Health and quality of life in patients with medication overuse headache syndrome after standardized inpatient rehabilitation

A cross-sectional pilot study

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Abstract

The aim of this pilot study was to determine health-related quality of life (HRQoL) in patients with history of medication overuse headache (MOH) after detoxification and a headache-specific inpatient rehabilitation program and to receive necessary information for future prospective studies.

HRQoL and headache-related disability were cross-sectionally measured by Short Form 36 (SF-36), Hospital Anxiety and Depression Scale (HADS), Migraine Disability Score (MIDAS), Coping Strategies Questionnaire (CSQ), and Symptom Checklist 90 revised (SCL-90-R). SF-36, HADS, and SCL-90-R data were compared to German population norms, stratified by age, sex, and comorbidities.

Fifty-one patients (72.5% females, mean age 47.3 years) were included with an average headache duration of 25.3 years. Moderate to high levels of headache were reported on the MIDAS VAS at 6.5 (range 0–10); SF-36 bodily pain was 40.3 (norm = 59.0, P < .001, 100 = best). Impaired functioning averaged at 78.4 (100 = no impairment) on the MIDAS. In contrast, SF-36 physical functioning was comparable to the norm (mean: 78.4, norm = 81.8, P = .63). All other SF-36 scales were significantly lower than expected from the norm (all P < .001). The scales depression, anxiety, obsessive-compulsive, and interpersonal sensitivity were significantly affected, whereas the levels of SCL-90-R schizophrenia nuclear and schizotypia were not lower than the norm. Coping with pain was moderate.

This pilot study is the first that presents a comprehensive and simultaneously specific assessment of health and quality of life of MOH patients after detoxification and inpatient rehabilitation. Moderate to high levels of pain and self-reported disability owing to headache were observed, whereas physical function on the SF-36 was not different from the expected level of the norm. Mental health was substantially affected in several dimensions, which had been described to reduce the ability to cope with pain. MOH patients seem to have high expectations of functionality, low symptomatology, and intact well-being.

Abbreviations: CSQ = coping strategies questionnaire, ES = effect size, HADS = Hospital Anxiety and Depression Scale, m = Mean, max = maximum, MCS = mental component summary, MIDAS = Migraine Disability Assessment Score, MOH = medication overuse headache, NRS = numeric rating scale, NSAIDs = non-steroidal anti-inflammatory drugs, p = type I error of the Wilcoxon test, PCS = physical component summary, PROM = patient-reported outcome measure, SCL-90-R = Symptom Checklist 90 revised, sd = standard deviation, SF-36 = short form 36, VAS = visual analogue scale, ZKP = Zurzach headache program (Zurzacher Kopfschmerz Programm).

Keywords: health, inpatient rehabilitation, medication overuse headache, outcome, quality of life

1. Introduction

Medication overuse headache (MOH) is a chronic headache syndrome associated with regular overuse of acute or symptom-
MOH patients improve after discontinuation of the overused substances.\[7\] Average medication use after detoxification decreased by about 14% and average annual costs for medication by 24%.\[10\] Inpatient and outpatient detoxification programs are effective in reducing medication and in decreasing headache frequency.\[11\] Dropout rates in outpatient programs are higher than in inpatient programs.\[11\] In particular, patients with “complicated MOH” benefit from a structured detoxification program.\[12\] Nevertheless, there is a lack of studies providing evidence for specific treatment strategies.\[13\] However, consensus reflected in guidelines highlights withdrawal as the major strategy.\[13,14\]

After detoxification, the relapse rate of MOH patients is high, especially in the first year, but decreases thereafter.\[15\] Experts recommend interdisciplinary rehabilitation after the detoxification to ensure long-lasting effects.\[16\] Further goals of multidisciplinary treatment are education, improvement of therapy to reduce headache frequency, and to increase quality of life.\[16\] There is a lack of evidence as to which therapeutic elements have to be part of a multidisciplinary treatment approach. There is only little empirical research about intensive multidisciplinary headache treatment programs for MOH, migraine, and tension-type headache, but some multimodal treatment approaches were shown to be effective for MOH and primary headaches.\[17,18\] The study of Gunreben et al.\[18\] presented pre- and postdata of intensive multidisciplinary pain program in terms of number of headache days per month, headache intensity, and depression. Pre- and post-data of standardized outcome instruments were not presented. At present, there are no published data on comprehensive health-related quality of life (HRQoL) from patients who underwent an intensive interdisciplinary MOH rehabilitation program after detoxification available.

