Effect of a Multi-Dimensional and Inter-Sectoral Intervention on the Adherence of Psychiatric Patients

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Abstract

Background
In psychiatry, hospital stays and transitions to the ambulatory sector are susceptible to major changes in drug therapy that lead to complex medication regimens and common non-adherence among psychiatric patients. A multi-dimensional and inter-sectoral intervention is hypothesized to improve the adherence of psychiatric patients to their pharmacotherapy.

Methods
269 patients from a German university hospital were included in a prospective, open, clinical trial with consecutive control and intervention groups. Control patients (09/2012-03/2013) received usual care, whereas intervention patients (05/2013-12/2013) underwent a program to enhance adherence during their stay and up to three months after discharge. The program consisted of therapy simplification and individualized patient education (multi-dimensional component) during the stay and at discharge, as well as subsequent phone calls after discharge (inter-sectoral component). Adherence was measured by the “Medication Adherence Report Scale” (MARS) and the “Drug Attitude Inventory” (DAI).

Results
The improvement in the MARS score between admission and three months after discharge was 1.33 points (95% CI: 0.73–1.93) higher in the intervention group compared to controls. In addition, the DAI score improved 1.93 points (95% CI: 1.15–2.72) more for intervention patients.

Conclusion
These two findings indicate significantly higher medication adherence following the investigated multi-dimensional and inter-sectoral program.
Introduction

Non-adherence to medication is one of the main causes for unsuccessful pharmacotherapy in psychiatry [1]. For example, the risk of hospitalization for non-adherent patients with schizophrenia is reported to be four times higher than for adherent patients [2]. It is assumed that increasing adherence might be the most efficacious way to further improve health care outcome [3].

Adherence is influenced by social and economic factors, the health care team/system, the characteristics of the disease, disease therapies and patient-related factors, as defined by the WHO [3]. Successful interventions should ideally target several of these dimensions because single-faceted interventions have frequently failed to show an impact [4]. Besides affecting practical barriers to adherence (e.g. dosing frequency), perceptual barriers, including the patient’s individual risk-benefit-assessment, should also be addressed [5]. Thus far, this dimension has only been considered in a few studies [5].

Successful research aimed at improving adherence in psychiatric patients and at developing a program for daily clinical practice in the hospital is rare. Previous investigations mainly enhanced the patient’s knowledge and insight [6,7] and focused on outpatients [8,9], but did not combine the beneficial therapy simplifications [10] with individualized patient education. The hospital stay and the time after discharge are periods that are susceptible to major changes in drug therapy that lead to patient uncertainty [11]. Hence, the transition between inpatient care and the ambulatory sector should be a point of focus by health care professionals.

Therefore, we investigated the effect of a multi-dimensional and inter-sectoral program on the patients’ adherence using a controlled study design in a psychiatric inpatient setting. The program started during the stay and ended three months after discharge. The additional program was administered only to patients in the intervention group. It included the provision of individualized verbal and written information regarding the patient’s specific psychiatric medication and diseases with subsequent telephone calls after discharge that focused on the patients’ concerns about their pharmacotherapy as well as maintaining adherence. Additionally, a medication review targeting the simplification of the pharmacotherapeutic regimen was performed. Control patients received usual care that included the dispensing of medication by ward staff and group counseling about medication once during their stay. The patient’s individual adherence was measured by the self-report questionnaires "Medication Adherence Report Scale” (MARS) [12] and ”Drug Attitude Inventory” (DAI) [13] at admission, discharge and three months after discharge.

Materials and Methods

Trial design

The study was designed as a non-randomized, prospective and open trial with a control and subsequent intervention group. The consecutive design was chosen to avoid a “carry-over effect” of methods from the intervention to the control group. Therefore, a parallel design with randomization was not possible.
Setting and participants

The study was conducted on two open wards with a total of 44 beds at the Department of Psychiatry and Psychotherapy of the maximum care University Hospital of Erlangen, Germany, which provides a total of 1300 beds. The Department of Psychiatry and Psychotherapy consists of two closed and two open wards as well as an ambulatory clinic and day clinic. An interdisciplinary team including psychiatrists, psychologists, nurses, occupational therapists, social workers and physiotherapists collaborate to help the patients during an acute disease crisis and with reintegration into society thereafter. All psychiatric diseases are treated with a particular focus on depression and anxiety disorders. The treatment comprises psychopharmacologic medication and non-drug interventions such as daily light, occupational and movement therapy as well as weekly psychological therapy.

