Anthroposophic Therapy for Migraine: A Two-Year Prospective Cohort Study in Routine Outpatient Settings

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Abstract: Background and Methods: Anthroposophic treatment for migraine is provided by physicians and includes special artistic and physical therapies and special medications. We conducted a prospective cohort study of 45 consecutive adult outpatients (89% women) starting anthroposophic treatment for migraine under routine conditions. Main outcomes were Average Migraine Severity (physician and patient ratings 0-10, primary outcome), Symptom Score (patient rating, 0-10), and quality of life (SF-36); main follow-up time point was after six months.

Results: The anthroposophic treatment modalities used were medications (67% of patients), eurythmy therapy (38%), art therapy (18%), and rhythmical massage therapy (13%). Median therapy duration was 105 days. In months 0-6, conventional prophylactic antimigraine medications were used by 14% (n=5/36) of evaluable patients.

From baseline to six-month follow-up, physician-rated Average Migraine Severity improved by 3.14 points (95% confidence interval 2.40-3.87, p<0.001); patient-rated Average Migraine Severity improved by 2.82 points (2.05-3.64, p<0.001); and Symptom Score improved by 2.32 points (1.68-2.95, p<0.001). In addition, three SF-36 scales (Social Functioning, Bodily Pain, Vitality), the SF-36 Physical Component summary measure, and the SF-36 Health Change item improved significantly. All improvements were maintained at last follow-up after 24 months. Patients not using conventional prophylactic antimigraine medications had improvements similar to the whole cohort.

Conclusions: Patients with migraine under anthroposophic treatment had long-term improvement of symptoms and quality of life. Although the pre-post design of the present study does not allow for conclusions about comparative effectiveness, study findings suggest that anthroposophic therapies may be useful in the long-term care of patients with migraine.

Keywords: Anthroposophy, combined modality therapy, drug therapy, eurythmy therapy, migraine, prospective studies.

INTRODUCTION

Migraine affects 14% of adults in Europe every year [1] and is associated with substantial disability, reduced quality of life, reduced work capacity, psychiatric and somatic comorbidity, and increased healthcare use [2-4]. The World Health Organization has ranked severe migraine attacks among the most disabling medical conditions, along with psychosis, dementia, and quadriplegia [5]. Up to three-fourths of patients with acute migraine attacks do not become pain free from 5-HT1 agonists ("triptanes") [6, 7] and half of patients will not respond to prophylactic medications [6]. Many migraine patients use complementary therapies [8-10], sometimes provided by their physicians.

Anthroposophic medicine (AM) is a physician-provided complementary therapy system founded by Rudolf Steiner and Ita Wegman in the 1920s [11]. AM extends the conception of human physiology beyond cellular and molecular mechanisms to holistic interactions of over-riding functional systems (downward causality [12]). In particular, a complex equilibrium exists between two polar systems: the 'nerve-sense system' (low metabolic rate, mediator of consciousness) and the 'metabolic system' of the abdominal organs (high metabolic rate, minimal consciousness). This equilibrium can be distorted in human disease. In migraine, a predominance of the metabolic system (with symptoms such as nausea, vomiting, diarrhea, abdominal cramps) leads to disturbances in the nerve-sense system (e.g. headache, sensory hypersensitivity, visual disturbances) [13-15]. This imbalance is sought to be regulated by special AM therapies (eurythmy movement exercises, art therapy, rhythmical massage therapy) [16] and special AM medications [14, 15, 17, 18].

Eurythmy therapy is an artistic exercise therapy involving cognitive, emotional and volitional elements. In eurythmy therapy sessions the patients exercise specific movements with the hands, the feet or the whole body. Eurythmy movements are related to the sounds of vowels and consonants, to music intervals or to soul gestures, e.g. sympathy-antipathy. Between therapy sessions the patients exercise eurythmy movements daily. In AM art therapy the patients engage in painting, drawing, clay modelling, music or speech exercises. Rhythmical massage therapy was devel-
opened from Swedish massage; special techniques include lifting movements, rhythmically undulating gliding movements, and complex movement patterns like lemniscates [19].