The goal of this study was to quantify comprehensively as well as condition-specifically HRQoL for patients with MOH after detoxification and subsequent MOH rehabilitation program using standardized patient-reported outcome measures (PROMs). Data were compared to available normative values obtained by general population surveys. The hypothesis was that there were still health deficits in some health dimensions, especially in mental and psychosocial health scales, even after successful detoxification and subsequent rehabilitation. The findings help to get an idea of the health and impairment of those patients as well as the usefulness and feasibility of the generic and condition-specific PROMs. This should give a basis to plan future prospective studies, which quantify intervention effects.

2. Methods

2.1. Patients

This cross-sectional pilot study examined patients having participated in the Zurzach headache program (Zurzacher Kopfschmerz Programm, ZKP) at the rehabilitation center “RehaClinic,” Bad Zurzach, Switzerland, between July 2012 and June 2014, by means of a postal survey. The conditions for entry to the ZKP were diagnosis of MOH, confirmed by board-certified neurologists. Cross-sectional assessment was performed 0.5 to 2.5 years after the end of the ZKP.

Exclusion criteria for the ZKP and the study were: Abuse of benzodiazepines; serious psychiatric comorbidity such as psychosis or suicidality; severe somatic illness requiring specific treatment and preventing participation in the ZKP, for example, cancer, inflammatory rheumatic disease, serious other neurological diseases that prevented participation in the program (e.g., dementia); nonadherence for the correct intake of prescribed medication or regular participation in all therapies of the ZKP; insufficient German language skills to understand the study questionnaire. The study protocol was approved by the independent local ethics commission (Health Department in Aarau, Switzerland, EK AG 2008/026). Written, signed informed consent was obtained from all participants.

2.2. Intervention

Withdrawal of overused medication was performed in an acute hospital or, for a few exceptions, on an outpatient basis and with neurological supervision, directly before admission. This means that detoxification was completed before inpatient rehabilitation. During detoxification, all analgesics and triptans were stopped on the first day and were replaced by prednisone 100 mg per day for 5 days.\[19,20\] Prophylactic medication was started on the first day according to the treatment recommendations of the Swiss Headache Society.\[21\] If an analgesic reserve was needed, a substance from a class of acute headache medication other than the previously (over)used was provided, for example, an NSAID instead of a triptan.

Immediately after acute withdrawal, patients were admitted to the ZKP, a comprehensive and multimodal inpatient rehabilitation lasting 2 to 3 weeks which was established in 2010. The concept includes multidisciplinary therapies provided by a team of neurologists, behavioural and clinical psychologists, physical therapists, and nurses. The concept is standardized and in line with international recommendations.\[16\] Each week, patients had 3 physician visits and 2 to 3 consultations with psychologists, including patient education. The standardized weekly program also included daily physical therapy and aerobic exercise, 2 sessions of relaxation therapy, 2 to 3 sessions of medical massage, 2 to 3 sessions of acupuncture if applicable, individual use of thermal water, and medical training therapy. The treatment team met weekly to discuss the progress of the patients. During the period of this study, the physicians and therapists remained stable.