Patients eligible for inclusion were ≥ 18 years who took at least one drug for a psychiatric condition, stayed a minimum of seven days on the ward and were able to communicate in German. Patients who met the inclusion criteria and gave written informed consent were assigned to the control or intervention group depending on the date of their admission. The control patients were included between September 2012 and March 2013 and the intervention patients between May and December 2013.

Ethics statement

The Ethics Committee of the Friedrich-Alexander University Erlangen-Nuremberg (FAU) approved the study protocol on 26.07.2012 (174_12 B). The trial was registered with the German Clinical Trials Register (DRKS00006358) and can be accessed via https://drks-neu.uniklinik-freiburg.de. The study was registered after enrolment of patients started because the responsible persons weren’t aware of the need to register before. No significant changes were made to the study protocol after approval of the ethical committee. The authors confirm that all ongoing and related trials for this intervention are registered.

Usual care

Usual care concerning medication included daily dispensing of medication by nurses in tablet boxes with four sections for morning, noon, evening and night medication respectively. The patient information leaflet is provided upon request of the patient. Medication can be a subject in the weekly medical consultation, but is not standard. General, non-individualized information about psychiatric medication is provided in a group session once during the stay.

Intervention

A multi-dimensional intervention was chosen due to its reported superiority to single-dimensional interventions [4,14,15]. The intervention was designed to create a program that addressed more than one of the five aforementioned factors influencing adherence [3]. The resulting multi-dimensional program focused on patient- and medication-related factors.

Since, according to the published literature, [16–18] insight is one of the most important factors that improves adherence in mental health, the program was based on a psychoeducational intervention. However, because standardized psychoeducation alone is not always efficacious, the intervention was individualized and combined with a simplification of drug regimens [19]. Additionally, phone calls provided closure to the inter-sectoral gap.[20]

The multi-dimensional and inter-sectoral program included 1) medication management with a focus on a simplification of therapy regimen, 2) counseling of patients during the hospital stay with verbal and written information and a medication plan at discharge, 3) phone calls...
after discharge. As recommended in the literature [1,21,22], the intervention was personalized to meet the individual requirements of each patient. For example, the description of the side effects in the information leaflets was adjusted for every patient.

1. Weekly, two pharmacists on the ward conducted a chart review to assess reasonable simplifications of medication regimens. Simplifications included reducing dosing frequency when possible, discontinuing unnecessary drugs and adjusting the times that drugs were taken in order to achieve optimal efficacy, few side effects and a simple therapy regimen. The simplifications were adjusted to the individual needs of each patient. In the intervention phase, these recommendations were discussed with the attending ward physicians who, together with the patient, decided the final implementation of the changes. For example, a common simplification concerned the once-daily dosage of Venlafaxine as a retard preparation that had been frequently administered twice daily prior to the intervention period.

In addition, the two pharmacists conducted a medication management to detect and solve drug-related problems, including drug-drug interactions or inappropriate dosing. Further information is provided elsewhere (unpublished data Wolf et al, 2015).

2. All intervention group patients received an individual consultation about their respective psychiatric disease and drugs. The patients were counseled about indication, mechanism of action, doses, side effects, interactions and instructions for use by a pharmacist. Additionally, the patients were informed about the delayed onset of effect and, if applicable, about the necessity to take the drug in order to prevent a relapse after recovery. Every patient received individualized written material based on the current state of the literature [23]. The duration of the consultation was adjusted based on the patient’s prior knowledge and desire to ask questions.

At discharge, the pharmacists provided intensive discharge counseling including a medication plan with information that included the trade name, active component, indication, required dosage and dosing schedule as well as administering information. The structure of the medication plan was based on the planned standardized medication plan for Germany [24].

3. During the follow up after discharge, the pharmacists telephoned the intervention group patients twice. The calls occurred 1.5 weeks and 6 weeks after discharge and were aimed at motivating the patient to maintain the pharmacotherapy, answering questions and being informed about possible changes made by the primary care provider. If necessary, for example, when a tendency toward suicidality was observed, a hospital psychiatrist was involved.

