associated after multiple adjustment. This is probably because of collinearity between suicide risk and the number of health problems, which are significantly reduced during methadone treatment [48]. Our results suggest the need for routine assessment of pain and suicide risk by physicians, particularly in opioid-dependent patients. This is particularly relevant for patients classified as polydrug users and those with harmful alcohol consumption, considering the relatively high frequency of suicide and suicide attempts—including voluntary overdoses—in people who use drugs, whether or not they are on methadone treatment.

ADHD was associated in univariable analyses with pain intensity and interference. A recent study showed that patients with ADHD are more likely to experience pain, and that while common mental disorder influences this association, it does not fully explain the reason for this [49].

The challenge of treatment decisions by physicians regarding pain in opioid-dependent populations is compounded by the conflicting literature on the association between pain and addiction. Although some previous surveys demonstrated a link between pain and drug use behaviors [7,10,13,50–52], the present study, like others [3,8,53,54], did not find that patients with pain have a higher risk of substance use. However, univariable analyses in our study showed that pain intensity and interference were associated with non-prescribed opioid use. Whether or not treating pain in opioid-dependent patients leads to a reduction in substance use remains an important issue for future research.

Our study has strengths and limitations. The main strengths are that as a pragmatic trial, the study population is representative of people seeking care for opioid dependence in France. Second, the assessment of pain is based on a validated scale, the BPI-short Form. In terms of limitations, the study is based on self-reports to measure medication and drug use. However, the reliability of self-reports in people who use drugs has already been widely demonstrated [55].

Conclusion

Methadone in opioid-dependent patients can reduce pain. However, undertreatment of pain in MMT patients remains a major clinical concern. Patients presenting pain and reporting interference with their daily life are at higher risk of suicide. Adequate screening and management of pain in this population is a priority and needs to be incorporated in routine comprehensive care.

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