2.3. Measures

All medical records from neurologists and other physicians were obtained to confirm diagnosis of MOH, determine comorbidities, body height, and weight. Sociodemographic and disease-relevant data were measured using a standardized questionnaire that has already shown its usefulness in other studies.\[22,23\] The pain Numeric Rating Scale (NRS) was used, as it had been described to be easier to comprehend and to handle than the visual analogue scale (VAS), with comparable psychometric properties.\[24,25\]

The Short Form 36 (SF-36) is a self-assessment questionnaire that is used to measure health and HRQoL on 4 physical/somatic scales (physical functioning, role physical, bodily pain, general health) and 4 psychosocial/mental scales (vitality, social functioning, role emotional, mental health) comprising 36 items.\[26,27\] A complex linear combination of all 8 scales provides the summary scales, with specific weights for the physical component summary (PCS) and different ones for the mental component summary (MCS).\[27\] Population survey-based normative data for the SF-36 exist and allow quantification stratified by sex, age (5-year classes), and comorbidity (present/absent).\[28\] These data had been collected in Germany, having comparable language and cultural properties, as in the German-speaking part.
of Switzerland. The SF-36 is the best tested and most often used generic outcome measure having proven its validity and reliability in numerous studies. Validity in measurement of pain, physical and social function, depression, and anxiety in chronic pain conditions have been proven.

The Migraine Disability Assessment Score (MIDAS) was used to assess the condition-specific disability. It records headache-related disability by measuring missing days because of headache in the last 3 months on 5 items: school or paid work; number of days missed, number of days with half performance or lower; household work: number of days missed, number of days with half performance or lower; family, social or leisure activities: number of days missed. The number of days is summed up using items 1 to 5 and rescaled to a score of 0 = all days missed/disabled and 100 = full-working capacity on all days. In addition, the number of days with headache in the last 3 months is recorded, and pain is quantified on an NRS 0 to 10.

The Coping Strategies Questionnaire (CSQ) evaluates active and passive coping strategies and is internationally used in chronic pain conditions. The German version is cross-culturally adapted and validated. Catastrophizing and the 2 items “control over pain” and “ability to decrease pain” were included in this study because their clinical significance for MOH was considered to be most appropriate and the responsiveness to change was optimal.

Psychopathological assessment was performed by the Hospital Anxiety and Depression Scale (HADS) and the Symptom Checklist 90 revised (SCL-90-R). This was done because the association between psychopathological symptoms and headache had been shown to be elevated in migraine patients when compared to controls without migraine, whereas this was not the case in patients with tension-type headache.

The HADS is a questionnaire for patients of nonpsychiatric settings consisting of 14 items, 7 for anxiety and 7 for depression. German population survey-based normative data are stratified by sex and age (10-year classes). The SCL-90-R is a self-report of psychopathological symptoms, especially for nonpsychiatric populations. Patients rate distress on a 5-step Likert-scale on 90 items. Based on our clinical experience, the following specific scales that are assumed to be most important for MOH were selected: obsessive-compulsive, interpersonal sensitivity, schizotypy, and schizotypic nuclear. The latter 2 scales are constructs based on a large population-based survey. Depression and anxiety were already covered by the HADS. Population norms, stratified by sex and age using 10-year intervals from Germany, were used for comparison. The normative data from the scale paranoid ideation were compared with the scale schizophrenia nuclear and those of the scale psychotism were compared with the scale schizotypia.

2.4. Statistical analysis

Unless indicated differently (e.g., for the pain NRS), all instrument scores were scaled from 0 = worst to 100 = best health/function/ability in accordance with the original scoring of the SF-36 to ease comparison with scores obtained from other instruments. The specific “missing rules” of the instruments had to be fulfilled for determination of the scales. This means that at least 50% of the items had to be completed for each of the SF-36 scales and 6 of 7 for each of the HADS scales. For the CSQ, MIDAS, and the SCL-90-R, no “missing rules” were specified in the original reports; thus, the “2 of 3 (67%) rule” was applied as described for other instruments. Owing to the metric properties of the determination of the scales and to have a useful minimal number for metric statistics, a minimal number of n > 30 was set to be required with a wishful number of n ≥ 50 for this pilot study.

All analyses were performed using the statistical package IBM SPSS 22.0 for Windows (SPSS Inc, Chicago, IL). Two-tailed significance tests were performed by the nonparametric Wilcoxon test for continuous data to ensure appropriateness for all distributions, not only the Gaussian.

