Long-Term Follow-Up of Orally Administered Diacetylmorphine Substitution Treatment

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Key Words
Opioid dependence · Heroin substitution treatment · Oral application · Open-label study · Prospective cohort · Bias

Abstract

Background: To assess the long-term course of the feasibility and safety of orally administered heroin [diacetylmorphine (DAM)] tablets in substitution treatment of severely addicted opioid users. Design: Open-label, prospective cohort study with 2 non-randomly assigned treatment arms: DAM tablets only (n = 128) or DAM tablets combined with injected DAM and/or other opioids (n = 237). The average duration of the observation period was 62 months. Study endpoints were the time to discharge from treatment and the number of serious adverse events. Results: Both patient groups had a higher than 70% retention rate after the first 48 months of treatment, with similar long-term retention rates (after 8 years both groups had retention over 50%). The physician-verified rate of serious adverse events was 0.01 events per application year among the exclusively oral substitution group (intention-to-treat analysis) during the last year of observation, and 0.005 events per application year in the other group. Conclusions: Because of their feasibility and safety over years, DAM tablets may be a valuable long-term therapeutic alternative.

Introduction

Maintenance treatment, especially with methadone, has become the standard treatment for opioid dependence in numerous countries and has been recommended by international organizations [1–4]. Not all opioid-dependent patients, however, fully respond to methadone maintenance treatment [5]. The medical prescription of intravenous diacetylmorphine (DAM) to chronic treatment refractory heroin-dependent patients has been considered as a second-line therapy in several countries and its feasibility and effectiveness has been confirmed in cohort studies [6, 7] and controlled trials [8–10]. Patients who had never or not regularly injected had to be excluded from intravenous heroin substitution for ethical con-
oral morphine, leading to higher \( C_{\text{max}} \) and shorter \( t_{\text{max}} \). Absorption from oral DAM was found to be faster than from oral morphine. Therefore, oral DAM could be seen as an active pro-drug of morphine reaching a higher level of systemic morphine faster than from morphine itself [14].

As previously mentioned [16], the results of an intention-to-treat analysis of the current open-label prospective cohort study for DAM tablets within the Swiss HAMT after an observation period of 1 year have been reported. Data p.o. treatment was part of the overall Swiss HAMT that addressed the treatment needs of patients ineligible for DAM i.v. or who had or wanted to switch to DAM p.o. (see below for details). In the Frick et al. study [16], the retention rates after 1 year in DAM tablets-only group (0.804, 95% CI: 0.735–0.873) and in the subgroup combining oral application of DAM with intravenous application or other opioids (0.843, 95% CI: 0.797–0.889) were higher than in the historical controls, i.e. the Swiss cohort of patients that had been intravenously substituted with DAM (1-year retention rate = 0.70). In addition, rates of serious adverse events under study medication (tablets only = 0.038 per application year; tablets in combination = 0.028 per application year) were comparable to the historical rate of the Swiss heroin-assisted treatment (0.043).

The present analyses report the results for the feasibility and safety of oral DAM using the same study endpoints with prolonged observation period: for feasibility, the period between a patient’s individual start date of oral medication and July 2008 was evaluated; for safety, the first 2 years after the start of the formal drug approval study were used.

**Methods**

The study design was an open-label prospective cohort study with 2 non-randomly allocated treatment arms to be compared. Additionally, the pooled cohort of both treatment arms is compared in its long-term outcome (retention from first tablet to July 28, 2008) to the complete population of injecting DAM users in the Swiss HAMT (from individual treatment entry to the same time point).

**Eligibility**

Patients who were already admitted to the Swiss heroin-assisted treatment program (for general eligibility criteria for this treatment program for severely addicted opioid users, see [19]) were
eligible to be enrolled into this study if one or more of the following conditions were fulfilled: (1) the status of their veins prohibited further intravenous application of DAM, (2) they had used opioids prior to the substitution program exclusively via inhalation or sniffing, and intravenous use, therefore, meant an increase in health risk for them, or (3) they wished to reduce their health risk gradually or abruptly, but in any case permanently shifting from intravenous to the oral application of DAM.