AM medications are prepared from plants, minerals, animals and from chemically defined substances and can be prepared in concentrated or potentised form. Potentization implies a successive dilution, each dilution step involving a rhythmic succussion (repeated shaking of liquids) or trituration (grinding of solids into lactose monohydrate). For example, a D6 potency (also called 6X) has been potentized in a 1:10 dilution six times, resulting in a 1:10⁻⁶ dilution [20]. Since potencies beyond D23 do not contain any molecules of the original substance, effects cannot readily be explained by molecular mechanisms. However, a systematic review of in vitro studies found biological effects of potencies ≥D23 in nearly three-fourths of the studies and in more than two-thirds of the studies with highest quality [21]. All AM medications are manufactured according to Good Manufacturing Practice and national drug regulations; quality standards of raw materials and manufacturing methods are described in the Anthroposophic Pharmaceutical Codex [20]. The available evidence suggests that AM medications and therapies are generally well tolerated with infrequent adverse reactions of mostly mild to moderate severity [19, 22].

AM therapy is practised by physicians (counselling, AM medication) and non-medical therapists (eurythmy, art, rhythmical massage) in 67 countries worldwide [23]. AM therapy providers are certified following structured training programs according to international, standardized curricula. Therapy guidelines exist for AM medical therapy [24], eurythmy therapy [25], and AM art therapy [26]. AM is practised in general practice and in specialist fields, including neurology, and is also integrated into intensive care and emergency medicine provided in hospitals, including university teaching hospitals [19]. For patients with migraine, the physician will choose among the available AM therapy modalities in order to tailor the treatment to individual needs of the patient. AM treatments can be administered alone or combined with conventional antimigraine medications [14].

Several studies have evaluated acute or prophylactic therapy for migraine with Ferrum-Quartz [19], a mixture of ferrous sulphate, honey and quartz, available as 150 mg capsules and 10 mg tablets ('Kephalodor' or 'Biodoron' tablets) [20]. Here we present a pre-planned subgroup analysis of migraine patients from a study of a broader range of AM therapy options (eurythmy therapy, art therapy, rhythmical massage therapy, medications) [27].

For complementary therapy systems in widespread use, regardless of whether evidence from randomized trials exists, it has been argued that the conventional drug research strategy – starting with studies of biological mechanisms and moving through Phase I, II and III clinical trials – is not optimal [28]. Another, more appropriate strategy has been proposed, moving from descriptive studies ('Phase 1') towards comparative studies of the whole system and its parts, and ending with studies of biological mechanisms ('Phase 5') [28]. In the context of this reversed strategy, the present analysis addressed topics in the Phases 1-2 (therapy paradigms, therapy use, clinical outcomes, perceived benefit, safety).

MATERIALS AND METHODOLOGY

Study Design and Objective

This is a prospective two-year cohort study in a real-world medical setting. The study was initiated by the health insurance company Innungskrankenkasse Hamburg as part of a research project on the effectiveness, costs, and safety of AM therapies in outpatients with chronic disease (Anthroposophic Medicine Outcomes Study, AMOS) [27, 29, 30]. The present pre-planned analysis concerned the subgroup of adult patients with migraine. Since this was the first prospective study of comprehensive AM treatment for this indication, the primary objective was to describe AM therapy for migraine (characteristics of AM therapy users, spectrum of AM therapy modalities used, extent of combination with conventional anti-migraine therapy) as well as clinical outcomes after AM treatment under routine conditions. Further research questions addressed the use of health services, adverse reactions, and therapy satisfaction.