3. Results

3.1. Patients

Between July 2012 and June 2014, 106 patients with a confirmed MOH diagnosis were treated in the program, of which 83 (78%) could be contacted by telephone (max. 7 attempts). The remaining 23 received the assessment set without previous contact. Of those who were contacted by phone, 8 refused participation, and 7 had insufficient German language skills. Of the 91 mailed sets, 51 (36%) were returned with complete data. Of the 40 sets that were not returned, 4 patients withdrew willingness to participate; for 36 sets, no reasons for decline were given (Fig. 1). Time since the end of the pain program ranged between 0.5 and 2.5 years (Table 1).

The median study participant was female, middle-aged, lived with a partner or spouse, was educated to the level of vocational training or less, worked full-time, did not smoke, and performed sports up to 2 hours/week (Table 1). The individuals had suffered from headache for 25 years, had taken 1 substance against headache in the last 2 weeks before assessment, and felt much better or slightly better in comparison to their state before the pain program.

In the last 14 days before assessment, 2 or 3 different pain medications (either acute or prophylactic) were taken by 45% of patients. A minority still took acute headache medication: 24% paracetamol/NSAIDs, 14% triptans, and 2% opioids. Prophylactic medication (e.g., riboflavin or topiramate) was taken by 45%.

3.2. Outcome

Pain intensity levels of the pain scales were moderate at the time of the assessment (mean: 3.3/10 points = maximal pain) up to severe (6.8/10) at the worst moment of the last 7 days before, corresponding to a score of 6.5 of 10 on the MIDAS (Table 2). The MIDAS revealed on average 40.1% (37.7/92) days with headache within the last 3 months. Of the 51 patients, 33 (65%) reported <45 days with headache in the last 3 months. The number of days with impaired function owing to headache (sum of the items 1–5) averaged to a value of 59.5 (of 276), which would correspond to 6.0 on the VAS and is significantly higher than expected from the normative data for the general population (mean: 38.1, P < 0.001).

Physical functioning was not affected on the SF-36 when compared to the norm (means: 78.4 vs. 81.8, P = 0.66; 100 = best). All other scales of the SF-36 showed highly significant (P ≤ 0.001) impairments with levels far from the expected scores, for example, general health 48.7 versus 62.1 and social functioning 56.8 versus 82.1. The only exception is SF-36 mental health: 59.7 versus 68.5 = norm, P = 0.4.
Consistently, HADS depression and anxiety were much higher than expected from the normative levels: 67.6 versus 78.1, \( P = .02 \) and 63.3 versus 76.5, \( P < .001 \). The same was true for the obsessive/compulsive scale (74.1 vs. 87.4) and, somewhat less, for the interpersonal sensitivity scale (79.9 vs. 89.6) of the SCL-90-R. In contrast, patients were not more schizotypal (84.4 vs. 88.8, \( P = .81 \)), but were less “schizophrenic” (schizophrenia nuclear: 94.4 vs. 95.1) than expected. In the SCL-90-R schizophrenia nuclear, the median of the patients was 100, whereas the mean norm was 95.3, which led to a significant difference (\( P = .03 \)) in favor of the patients.

Coping with pain showed moderate levels in the middle range of the scale on the CSQ (catastrophizing: 53.1). This corresponds to the 37% replies of “much better” and 31% “slightly better” on the transition item (Table 1).

### Table 1

#### Sociodemographic and disease-relevant data (n = 51).

| Characteristics                      | %       |
|--------------------------------------|---------|
| Sex                                  | Female  |
| Education                            | Basic school (8–9 years) |
|                                      | Vocational training    |
|                                      | college/high school/college |
| Living                               | With partner/spouse |
| Smoking                              | Yes     |
| Sports                               | >2 h/wk |
|                                      | >0–2 h/wk |
|                                      | None    |
| Working capacity                     | Full (42 h/wk) |
|                                      | Part    |
|                                      | None    |
|                                      | Retired |
| Primary type of headache             | Migraine |
| (before MOH therapy)                 | Tension-type headache |
| Medication: number                   | None    |
| (last 14 days)                       | 1 Substance |
|                                      | 2 Substances |
|                                      | 3 Substances |
| Medication: therapeutic substances   | Paracetamol/NSAIDs |
| (last 14 days)                       | Opioids |
|                                      | Triptans |
| Medication: prophylactic substances  | Riboflavin |
| (last 14 days)                       | Topiramate |
| Medication                           | Others |
| Transition                           | Much better |
| (compared to the state before the pain program) | Slightly better |
|                                      | Almost the same |
|                                      | Slightly worse |
|                                      | Much worse |