Outcome measures

The primary outcome was the change in adherence from baseline to follow-up three months after discharge as measured by the MARS [12]. The secondary outcome measure was the difference in attitude towards psychotropic drugs as determined by the DAI [13].

MARS. The MARS (Medication Adherence Report Scale) is a 5-item self-report questionnaire that asks about adherence in a continuous and sympathetic way [12]. It was translated into German, validated and considered reliable [13]. The MARS has been used to assess non-adherence over a variety of medical conditions including psychiatric diseases [25].

The patients score their own medication taking behavior on a Likert-scale ranging from ‘always’ (1 point), ‘often’, ‘sometimes’, ‘rarely’ to ‘never’ (5 points) in terms of the following parameters: forgetting medication, altering or omitting doses, taking less than instructed or
discontinuing the medication. In the end, scores are summed up with a maximum of 25 and a minimum of 5. A higher score indicates more adherent behavior. To calculate a change score as a percentage, the maximal possible difference of 20 points is defined as 100%.

Item 1 of the MARS assesses unintentional non-adherence (forgetfulness), whereas items 2–5 determine intentional non-adherence such as deliberate altering of the dose or suspending the medications.

A cut-off value is not defined by the original initiators of the MARS. However, cut-off values between 20 and 24 have been used previously [25–27]. By applying a cut-off value to the MARS scores, the percentage of adherent patients can be determined. This can be advantageous when comparing adherence rates measured by different methods. Due to the skewed distribution and its utilization in previous studies [28], a cut-off value of 24 was defined.

DAI. The Drug Attitude Inventory (DAI) [13] is a widely used and well-established self-rating instrument that assesses a patient’s attitude towards psychotropic drugs. The DAI score is associated with the degree of non-adherence [29]. Although the DAI was primarily designed for antipsychotic drugs, it is by now also used for a variety of psychiatric diseases such as depression [30], thereby justifying its use in this trial.

The DAI originally contained 30 items, but a shortened version with 10 items that are considered to be predictive for the entire set, was developed. Both versions show good test-retest reliability and discriminative validity as well as a high internal consistency [31,32]. The DAI-10 is more frequently used due to its simplicity and equal psychometric properties.

The DAI consists of 10 statements by patients with schizophrenia about their medications and includes subjective feelings as well as attitudes [13].

For every statement, the patients score whether it is true or false according to their own experiences with their drugs during the last four weeks. An answer indicating a positive attitude scores +1 point whereas an answer suggesting a negative attitude scores -1 point. A positive or negative sum score demonstrates a positive or negative attitude associated with (non-) adherence, respectively.

Collection of baseline and outcome data
Baseline data was collected after written informed consent of the patient and enrollment in the study. The pharmacists conducted a patient interview to assess demographic details, the therapy regimen and baseline adherence values measured by the MARS and DAI.

At discharge and three months after discharge (follow-up), the current medication profile for each patient was recorded and assessed by a pharmacist. The MARS and DAI were completed by the patients again. The data collected three months after discharge was usually obtained by telephone unless the patient wanted to return to the hospital to talk in person.

Statistical analysis
The sample size estimation was based on a study conducted by Finley et al that reported a 15% difference of adherence in control and intervention groups [33]. A minimum sample size of 123 patients was computed to detect a difference that was statistically significant at the 5% level with a power of at least 80% [34]. The study of Finley et al was chosen because it involved a similar program consisting of medication management, patient education and telephone follow-up [33].

Descriptive statistics are presented as the mean with standard deviation or the median with interquartile ranges for continuous variables. Categorical variables were presented as numbers and percentages. To check for group differences due to the allocation by time of admission,
Chi-Square-Test or Fisher’s-Exact-Test were applied for categorical variables, while Student’s-
t-Test and Mann-Whitney-U-Test were used for continuous variables.

The analysis of the primary and secondary outcomes was conducted following the inten-
tion-to-treat principle. Missing outcome values were imputed with the last valid observation of
the patient (e.g., the baseline value). A sensitivity analysis showed that this imputation
approach was more conservative than a corresponding analysis of the complete sample that
resulted in higher effect sizes (results not shown).