All 402 patients already receiving DAM tablets in the Swiss heroin-assisted treatment, or scheduled to receive such tablets from the reference date of each treatment center, were enrolled in the study (fig. 1). Twenty-one of the 23 existing Swiss outpatient treatment centers recruited patients. For the remaining 2 centers, there were no patients fulfilling the inclusion criteria.

Treatment Allocation and Study Groups

All patients were treated according to Swiss heroin-assisted treatment protocol, as specified by and under permission of the Federal Office of Public Health on a patient-named basis. Treatment included psychosocial services at least once per month, and somatic interventions if necessary. Opiate dosage and number of administrations were prescribed according to the patient’s needs. The treatment centers were open 3 times per day, 7 days per week. In groups 1 and 2, no other substitution medication was given (for protocol violations, see below). Consumption was monitored by a self-reported questionnaire and urinalysis.

Groups 1 and 2 (Tablets Only). A total of 128 patients received a medication regimen providing DAM exclusively in tablet form [either immediate-release tablets alone (group 1) or combining immediate release and slow-release tablets (group 2)].

Group 3 (Tablets and Other Opioids). A combination of DAM tablets with either injectable DAM or other noninjectable opioids, like morphine (as a tablet), was given (≥1 dose) to 237 patients (group 3). These patients received intravenous applications only if there was clear evidence of intravenous use prior to the study. With regard to medications other than diamorphine, study group 3 was designed to be a ‘left over’ category, comprising all combinations of substitution-medication including combinations with methadone or morphine tablets.

At baseline, 26.5% of all patients in groups 1 and 2, but only 9.7% of group 3, had never practiced intravenous application of heroin or other opioids (χ² test; p < 0.001). With respect to variables predictive for treatment outcome in the cohort entering the Swiss HAMT program between January 1, 2001 and February 29, 2004 [20], no differences were found between patients receiving tablets and those receiving only intravenous applications of heroin.

Within the groups described, 25 patients in groups 1 and 2 (tablets only) and 49 patients in group 3 had not received DAM tablets prior to the start of the study. The remaining patients had some experience with DAM tablets as part of the Swiss heroin-assisted treatment. Therefore, this article uses the longest available observation period per patient for evaluation, and counts the duration of oral substitution treatment from the very first day that the patient received a DAM tablet.

Study Period

The feasibility of oral DAM medication was evaluated for each patient’s longest possible observation period, irrespective of the time when he/she was included into the formal drug approval study. Thus, the duration of the observation period was counted as the days from each patient’s first medication with DAM tablets until July 18, 2008 when the database was frozen. On average, patients in the tablets only groups had an observation period of 2,253 days (SD 627), while patients in group 3 (tablets and other opioids) had 2,237 days (SD 528). Analyses on adverse events were
performing for the second year after the start of the formal drug approval study (in most cases: January 2005 to January 2006) in order to compare the risk for serious adverse events per application year to the risk calculated for the observation period (November 2003 to December 2004) that was reported by Frick et al. [16].

Study Endpoints

The main endpoint of the 1-year study by Frick et al. [16] was the retention rate, which was to be compared to the known historical 1-year retention rate of the Swiss heroin substitution program, reported to be a proportion of 0.7 [6]. Sample size calculations were based on this comparison [16, 21]. Discharge from HAMT was defined as either an explicit ending of treatment by a respective declaration of the patient, formally excluding the patient from the treatment for disciplinary reasons according to the rules of HAMT

The main study endpoint of the prospective cohort study was the retention rate. Figure 3 shows the temporal course of this endpoint.

The retention rate 24 months after the individual start date of oral medication for groups 1 and 2 was 80.7% (95% CI: 73.6–87.8%), which was somewhat lower than the rate of group 3 (87.6%; 95% CI: 83.2–92.0%). For both groups,
however, these retention rates were significantly higher after more than 2 years than the 1-year retention rate in the historical sample of intravenous heroin-assisted treatment (70%).