Setting, Participants, and Therapy

All physicians certified by the Physicians’ Association for Anthroposophical Medicine in Germany and working in an office-based practice or outpatient clinic were invited to participate in the AMOS study. Certification as an AM physician required a completed medical degree and a three-year structured postgraduate training. The participating physicians recruited consecutive patients starting AM therapy under routine clinical conditions. Patients enrolled from 1 January 1999 to 31 December 2005 were included in the analysis if they fulfilled the eligibility criteria. Inclusion criteria were:

1. Age 17-75 years.
2. A diagnosis of migraine according to the criteria of the International Headache Society [31], assessed by the Kieler Kopfschmerzfragebogen, a standardized headache questionnaire in the German language [32;33], documented by the patients (for patients enrolled before April 2001: physician’s diagnosis, assisted by a brief written clinical description). In accordance with the clinical practice guidelines of the Drug Commission of the German Medical Association [34], neuroimaging was not required for routine diagnostics of all patients.
3. Starting AM therapy for migraine:
   - AM-related consultation of at least 30 minutes followed by new prescription of AM medication,
   - or new referral to AM therapy (art, eurythmy or rhythmical massage).

Patients were excluded if they had previously received the AM therapy in question (see inclusion criteria 3) for migraine. Patients were treated according to the physician’s discretion. AM therapy was evaluated as a whole system [35]. Additional costs for AM treatments were 0.1-5.2 Euro per daily dose of an AM medication (51 different price groups, median 0.2 Euro, mean 0.8 Euro); 20-32 Euro per AM therapy session; and 46 Euro and 92 Euro for AM consultations with physicians of 30 min and 60 min duration, respectively. These costs were reimbursed by some but not all German health insurance companies.
The data were entered twice by two different persons into patients received no compensation. Physicians were compensated 40 Euro (after March 2001: 60 Euro) per included and fully documented patient, while the therapists were compensated 10 Euro. The response to medications or therapies were not made available to the physicians. The therapists documented eligibility criteria; the therapists documented adverse reactions to medications or therapies were documented by the physicians throughout the study. Serious adverse events were documented by the physicians after 6 and 12 months. Adverse reactions to medications or therapies were documented by the patients after 6, 12, 18 and 24 months and by the physicians after 6 months. The documentation included cause, intensity (mild / moderate / severe = no / some / complete impairment of normal daily activities), and therapy withdrawal due to adverse reactions. Serious adverse events were documented by the physicians throughout the study.

Secondary clinical outcomes were Symptom Score and quality of life. Symptom Score, the severity of one to six most relevant symptoms present at baseline, was assessed by patients on numerical rating scales from 0 (“not present”) to 10 (“worst possible”). Quality of life was assessed with the SF-36 Health Survey [37] (Physical and Mental Component summary measures, eight scales, Health Change item).

Physician-rated Average Migraine Severity was documented after 0 and 6 months, while all other clinical outcomes were documented after 0, 3, 6, 12, 18, and 24 months.

### Other Outcomes

Therapy outcome rating (0-10), satisfaction with therapy (0-10) and therapy effectiveness rating (“very effective”, “less effective”, “ineffective” or “not evaluable”) were documented by the patients after 6 and 12 months.

Adverse reactions to medications or therapies were documented by the patients after 6, 12, 18 and 24 months and by the physicians after 6 months. The documentation included cause, intensity (mild / moderate / severe = no / some / complete impairment of normal daily activities), and therapy withdrawal due to adverse reactions. Serious adverse events were documented by the physicians throughout the study.

### Data Collection

All data were documented with questionnaires returned in sealed envelopes to the study office. The physicians documented eligibility criteria; the therapists documented AM therapy administration; all other items were documented by the patients, unless otherwise stated. The patient responses were not made available to the physicians. The physicians were compensated 40 Euro (after March 2001: 60 Euro) per included and fully documented patient, while the patients received no compensation.

The data were entered twice by two different persons into Microsoft® Access 97 (Microsoft Corp., Redmond, WA, USA). The two datasets were compared and discrepancies resolved by checking with the original data. Non-AM medications were coded according to the Anatomical Therapeutic Chemical Classification Index.