| Characteristic                        | Mean (sd) |
|---------------------------------------|-----------|
| Age, y                                | 47.3 (11.8) |
| Body mass index, kg/m²                | 25.3 (14.4) |
| Duration headache, y                  | 24.8 (4.8)  |
| Time since pain program, y            | 1.3 (0.6)   |

MOH = medication overuse headache, NSAIDs = nonsteroidal anti-inflammatory drugs, sd = standard deviation.

* Not related to headache.

Figure 1. Patient selection. MOH = medication overuse headache, ZKP = Zurzach headache program (Zurzacher Kopfschmerz Programm).

4. Discussion

This cross-sectional pilot study quantified health and HRQoL in patients with history of MOH after detoxification and a subsequent headache-specific inpatient rehabilitation program with a comprehensive set of generic and condition-specific questionnaires. The study did not aim to quantify treatment effects, but to provide data about patients’ state of health and appropriateness of the assessment as base for planning future prospective studies. Where possible, empirical data were compared to normative values obtained by general population surveys. The hypothesis that deficits persisted in some health dimensions after withdrawal and subsequent rehabilitation was confirmed and psychopathological symptoms persisted. In our sample, MOH was associated with female sex, middle age (40–50 years), and relatively low education level. These characteristics are consistent with many other European studies.[6,18,47–51]

Substantial burden of headache was measured by the NRS, the MIDAS, and the SF-36—the score of SF-36 bodily pain showed...
statistically significant difference from the norm. The same was true for all other scales of the SF-36, except for the SF-36 physical functioning. The number of days with headache and with consequent limitations at work and in daily life on the MIDAS showed levels compatible with significant burden. Comparison to norms is not possible because of lack of data. Affective symptoms as well as obsessive and compulsive symptoms and interpersonal sensitivity were significantly increased, whereas schizophrenial or schizophrenic symptoms were not. Finally, ability to cope with pain was moderate and comparable to other chronic pain patients.[26]

The pattern of our empirical SF-36 scores compared with that of the norm scores was similar to that of previous studies: the highest differences were in role physical, bodily pain, social functioning, and the lowest were in physical functioning. This pattern was consistent with a population-based sample in Spain (n = 74 MOH).[6] However, in that study, the smallest difference in SF-36 physical functioning still reached significance, whereas it did not in our data. A previously conducted survey by the same study group (n = 22 MOH) found SF-36 scores, which were much lower than those of our patients for physical functioning (59.2 vs. 78.4 in our study) and mental health (48.2 vs. 59.7 in our study), whereas the other scales were comparable.[25]

To the best of our knowledge, no further study exists, using either the SF-36 or the SF-12 to compare MOH (after detoxification and headache specific rehabilitation) with healthy controls, whereas several studies of chronic daily headache sufferers have been published. One review of comparative studies concluded that 3 of 5 studies having used SF-36 or SF-12 suggested that MOH patients have worse HRQoL than those with chronic headache without medication overuse.[53] However, the reviewed data presented mean differences between different studies in bar plots and did not provide the exact figures.[54] For this reason, it is important to know—besides the previous diagnosis apart from MOH—whether the patients had a history of MOH or not. The primary diagnosis of most patients in this study was migraine (92%, Table 1).