For both endpoints (MARS and DAI), statistical regression analysis was performed to esti-
mate the effect of the intervention while adjusting for group differences at baseline. The change
of the outcome variables between baseline and follow-up served as response variables, while
the grouping variables, age, sex, comorbidities, number of medications at admission and the
baseline score served as predictors. The estimated coefficient for the group variable therefore
represents the adjusted effect of the intervention and was reported with a corresponding 95%
CI. This procedure was repeated for both sum scores as well as for the single items.

Statistical analyses were performed using IBM SPSS Statistics for Windows Version 22.0
(SPSS Armonk, NY, USA: IBM Corp.) and the statistical programming environment R 3.0.2 (R
Foundation for Statistical Computing, http://www.R-project.org/).

**Results**

**Baseline characteristics**

A total of 269 patients were included. 136 patients admitted between September 2012 and
March 2013 and who met the inclusion criteria were allocated to the control group. Three
patients were discharged prior to the first pharmaceutical interview and consequently were not
included in the analyses of baseline and outcome variables. Recruitment for the intervention
group began in April 2013 and ended in December 2013. 133 patients were included in the
intervention group. Again, two patients were discharged early and therefore excluded from the
analysis. Mostly due to spontaneous discharge, the discharge assessment was not completed for
23 patients (15 control and 8 intervention). 109 control and 107 intervention group patients
were followed for three months after discharge. One patient did not take any medications
between discharge and follow-up. Consequently, the MARS and DAI score were not assessed
resulting in a total of 263 patients for analysis of the MARS. Six patients did not take psychiat-
ric drugs during the last four weeks before follow-up. The DAI was not assessed for them at fol-
low-up resulting in 258 evaluable questionnaires. All analyses were carried out following the
intention-to-treat principle with imputed values, if necessary, for 263 patients and 258 patients,
respectively. (Fig 1)

There were no significant differences between the two groups regarding demographic data
such as marital and work status, nationality or living arrangements. Furthermore, the two
groups did not differ significantly in terms of disease- and medication-related variables. The
control group consisted of more patients with schizophrenia (F2, 9.8%) and fewer patients
with anxiety disorders (F4, 15.0%) compared to the intervention group (6.9% and 19.8%,
respectively). Additionally, intervention group patients stayed longer in hospital than control
group patients. However, these differences were not statistically significant. Furthermore,
patients in the intervention group were slightly older than the control patients. The intervent-
ion group consisted of a higher percentage of female patients and had more secondary diagno-
ses. These differences were considered in the statistical analysis. Mean MARS and DAI scores
were similar in both groups at admission. (Table 1)
Primary outcome measures