### Quality Assurance, Adherence to Regulations

The study was approved by the Ethics Committee of the Faculty of Medicine Charité, Humboldt University, Berlin, Germany, and was conducted according to the Declaration of Helsinki and largely following the ICH Guideline for Good Clinical Practice E6. Written informed consent was obtained from all patients before enrolment.

### Data Analysis

The data analysis was performed for all patients fulfilling the eligibility criteria, using SPSS® 14.0.1 (SPSS Inc., Chicago, Ill, USA) and StatXact® 5.0.3 (Cytel Software Corporation, Cambridge, MA, USA). T-test was used for continu-ous data with normal distribution; Wilcoxon Signed-Rank test was used for paired continuous data with non-normal distribution; McNemar test and Fisher’s exact test were used for dichotomous data. All tests were two-tailed. Significance criterion was p < 0.05. Since this was a descriptive study, no adjustment for multiple comparisons was performed [38]. Pre-post effect sizes were calculated as Standardized Response Mean (= mean change score divided by the standard deviation of the change score) and classified as minimal (< 0.20), small (0.20-0.49), medium (0.50-0.79), and large (≥ 0.80) [39, 40]. Clinical outcomes were analyzed with replacement of missing values by the last value carried forward.

Two pre-planned sensitivity analyses (SA1-SA2) were performed to assess the influence of natural recovery and adjunctive therapy on the 0-6-month outcome of physician- and patient-rated Average Migraine Severity. SA1 concerned natural recovery, which was assumed to be unlikely in patients with disease duration of at least one year [41]. The sample was therefore restricted to patients with disease duration of at least 12 months prior to study enrolment. SA2 concerned the effects of prophylactic antimigraine medications: The sample was restricted to patients not using prophylactic antimigraine medications during the first six study months.

### RESULTS

#### Participating Physicians and Therapists

The patients were enrolled by 30 physicians with four different qualifications (27 general practitioners, one neurologist, one internist, and one gynaecologist). Comparing these physicians to AM-certified physicians in Germany with the same four qualifications but without study patients (n = 225), no significant differences were found regarding gender (60.0% vs. 58.5% males), number of years in practice (16.1 ± 6.9 vs. 19.3 ± 8.9), and the proportion of physicians working in primary care (90.0% vs. 92.0%). Mean age was 44.9 ± 5.9 and 48.3 ± 8.0 years in physicians with and without study patients, respectively (mean difference 3.4 years, 95% confidence interval 0.4-6.3 years, p = 0.027).

The patients were treated by 31 different AM therapists (art, eurythmy, rhythmical massage). Comparing these therapists to certified therapists without study patients (n = 1137), no significant differences were found regarding gender (87.1% vs. 80.7% women), age (mean 48.2 ± 9.3 vs. 50.3 ± 9.4 years) or the number of years since therapist qualification (10.5 ± 6.9 vs. 13.2 ± 8.7 years).

#### Patient Recruitment and Follow-Up

A total of 79 patients starting AM therapies for headache were screened for inclusion. Of these patients, 45 fulfilled all eligibility criteria and were included in the analysis. 34 patients were not included; in 27 of these patients the eligibility criteria were not fulfilled (the indication for AM therapy was not migraine but another headache disorder). The remaining seven patients were potentially eligible but not included for the following reasons: patients’ baseline questionnaire missing (n = 3), physician’s baseline documentation missing (n = 3), patients’ and physicians’ baseline questionnaire dated > 30 days apart (n = 1).

A total of 89% (40/45) of patients were enrolled by general practitioners, while 11% were enrolled by specialists.
The physicians’ settings were primary care practices (89% of evaluable patients, n = 39/44), referral practice (7%, n = 3), and outpatient clinics (5%, n = 2).