The SF-36 PCS and MCS scores of this present study were comparable to a study conducted at the 6-month follow-up after an inpatient withdrawal therapy for MOH: the mean PCS at follow-up was 43.2 versus 41.7 in our study, the MCS 42.2 versus 39.0, respectively.[18] Post-treatment SF-36 scores showed significant improvement when compared to baseline.[14]

In a study about the effectiveness of an intensive multidisciplinary headache treatment program lasting 96 hours, pretreatment PCS was 37.9 and MCS was 39.0.[14] Post-treatment SF-36 scores were not reported. However, patients significantly improved in the depression and in the number of days with headache, but not on pain intensity.

On the MIDAS, MOH-related disability continuously decreased during the follow-up period after an inpatient withdrawal treatment from an average of 70.8 (MIDAS sum of items 1–5) before treatment to 34.1 one year later and to 17.0 at 5 years.[55] These post-treatment levels were lower than ours at 59.5. In contrast, our data are comparable to the baseline MIDAS score of 59.9 of a multicenter, multinational study.[56] Those cases improved to a score of 25.7 at the 6-month follow-up. This means

Table 2

| Outcome | Test | Patient | Norm |
|---------|------|---------|------|
| Pain VAS | At the moment (10 = max) | 3.3 | 2.8 | — | — | — |
| Pain VAS | At its worst of 7 days (10 = max) | 6.8 | 2.5 | — | — | — |
| Pain VAS | At its best of 7 days (10 = max) | 1.9 | 1.9 | — | — | — |
| MIDAS | Sum of items 1–5 (276 = max) | 59.5 | 64.8 | — | — | — |
| MIDAS | Days with headache (92 = max) | 37.7 | 29.9 | — | — | — |
| MIDAS | Intensity of headache (10 = max) | 6.5 | 2.0 | — | — | — |
| MIDAS | Total score (100 = best) | 78.4 | 21.4 | 81.8 | 8.7 | .03 |
| SF-36 | Physical functioning | 78.4 | 21.4 | 75.7 | 9.8 | <.001 |
| SF-36 | Role physical | 50.5 | 23.8 | 58.1 | 9.2 | <.001 |
| SF-36 | Bodily pain | 40.3 | 20.3 | 62.1 | 6.8 | <.001 |
| SF-36 | General health | 48.7 | 21.5 | 55.5 | 3.6 | <.001 |
| SF-36 | Vitality | 41.3 | 21.8 | 81.8 | 3.6 | <.001 |
| SF-36 | Social functioning | 56.8 | 28.1 | 82.1 | 3.6 | <.001 |
| SF-36 | Role emotional | 64.2 | 29.5 | 84.2 | 5.1 | <.001 |
| SF-36 | Mental health | 59.7 | 22.3 | 68.5 | 3.5 | <.001 |
| SF-36 | Physical component summary | 41.7 | 6.7 | 45.9 | 4.0 | <.001 |
| SF-36 | Mental component summary | 39.4 | 13.9 | 49.1 | 1.3 | <.001 |
| HADS | Depression | 67.6 | 24.1 | 78.1 | 4.0 | <.001 |
| HADS | Anxiety | 63.3 | 19.4 | 76.5 | 1.9 | <.001 |
| SCL-90-R | Schizophrenia nuclear | 94.4 | 12.2 | 95.1 | 1.7 | <.001 |
| SCL-90-R | Schizotypia | 84.4 | 16.8 | 88.8 | 1.8 | <.001 |
| SCL-90-R | Interpersonal sensitivity | 79.9 | 16.8 | 89.6 | 2.1 | <.001 |
| SCL-90-R | Obsessive compulsive | 74.1 | 19.1 | 87.4 | 2.1 | <.001 |
| CSQ | Catastrophizing | 53.1 | 23.5 | — | — | — |
| CSQ | Control pain | 49.7 | 23.2 | — | — | — |
| CSQ | Decrease pain | 49.7 | 23.5 | — | — | — |