Self-reported adherence was measured by the MARS, which can range between 5 and 25 with a higher score indicating a more adherent behavior [12]. The mean (SD) MARS score of control patients increased during hospitalization from 22.23 (2.87) by 1.21 points (6.05%) to 23.44 (2.34). Three months after discharge, the MARS score decreased by 0.97 points (4.85%) near to its level at baseline, 22.47 (2.99). In the intervention group, the MARS score improved from 22.02 (3.42) at admission by 2.3 points (11.6%) to 24.34 (1.61) at discharge. This value declined by 0.59 points (2.95%) to 23.75 (2.08) at the follow-up. Considering the improvement of the MARS score from baseline to follow-up, the adjusted effect of the intervention was 1.33 points (95% CI 0.73–1.93). (Fig 2) Importantly, 53.44% of the intervention group patients in
| Characteristic                                | Control (n = 133) | Intervention (n = 131) | p-Value |
|----------------------------------------------|-------------------|------------------------|---------|
| **Gender (n women (%))**                     |                   |                        |         |
| Control                                      | 43 (32.33%)       | 61 (46.56%)            | 0.018a  |
| Intervention                                 |                   |                        |         |
| **Age (years (±SD))**                        | 45.44 (± 14.67)   | 49.10 (± 15.26)        | 0.048b  |
| **Nationality (n (%))**                      |                   |                        |         |
| German                                       | 120 (90.2%)       | 122 (93.1%)            | 0.393a  |
| other                                        | 13 (9.8%)         | 9 (6.9%)               |         |
| **Living arrangements (n (%))**              |                   |                        |         |
| living with family                           | 82 (61.7%)        | 89 (67.9%)             | 0.626d  |
| living alone                                 | 47 (35.3%)        | 39 (29.8%)             |         |
| living in institution/sharing a flat         | 4 (3.0%)          | 3 (2.3%)               |         |
| **Marital status (n (%))**                   |                   |                        |         |
| married                                      | 67 (50.4%)        | 68 (51.9%)             | 0.405d  |
| single                                       | 44 (33.1%)        | 33 (25.2%)             |         |
| divorced                                     | 17 (12.8%)        | 22 (16.8%)             |         |
| widowed                                      | 5 (3.8%)          | 8 (6.1%)               |         |
| **Work status (n (%))**                      |                   |                        |         |
| employee                                     | 37 (27.8%)        | 49 (37.4%)             |         |
| unemployed                                   | 25 (18.8%)        | 19 (14.5%)             |         |
| pensioners                                   | 21 (15.8%)        | 23 (17.6%)             |         |
| incapacitated                                | 16 (12.0%)        | 14 (10.7%)             |         |
| self-employed                                | 7 (5.3%)          | 6 (4.6%)               |         |
| official                                     | 7 (5.3%)          | 3 (2.3%)               |         |
| other                                        | 20 (15.0%)        | 17 (13.0%)             |         |
| **Education (n (%))**                        |                   |                        |         |
| high-level                                   | 50 (37.6%)        | 44 (34.1%)             | 0.604d  |
| intermediate-level                           | 72 (54.1%)        | 74 (57.4%)             |         |
| low-level                                    | 7 (5.3%)          | 3 (2.3%)               |         |
| other                                        | 4 (3.0%)          | 8 (6.2%)               |         |
| **Prior psychiatric hospitalisations (n (%))** |                   |                        |         |
| 0–1                                          | 73 (54.9%)        | 77 (58.8%)             | 0.523a  |
| ≥ 2                                          | 60 (45.1%)        | 54 (41.2%)             |         |
| **Number of drugs (Median (IQR))**           |                   |                        |         |
| at admission                                 | 4 (2–6)           | 3 (2–5)                | 0.769c  |
| at discharge                                 | 4 (2–6)           | 4 (2–6)                | 0.787c  |
| 3 months after discharge                     | 4 (2–6)           | 3 (2–6)                | 0.577c  |
| **Psychiatric Diagnosis (n (%))**            |                   |                        |         |
| Mood (affective) disorder (F30-F39)          | 89 (66.9%)        | 88 (67.2%)             |         |
| Neurotic, stress-related and somatoform disorders (F40-F48) | 20 (15.0%)        | 26 (19.8%)             |         |
| Schizophrenia, schizotypal and delusional disorders (F20-F29) | 13 (9.8%)        | 9 (6.9%)               |         |
| Mental and behavioral disorders due to psychoactive substance use (F10-F19) | 7 (5.3%)          | 5 (3.8%)               |         |
| Others                                        | 4 (3.0%)          | 3 (2.3%)               |         |
| **(Continued)**                              |                   |                        |         |
comparison to 31.06% of patients in the control group achieved the possible maximum of 25 points in the MARS indicating a perfect adherent behavior. (Fig 3)

Intentional and non-intentional non-adherence were increased similarly. The adjusted effect size for improvement in non-intentional non-adherence (item 1) was 0.27 points (95% CI 0.05–0.49), for intentional non-adherence (items 2–5) 0.96 points (95% CI 0.39–1.51). (Table 2)

Secondary outcome measures
Adherence in psychiatric patients is strongly affected by their attitude towards their medication [13]. Therefore, we used the DAI to investigate the self-reported attitude of our sample to further clarify the causes for non-adherence. The DAI ranges between -10 (negative attitude) and +10 (positive attitude) [13].