The last patient follow-up ensued on 9 May 2007. A total of 96% (n = 43/45) of patients returned at least one follow-up questionnaire. The patients were administered a total of 225 follow-up questionnaires, out of which 157 (70%) were returned. Follow-up rates were 89% (n = 40/45), 76%, 69%, 60%, and 56% after 3, 6, 12, 18, and 24 months, respectively. Respondents (n = 34) and non-respondents (n = 11) of the six-month follow-up questionnaire did not differ significantly regarding age, gender, disease duration or baseline parameters (physician- and patient-rated Average Migraine Severity, Symptom Score). Corresponding dropout analysis for the 24-month follow-up also showed no significant differences between respondents (n = 25) and non-respondents (n = 20). The physician six-month follow-up documentation was available for 82% (n = 37/45) of patients.

Baseline Characteristics

The patients were recruited from 12 of 16 German federal states. Mean age was 39.5 ± 11.1 years (range 17-61 years, Table 1). A total of 89% (n = 40/45) of the patients were women. Median duration of the migraine disorder was 12.0 years (interquartile range [IQR] 2.5-20.0, range 4 months to 40 years, mean 13.6 ± 10.1 years). The number of days with headache in the past month was documented in patients enrolled after March 2001 (n = 23) and was 1 day in 4% (n = 1/23), 2-6 days in 61% (n = 14), 7-10 days in 26% (n = 6), 11-15 days in 0% (n = 0), and > 15 days in 9% (n = 2), with a median number of 5.0 days (mean 6.8 ± 6.3 days).

A current comorbid disease was present in 78% (35/45) of patients. Most common comorbid diseases, classified by the International Statistical Classification of Diseases and Related Health Problems, 10th Revision, were F00-F99 Mental and Behavioural Disorders (24%, n = 17 of 70 diagnoses), N00-N99 Diseases of the Genito-urinary System (17%), L00-L99 Diseases of the Skin and Subcutaneous Tissue (13%), and M00-M99 Musculoskeletal Diseases (9%). Further baseline data are presented in Table 1.

Therapy

AM Therapies

At study enrolment, the duration of the consultation with the AM physician was < 30 min in 40% (n = 18/45) of patients, 30-44 min in 40%, 45-59 min in 9%, and 60 min in 11%. At enrolment 22% (n = 10/45) of patients fulfilled In-
clusion Criterion 3a (AM-related consultation of ≥ 30 minutes followed by new prescription of AM medication), 38% (n = 17) fulfilled Inclusion Criterion 3b (referral to AM art, eurythmy or rhythmic massage therapy), and 40% (n = 18) fulfilled Inclusion Criteria 3a and 3b. Of the 35 patients referred to AM art/eurythmy or massage therapy, 89% (n = 31) had the planned AM therapy, 3% (n = 1) did not have AM therapy, and for 9% (n = 3) the AM therapy documentation is incomplete. AM therapies used were eurythmy therapy (n = 17 patients), rhythmical massage therapy (n = 6), and art therapy (n = 8). The AM eurythmy/art/massage therapy started median 9 (IQR 0-35) days after enrolment. Median therapy duration was 105 days (IQR 77-246 days), median number of therapy sessions was 12 (IQR 10-20). AM medications were used by 67% (n = 30/45) of patients. The most frequently used AM medications in months 0-6 were Ferrum-Quartz (n = 14 patients, thereof n = 7 using 150 mg capsules and n = 7 using 10 mg tablets), and Phosphorous dilution (n = 3 patients – contains potentised yellow phosphorus; used by the three patients in a D5, D6, and D20 potency, respectively).

Use of Conventional Antimigraine Medications in Months 0-6

Medications to treat acute migraine attacks were used by 53% (n = 19/36) of evaluable patients; medication groups used were non-opioid analgesics (44%), non-steroid anti-inflammatory agents (6%), and 5-HT1 agonists (14%). The total number of days with use of medication to treat acute migraine attacks in months 0-6 was 1-9 days (n = 9 patients), 10-19 days (n = 3), and 20-90 days (n = 19). Prophylactic antimigraine medications were used by 14% (n = 5/36); medications used were antidepressants (8%), cyclandelate (3%), and flunarazine (3%). The total number of days of use of prophylactic medications was 5 days (n = 1), 60-180 days (n = 3), and unknown (n = 1).