At day 1,349 (3.7 years or 45 months after initial tablets medication) survival curves of both conditions intersected at a retention rate of 72.5% (95% CI for group 3: 67.0–78.8%). Thereafter, groups 1 and 2 displayed a somewhat more favorable retention rate until day 2,472 (after 82 months or 6.8 years). Then both survival curves intersected a second time at a rate of 56.0% (95% CI for groups 1 and 2: 44.4–67.7%). Groups 1 and 2 displayed a very stable treatment retention rate after this period lasting at 56.0% for more than 12 years (or 150 months) for the longest observed patients. Group 3 displayed a retention rate of 50.3% (95% CI: 39.0–61.4%) after 8.8 years (or 106 months), which remained stable to the last observation after 11.4 years (or 139 months). Both retention rates can be seen as remarkably high given the length of the treatment period. A formal log-rank test to compare the groups was not possible because of the nonproportional hazard rates. The Kolmogorov-Smirnov test for differences between the shapes of both survival curves yielded a value of p < 0.001. This indicates potential disadvantages for the tablets only medication regimen during the first 3 years which were outweighed by the advantages during the years 4–6; there were no meaningful differences in the long run after 6 years.

Figure 2 shows treatment retention for all DAM p.o. groups combined in comparison to DAM i.v. DAM p.o. was associated with longer treatment retention. After 12 years, the respective estimates for treatment retention were: DAM p.o. patients, 52.1% (95% CI at day 3,205: 44.6–59.5%); DAM i.v. patients, 15.7% (95% CI at day 4,373: 14.1–17.3%).

Retention in a maintenance study is often time-dependent, with the highest dropout rate occurring shortly after receiving a new study medication. As the majority of patients in our study had already been using the study medication before the start of the study, this may have resulted in self-selection bias. It may be possible that in the patients who had been using the study medication before, people at higher risk of dropout had already dropped out by the time the study began, and thus, the high retention rate was caused mainly by these long-term heroin tablet users. We tested this potential bias by comparing the two groups, i.e. those who had been on DAM p.o. before the study began and those who started with DAM p.o. on the first day of the study. Though the confidence intervals overlapped over the full study period (fig. 4), a formal log-rank test yielded a p value of 0.0326, pointing at a somewhat faster dropout of patients set on tablets in October 2003.

**Major Study Endpoint: Adverse Events**

The physician-verified rate of serious adverse events was 0.01 events per application year, or 1% of affected patients among the exclusively oral substitution group (ITT analysis) during the second year of observation (table 1, which also gives adverse events during first observation year). This rate was lower than during the first observation year. Combined medication strategies were associated with a rate of 0.005 events per application year (0.5% of affected patients), again lower than the first observation year. Neither group deviated significantly from the a priori fixed tolerance limit of 4.3% of affected patients.

The two serious events (death of one patient in India on study leave substituted by methadone; admission to psychiatric hospital for stabilization and cocaine withdrawal) were not causally related to the study medication. After January 2006 (end of second observation year), registration and monitoring of adverse events followed the routine procedures of Swiss HAMT and, thus, were not available for the purpose of this study.
Table 1. Incidence of serious adverse events