CSQ = Coping strategies questionnaire, HADS = Hospital Anxiety and Depression Scale, m = mean, max = maximum, MIDAS = Migraine Disability Questionnaire, p = type I error of the Wilcoxon test: patient versus norm data, SCL-90-R = Symptom checklist 90 revised, sd = standard deviation, SF-36 = Short form 36, VAS = visual analogue scale. Scaling (where not indicated differently): 0 = worst, maximal symptoms to 100 = best, no symptoms.
that the posttreatment scores of our patients indicated much greater disability because of headache than in the 2 aforementioned studies. Based on the observed improvements on the MIDAS score as reported in 4 treatment studies, it may be hypothesized that our sample was a selection of highly disabled patients admitted to intensive inpatient rehabilitation.[55–58] Scores for schizophrenia, and equally schizotypia, were significantly lower, but scores for interpersonal sensitivity and obsessive compulsive (symptoms) were significantly higher, and scores for depression and anxiety were higher than expected from the norm. Schizophrenia nuclear is a milder version of the core diagnostic symptoms of psychosis, which is thought to be a biologically based perception disorder.[63] Schizotypal signs are characterized by generalized distrust, odd interpersonal beliefs, and paranoid ideation, which lead to a distorted perception of the environment, influenced by psychosocial conditions.[43] The 2 symptom dimensions show relatively high overlap on the psychopathological level, but coincide only partly.[43] They are expression of psychotic symptoms with different levels of severity and persistence on the continuum of the psychotic phenotype.[65] This is consistent with previous studies. MOH was associated with obsessive-compulsive disorders; in one study, the prevalence of the latter was 28%.[60] Psychiatric comorbidities are predictors for prognosis and response to treatment.[50–52] A small study showed a significantly increased risk of suffering from overall mood disorders (odds ratio, OR = 4.5), anxiety (OR = 5.0) and disorders associated with the use of psychoactive substances other than analogics (OR = 7.6) in MOH sufferers when compared to patients with migraine only (n = 41 in both groups).[63] The HADS anxiety score of 63.3 in our study is comparable to a 3-month follow-up score of 66.7[62] and a 6-month follow-up score of 66.2,[56] whereas the depression score in our study (67.6) is lower when compared to the aforementioned studies (81.0 at 3-month follow-up; 80.4 at 6-month follow-up).

In this cross-sectional pilot study, the combination of generic and condition-specific assessments provided comprehensive but also disease-specific measurements of health and HRQoL. This is the strength of the study. The questionnaire set gave moderate respondent and administrative burden and the single instruments turned out to be appropriate for MOH. Furthermore, the instruments were internationally used, validated, and standardized, and allowed comparison of empirical with normative data for some instruments. Even after completion of the therapies, deficits on various health dimensions could still be measured as compared to normative data. This means that the instruments used did not reveal ceiling effects, which may have reduced validity of the measurements. Further strengths were a compact cohort with representative sociodemographic characteristics and a standardized intervention throughout the whole period of treatment.

The cross-sectional design of this study does not allow the evaluation of health changes or evaluation of the therapeutic effects of the rehabilitation program. Furthermore, this study does not allow a conclusion about the compliance of the patients to the recommended treatment and instructions after the rehabilitation program. However, the study aimed to provide basic data on the state of health and quality of life after specific treatment for MOH in a sample of severely affected patients. A further limitation is that the time interval between the completion of the headache program and the postal survey varied between 0.5 and 2.5 years. The sample size of 51 out of a patient population of 106 with confirmed MOH diagnosis is limited and cannot exclude a selection bias, for example, towards patients with better health status, which is an inherent problem in every voluntary survey.

This study is the first that presents a comprehensive and simultaneously specific assessment of health and quality of life of MOH patients after detoxification and inpatient rehabilitation. Moderate to high levels of pain and self-reported disability owing to headache were observed, whereas physical function on the SF-36 was not different from the expected level of the norm. This may be because of high expectations of functionality and corresponds to low self-reported coping with pain, relatively high catastrophizing, depression, and anxiety. The data of this cross-sectional pilot study sampled over 2 years can be taken as valid estimate of the outcome of the pain program in our clinic. Generalizability to other pain programs should be done with care because comparable data with standardized instruments are lacking from other programs. Future longitudinal data will provide quantification of treatment effects and refine insight into processes of treatment, individual developments, and need specific to MOH during the course of treatment and recovery.

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