Clinical Outcomes

All clinical outcomes were improved from baseline at all subsequent follow-ups (exception: the SF-36 scale Role Emotional in months 0-6) (Table 2, Figs. 1-3). For seven of the 14 outcomes (physician- and patient-rated Average Migraine Severity, Symptom Score, the SF-36 scales Social Functioning, Bodily Pain, and Vitality and the SF-36 Health Change item) all improvements from baseline were significant at all follow-ups. Three further outcomes (SF-36 Physical and Mental Component summary measures, SF-36 scale Mental Health) were significantly improved at three or four out of five follow-ups.

At six-month follow-up, an improvement of ≥ 50% of baseline Average Migraine Severity was observed in 52% (n =23/44) of evaluable patients for physician rating and 45% (n = 20/44) for patient rating. Effect sizes for the 0-6-month comparison were large for three outcomes (physician- and patient-rated Average Migraine Severity, Symptom Score), medium for three outcomes (SF-36 Social Function, Bodily Pain, Health Change), small for six outcomes, and minimal for two outcomes (Table 2).

We performed two sensitivity analyses of the 0-6-month outcome of physician- and patient-rated Average Migraine Severity (Table 3: SA1-SA2; see Methods for further description). For physician rating, the individual analyses resulted in a reduction of the average 0-6 month improvement.

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<tr>
<th>Table 2. Clinical Outcomes 0-6 Months</th>
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<td>SF-36 Health Change**</td>
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*SPositive differences indicate improvement. **SF-36 Health Change: range from 1 ("much better now than one year ago") to 5 ("much worse now than one year ago"). SRM: Standardized Response Mean effect size (minimal: < 0.20, small: 0.20-0.49, medium: 0.50-0.79, large: 0.80).
Fig. (1). Average Migraine Severity.
Range: 0 “not present”, 10 “worst possible”. Physician and patient ratings.

Fig. (2). Symptom Score.
Range: 0 “not present”, 10 “worst possible”.

of maximum 6% (3.14±2.95 points). Combining both analyses, the improvement was reduced by 10% (3.14±2.82 points). For patient rating, the individual analyses resulted in a reduction of the improvement of maximum 8% (2.94±2.61 points). Combining both analyses, the improvement was reduced by 2% (2.84±2.78 points).

Other Outcomes
At six-month follow-up, the patients’ average therapy outcome rating (numeric scale from 0 “no help at all” to 10 “helped very well”) was 7.33 ±2.42 and patient satisfaction with therapy (from 0 “very dissatisfied” to 10 “very satisfied”) was 7.72 ±2.05. The patients’ effectiveness rating of eurhythm, art or rhythmical massage therapy was positive (“very effective” or “effective”) in 76% (n = 19/25) of evaluable patients who had started therapy, and negative (“less effective”, “ineffective” or “not evaluable”) in 24%. The ratings of therapy outcome, satisfaction and effectiveness did not change significantly between 6- and 12-month follow-up.
Fig. (3). SF-36 Scales, SF-36 Health Change Item.

SF-36 scales: Range 0-100, higher scores indicate better health. SF-36 Health Change: Range from 1 (“much better now than one year ago”) to 5 (“much worse now than one year ago”).
The frequency of reported adverse drug reactions was 3% (n = 1 of 30 users) for AM medications and 9% (n = 3 of 34 users) for non-AM medications (p = 0.616). Adverse drug reactions of severe intensity were reported in three patients using non-AM medications (unspecified psychic symptoms from homoeopathic arsenicum album; nightmares from amitriptyline; skin blisters from estradiol/norethisterone tablets), while medication was stopped due to reported adverse reactions in one patient using AM medication (skin blisters from rosemary ointment). No adverse reactions from non-medication therapies occurred. No serious adverse events occurred.