<table>
<thead>
<tr>
<th></th>
<th>Group 1 (only IR tablets)</th>
<th>Group 2 (IR + SR tablets)</th>
<th>Groups 1 and 2 combined</th>
<th>Group 3 (combination of tablets with opioid medication)</th>
<th>Total</th>
</tr>
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<tbody>
<tr>
<td>Observation period 1: November 1, 2003 to January 5, 2005</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Events</td>
<td>0</td>
<td>5</td>
<td>5</td>
<td>7</td>
<td>12</td>
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<tr>
<td>Patients with event</td>
<td>0</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>10</td>
</tr>
<tr>
<td>Patients under risk</td>
<td>40</td>
<td>88</td>
<td>128</td>
<td>237</td>
<td>365</td>
</tr>
<tr>
<td>Days under risk</td>
<td>15,999</td>
<td>32,434</td>
<td>48,433</td>
<td>90,608</td>
<td>139,041</td>
</tr>
<tr>
<td>Events per year</td>
<td>0.000</td>
<td>0.056</td>
<td>0.038</td>
<td>0.028</td>
<td>0.032</td>
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<tr>
<td>95% CI (Poisson) lower limit</td>
<td>0.000</td>
<td>0.019</td>
<td>0.012</td>
<td>0.011</td>
<td>0.016</td>
</tr>
<tr>
<td>95% CI (Poisson) upper limit</td>
<td>0.081</td>
<td>0.126</td>
<td>0.086</td>
<td>0.057</td>
<td>0.054</td>
</tr>
<tr>
<td>Rate of persons with serious events per year</td>
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<td>0.056</td>
<td>0.038</td>
<td>0.020</td>
<td>0.026</td>
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<tr>
<td>95% CI (Poisson) lower limit</td>
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<td>95% CI (Poisson) upper limit</td>
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<td>0.126</td>
<td>0.086</td>
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<tr>
<td>Mean observation time per patient, years</td>
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<td>1.00</td>
<td>1.03</td>
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<td>1.04</td>
</tr>
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</table>

Observation period 2: January 6, 2005 to January 5, 2006

<table>
<thead>
<tr>
<th></th>
<th>Group 1 (only IR tablets)</th>
<th>Group 2 (IR + SR tablets)</th>
<th>Groups 1 and 2 combined</th>
<th>Group 3 (combination of tablets with opioid medication)</th>
<th>Total</th>
</tr>
</thead>
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<tr>
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<td>1</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Patients with event</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Patients under risk</td>
<td>37</td>
<td>68</td>
<td>105</td>
<td>203</td>
<td>308</td>
</tr>
<tr>
<td>Days under risk</td>
<td>12,860</td>
<td>24,121</td>
<td>36,981</td>
<td>68,137</td>
<td>105,118</td>
</tr>
<tr>
<td>Events per year</td>
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<td>0.010</td>
<td>0.005</td>
<td>0.007</td>
</tr>
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<td>0.000</td>
<td>0.000</td>
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</tr>
<tr>
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<td>0.019</td>
</tr>
<tr>
<td>Rate of persons with serious events per year</td>
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<td>0.015</td>
<td>0.010</td>
<td>0.005</td>
<td>0.006</td>
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<tr>
<td>95% CI (Poisson) lower limit</td>
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<tr>
<td>95% CI (Poisson) upper limit</td>
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<td>0.051</td>
<td>0.033</td>
<td>0.017</td>
<td>0.017</td>
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<tr>
<td>Mean observation time per patient, years</td>
<td>0.95</td>
<td>0.97</td>
<td>0.96</td>
<td>0.92</td>
<td>0.94</td>
</tr>
</tbody>
</table>

Fig. 4. Retention rate of patients receiving DAM tablets prior to October 2003 (start of drug approval study) versus the rate since October 2003.
Discussion

Chronic dependent opioid users are characterized by the persistence of opioid use regardless of the difficulties they experience with their health, the law and in their social environment. DAM tablets offer an effective maintenance treatment for the specific subgroup of refractory and severely addicted opioid users who do not inject. Overall, DAM p.o. substitution has proven to be a safe and feasible treatment for severely opioid-dependent people. Thus, our results may be generalizable to a selection of patients willing to try oral administration. Safety was comparable to DAM i.v., which is already accepted as a regular medication by the Swiss authorities for the treatment of this group. Feasibility, as measured by treatment retention, was even significantly better in DAM p.o. substitution.