The mean (SD) DAI score increased in the control patients during their hospitalization from 1.08 (3.45) at baseline by 1.04 points (5.2%) to 2.12 (3.82) at discharge. After discharge, this value increased slightly until follow-up by 0.33 points (1.65%) to 2.45 (4.02). In the intervention patients, the DAI score improved from 1.63 (3.75) at baseline by 2.79 points (Table 2)

![Fig 2. MARS scores. Development of MARS Scores with median and interquartile ranges for control and intervention group from baseline to follow-up three months after discharge. MARS, Medication Adherence Report Scale.](image-url)
(13.95%) to 4.42 (3.35) at discharge and by 0.37 points (1.85%) to 4.79 (3.39) at follow-up. Conclusively, the adjusted effect of the intervention on the improvement of the DAI score to follow-up was 1.93 points (95% CI 1.15–2.72). At admission, 57.36% of the control group patients and 62.02% of the intervention group patients were classified as adherent (DAI > 0). In the control group, the percentage of patients regarded as adherent at discharge was 69.77% and three months after discharge, 72.09%. In contrast, the proportion of adherent patients in the intervention group increased to 87.60%, which was maintained until follow-up (= 87.60%). (Fig 4)

Regarding the individual items of the DAI, an (adjusted) effect can be observed for items 3 (“I take medications of my free choice.”), 6 (“I take medication only when I feel ill.”), 7 (“I feel more normal on medication.”) and 9 (“My thoughts are clearer on medication.”). (Table 3)

**Table 2. Description of the individual items of the MARS.**

| MARS item (n=133) | Baseline Follow-up | Intervention mean (SD) | Adjusted effect* | 95% CI |
|------------------|-------------------|----------------------|-----------------|-------|
| Control          |                   |                      |                 |       |
| I forgot to take them. | 4.24 (0.87) | 4.26 (0.88) | 4.37 (0.80) | 4.68 (0.57) | 0.27 | 0.05–0.49 |
| I alter the dose. | 4.36 (0.87) | 4.45 (0.79) | 4.31 (0.88) | 4.74 (0.59) | 0.32 | 0.13–0.51 |
| I stop taking them for a while. | 4.65 (0.66) | 4.67 (0.65) | 4.52 (0.80) | 4.84 (0.46) | 0.26 | 0.11–0.42 |
| I decide to miss out a dose. | 4.52 (0.75) | 4.58 (0.71) | 4.47 (0.80) | 4.78 (0.50) | 0.21 | 0.06–0.36 |
| I take less than instructed. | 4.45 (0.77) | 4.52 (0.76) | 4.35 (0.84) | 4.71 (0.61) | 0.26 | 0.09–0.43 |
| Intentional Non-Adherence (Sum Score 2–5) | 17.99 (2.72) | 18.21 (2.70) | 17.65 (3.12) | 19.07 (2.04) | 0.96 | 0.39–1.51 |
| Sum Score MARS | 22.23 (2.87) | 22.47 (2.99) | 22.02 (3.42) | 23.75 (2.08) | 1.33 | 0.73–1.93 |

CI, Confidence Interval. MARS, Medication Adherence Report Scale. SD, Standard Deviation.

*Estimated treatment effect from statistical regression model, adjusted for sex, age, comorbidities, number of medications at admission and the baseline MARS score.
Discussion

To our knowledge, this is the first controlled study that assessed the impact of a multi-dimensional and inter-sectoral program during hospitalization and after discharge on the adherence of psychiatric patients. We used the period of hospitalization for adherence enhancing interventions. In addition, the patients received telephone calls after discharge.

The complex intervention chosen for this study enhanced both non-intentional (item 1 of the MARS) and intentional non-adherence (item 2–5 of the MARS). Although the difference between the control and intervention groups is highly significant at discharge and three months thereafter, it is striking that during hospitalization 26% of the intervention and 52% of the control patients were non-adherent. A study conducted with patients hospitalized in a department of internal medicine with an average length of stay of 9.3±6.8 days suggests that about 23% of patients are at least partly non-adherent during hospitalization [35]. As inpatient non-adherence is rarely investigated, possibly because it is perceived to be absent, we can only hypothesize that the higher percentage in our study could be explained by the longer duration of hospitalization of psychiatric patients e.g., 38.73 [29.25] days in our trial for the whole study population. It is conceivable that an extension or intensification of the interventions would achieve an even higher number of completely adherent patients.