**DISCUSSION**

This is the first prospective evaluation of comprehensive AM treatment for migraine. We aimed to obtain information on AM therapy for this indication under routine outpatient conditions in Germany and studied adults starting AM therapy for migraine. Two-thirds of patients used non-medication AM therapies; two-thirds used AM medications; one in seven patients also used conventional prophylactic medications. Under AM treatment, significant and sustained improvements of migraine severity and quality of life were observed.

Strengths of this study include a detailed assessment of the therapy setting and therapy-related factors, a long follow-up period, and high representativeness: 11% of all AM-certified physicians seeing migraine patients in Germany participated; the participating AM physicians and therapists resembled all eligible physicians and therapists with respect to socio-demographic characteristics; and 87% of screened and eligible patients were enrolled. These features suggest that the study mirrors contemporary AM practice to a high degree.

To assess the routine clinical practice, where the selection of AM therapy options will vary according to individual needs, we analyzed AM as a whole system [35]. The sample size of AM therapy modality subgroups (eurythmy, art, rhythmical massage, medical) did not allow for subgroup analysis. The influence of the AM therapy modality and other therapy variables (e.g., duration of the consultation with the physician at study enrolment, number of AM therapy sessions) on clinical outcomes has been assessed in a multivariate analysis of adult AMOS patients with migraine and other chronic indications [52].

Another limitation in regard to sample size applies to the number of days with headache. This outcome, widely used in migraine studies, was only documented in a subset of patients, the documentation did not distinguish between migraine and interval headaches, and the number of patients with evaluable data at six-month follow-up (n = 13) did not allow for outcome analysis.

Since the study had a long recruitment period, the study physicians were not able to participate throughout the period and to screen and enrol all eligible patients (criteria: see Methods section). For a different subset of patients from the AMOS project (patients referred to AM therapies for any chronic indication and enrolled before 1 April 2001), it was estimated that physicians enrolled every fourth eligible patient [53]. This selection could bias results if physicians were able to predict therapy response and if they preferentially screened and enrolled such patients for whom they expected a particularly favorable outcome. In this case one would expect the degree of selection (= the proportion of eligible vs. enrolled patients) to correlate positively with clinical outcomes. That was not the case, the correlation was almost zero (Spearman-Rho -0.04, p = 0.496, n = 364 [53]). Likewise, another analysis of 500 adult patients enrolled into AMOS in the period 1999-2005 showed no correlation between the degree of selection and the 0-6 month difference of Symptom Score [52]. These analyses do not suggest that the participating physicians' expectations of future therapy response led to any selection bias affecting clinical outcomes.

A limitation of the study is the absence of a comparison group receiving conventional treatment or no therapy. Accordingly, one must consider several other causes for the observed improvements apart from the AM treatment. We therefore conducted a sensitivity analysis of physician-
Because 14 clinical outcomes were analysed (Table 2), the issue of multiple hypothesis testing arises [38]. However, eight of the 14 comparisons at six-month follow-up showed significant improvements and six comparisons had p-values ≤ 0.001 – a constellation that would not be expected to occur by chance. (E. g. a Bonferroni adjustment for 14 tests would have indicated p < 0.004 as significance level).

Since AM treatment was to be evaluated under routine conditions, therapy was administered at the discretion of the physicians and therapists and not according to standardized protocols. This raises the question of whether study interventions would be replicable in future studies. However, AM therapy providers worldwide are trained according to highly standardized curricula, specifying e. g. individual eurythmy movements for specific diseases, constitution types, and movement patterns. Therefore, relevant therapy differences across settings would not be expected. Moreover, in this study, any local therapy differences would probably be offset by the large number of participating AM physicians and therapists.