There may be alternative settings for nonintravenous DAM substitution. For instance, oral morphine has been suggested as a second-line substitution medication [12], and inhalable DAM has been tested in the Netherlands with success [11]. We hope that these and other alternatives will be tested against DAM p.o. in randomized trials. For inhalable DAM, different forms for inhalation from the Dutch may have to be found to be accepted as medication in some countries, but in principle this is possible. Our hypothesis would be that DAM p.o. would be more effective than other oral opioids including morphine, as the bioavailability is higher and a faster onset can be reached [18]. But these are hypotheses based on basic research; the effects have to be confirmed in randomized trials under real-life conditions.

There were some potential methodological limitations concerning the self-selection process of the participants in our study. A portion of our patients might have chosen oral application of DAM because their attitude towards risks associated with intravenous application had been changing and this change reflected a general process of dissociation from the illegal drug scene. This process could then have determined the low rate of dropouts as well as the low illegal consumption and the observed low rate of serious adverse events. This argument could only be countered by a randomized controlled clinical trial comparing intravenous versus oral application mode of DAM, which would be unethical.

We found hints on a faster dropout process of the 74 patients (18%) who started with tablets in October 2003. Whether this means the retention rates of the oral DAM groups are positively flawed by omitting potential former dropouts prior to our study cannot be concluded with certainty. It also seems plausible that the historical situation in autumn 2003 was understood by the treating physicians as a ‘last chance’ to change over the substitution medication of their patients to oral DAM for a long time period, and therefore they recruited in October 2003 a group of patients who were less suited for oral substitution than the previous subgroup.

It may also be argued that the reported retention was only possible because the patients were recruited mainly from patients already in maintenance treatment, thus excluding the high rate of dropouts in the initial treatment period [6, 10]. Three indirect arguments could be made: opioid addiction is a chronic relapsing disease and, thus, for many opioid-dependent patients, life has been a succession of episodes inside treatments and untreated episodes. Even people entering treatment after 1 year outside treatment have often had multiple treatment episodes before [10, 22], and vice versa: cohorts of originally untreated opioid users show a high prevalence of maintenance treatment episodes after some time [23]. The success of maintenance treatment seems to be relatively independent of prior history. A German trial did not find significant differences in effectiveness of heroin-assisted treatment between people already in methadone treatment or outside the treatment system [10]. Furthermore, in the Swiss study, the success of treatment, as defined by retention rate, was independent of whether it was a first, second or third treatment episode [24]. Therefore, we would hypothesize that the effectiveness of DAM p.o. is independent of whether it is a first treatment episode or if the patient enters this treatment from another treatment. Such a hypothesis must be further examined in new studies.

Finally, the quite ‘liberal’ definition of retention in the Swiss HAMT is also covering periods of methadone substitution during patients’ holidays, for instance. Therefore comparing these retention rates with stricter definitions from outside Switzerland might face a potential ‘overestimation’ of retention for Swiss patients. But this argument does not apply for comparisons within the Swiss HAMT.

Notwithstanding the methodological limitations mentioned above, the results of our study strongly support the continuation of this treatment for all patients with the mentioned indications for the oral application of HAMT. While other studies may clarify the optimal application of DAM tablets in combination with other opioids or in exclusive prescription, the current practice clearly seems justified by the clinical results.
Another open question concerns the application of oral DAM as a first-line treatment for refractory opioid users. Currently, intravenous application is the default for everybody, except for users of illegal opioids who do not inject. Given the results above, it should be considered whether the continuation of the status quo of providing a potentially more harmful treatment option (intravenous prescription) as a first-line treatment is justified. Of course, we would need more research with better design and including different endpoints before a truly evidence-based decision to make oral DAM a first-line treatment for refractory opioid users can be made.

Finally, we hope that our results with respect to DAM tablets will lead to a further ‘normalization’ of the role of HAMT within the continuum of care. Meanwhile diamorphine tablets received orphan drug status in Switzerland. In a time when substitution treatment finally has been accepted as the standard of evidence-based opioid treatment [1, 2], we see no reason why heroin tablets should be considered any different from other forms of opioid substitution. As already indicated above, further randomized trials using different types of opiates should be conducted to provide the necessary evidence of future guidelines.

References