The present analysis was restricted to clinical outcomes and did not assess costs. In an analysis of 487 adult patients from the AMOS study with migraine and other chronic indications, costs of AM medications and therapies in the first study year amounted to 12% of all costs of healthcare and productivity loss [57]. These extra costs of the new AM treatment in AMOS were offset by a reduction of inpatient hospital costs [29, 57]. Other, comparative evaluations indicate similar or lower costs in AM therapy settings, compared to conventional settings [19, 58].

This study provides the first data on comprehensive AM treatment for migraine in AM settings. The female/male ratio in this predominantly primary care sample (8.0/1.0) was higher than among migraineurs in the population (3.0/1.0) [59] and also higher than in primary care studies of AM medications for migraine (1.6-3.1/1.0) [60, 61]. The higher proportion of women in this sample might possibly reflect that women are more likely than men to engage in creative therapies such as AM art and eurythmy therapy. The use of non-opioid analgesics (44% of patients), 5-HT1 agonists (14%), and prophylactic antimigraine agents (14%) was similar to the use among migraineurs in Germany [62, 63] or France [64] during the study.

Baseline migraine disability, assessed by the number of days with headache in the past month (median 5.0 days), was higher than in the German population with migraine (median 2.2 days) [65], but the proportion of patients with >10 days with headache was similar (9% vs. 12%) [63]. Quality of life at baseline, assessed with the SF-36 Health Survey, was worse than in other migraine studies (median difference for all SF-36 scores in all evaluable studies: 0.42 standard deviations) [66]. The proportion of responders at six-month follow-up (patients with 50% improvement of Average Migraine Severity from baseline: 45-52%) was of the same order of magnitude as in trials of prophylactic antimigraine medications [6]. Notably, in prophylactic trials a response was defined as 50% reduction in number of migraine attacks, which was not documented in the present study; therefore this comparison is necessarily approximate. A systematic comparison of identical quality of life outcomes (SF-36) in migraine cohorts receiving other therapies showed improvements of largely of the same order of magnitude as in this study [66].

Previous studies have found beneficial effects of single AM medications on migraine [19]. In accordance with these findings, we found improvement of migraine symptoms and quality of life under comprehensive AM treatment. A notable finding was that three-fourths of study patients did not use anti-migraine medications in the first six months after enrolment, apart from non-opioid analgesics. Some migraine patients will not profit from medication; other patients stop taking medications due to adverse reactions or reject them, e. g. because they are passive. Even non-medication therapies for migraine are not suitable for all patients: some therapies are also passive (e. g. acupuncture or chiropractics) and others may be perceived as monotonous (e. g. biofeedback). In this context, the AM approach, with recourse to artistic exercise therapies as well as medication and rhythmic massage therapy, may be a welcome therapy option for some migraineurs.

CONCLUSIONS

In this study, adult migraine patients under AM treatment had long-term reduction of symptoms and improvement of quality of life. Improvements were similar in patients not using antimigraine prophylactic medications. Although the pre-post design of the present study does not allow for conclusions about comparative effectiveness, study findings suggest that AM therapies may be useful in the long-term care of patients with migraine.

TRIAL REGISTRATION

Not applicable (this is not a clinical trial).
ABBREVIATIONS

± = Standard deviation
AM = Anthroposophic Medicine
AMOS = Anthroposophic Medicine Outcomes Study
IQR = interquartile range
SA = Sensitivity analysis

CONFLICT OF INTERESTS

Within the last five years HJH and GSK have received restricted research grants from Wala-Heilmittel GmbH and Weleda AG, manufacturers of AM medications, otherwise all authors declare that they have no potential conflict of interests.

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AUTHORS’ CONTRIBUTIONS

HJH, CMW, GSK, SNW, and HK contributed to study design. HJH, AG, AR, and HK contributed to data collection. HJH, RZ, and HK wrote the analysis plan, HJH and AG analyzed data. HJH was principal author of the paper, had full access to all data, and is a guarantor. All authors contributed to manuscript drafting and revision and approved the final manuscript.

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