Previous research exploring socio-demographic predictors of adherence has provided inconsistent results so far. It is rather evident that increasing age contributes to a more adherent behavior until a threshold of about 75 years when forgetfulness and frailty impede adherence [5]. The few studies investigating the impact of gender on adherence provide inconsistent results with a slight trend towards women being more adherent [36]. Additionally, the exact
The impact of the psychiatric diagnosis on adherence is still indistinct [37,38], although the high rate of non-adherence in schizophrenia and its severe clinical impact have been well documented [39]. As generally chronic diseases have been associated with poorer adherence than acute diseases [3], the discrepancies in diagnoses between our control and intervention groups seem negligible. Although the statistical analysis has considered the statistically significant differences between the groups (age, gender and comorbidities), it cannot be excluded that the differences in terms of diagnosis or employment contribute to the better adherence in the intervention group.

It was interesting to observe that the adherence of control patients was increased during the hospitalization—possibly by the usual interventions of the ward staff such as group sessions about medication—but decreased after discharge to its original level. This implies that usual adherence supporting interventions are not completely sustainable. The valuable hospitalization time should be used to institute enduring interventions akin to the program described and assessed in this trial.

The difference in items 3 (“I take medications of my free choice”) could suggest a higher sense of participation in the decision making process in the intervention patients, which has been reported to increase adherence [40]. The improvement in item 6 (“I take medication only when I feel ill”) seems to demonstrate a better knowledge about drug instructions in the intervention patients, possibly a result of the counseling provided by the pharmacists. The differences in items 7 and 9 could be attributed to a better

Table 3. Description of the individual items of the DAI.

| DAI item                                                                 | Control mean (SD) Baseline | Follow-up | Intervention mean (SD) Baseline | Follow-up | Adjusted effect* | 95% CI                  |
|-------------------------------------------------------------------------|---------------------------|-----------|---------------------------------|-----------|-----------------|-------------------------|
| For me, the good things about medication outweigh the bad.              | 0.35 (0.52)               | 0.55 (0.56) | 0.44 (0.56)                    | 0.69 (0.44) | 0.03            | -0.11–0.40              |
| I feel strange, “doped up”, on medication.                             | 0.36 (0.61)               | 0.59 (0.66) | 0.54 (0.67)                    | 0.76 (0.45) | 0.24            | -0.04–0.51              |
| I take medications of my free choice.                                  | 0.66 (0.60)               | 0.76 (0.50) | 0.60 (0.70)                    | 0.93 (0.28) | 0.24            | 0.06–0.41               |
| Medications make me feel more relaxed.                                 | 0.75 (0.69)               | 0.18 (0.76) | -0.03 (0.72)                   | 0.22 (0.72) | 0.12            | -0.08–0.32              |
| Medications makes me feel tired and sluggish.                          | 0.06 (0.66)               | 0.29 (0.69) | 0.23 (0.68)                    | 0.45 (0.64) | 0.06            | -0.12–0.24              |
| I take medication only when I feel ill.                                | -0.16 (0.95)              | 0.03 (0.97) | -0.10 (0.96)                   | 0.75 (0.64) | 0.69            | 0.43–0.94               |
| I feel more normal on medication.                                      | -0.11 (0.74)              | -0.11 (0.82)| -0.14 (0.83)                   | 0.12 (0.86) | 0.26            | 0.04–0.49               |
| It is unnatural for my mind and body to be controlled by medications.  | 0.11 (0.79)               | 0.25 (0.82) | 0.19 (0.82)                    | 0.48 (0.74) | 0.15            | -0.06–0.36              |
| My thoughts are clearer on medication.                                 | -0.26 (0.71)              | -0.28 (0.78)| -0.28 (0.78)                   | -0.05 (0.83) | 0.26            | 0.07–0.46               |
| Taking medication will prevent me from having a breakdown.             | -0.19 (0.76)              | 0.19 (0.81) | 0.19 (0.84)                    | 0.43 (0.74) | 0.08            | -0.12–0.28              |
| Sum Score DAI                                                           | 1.08 (3.45)               | 2.45 (4.02) | 1.63 (3.75)                    | 4.79 (3.39) | 1.93            | 1.15–2.72               |

CI, Confidence Interval. DAI, Drug Attitude Inventory. SD, Standard Deviation.
*Estimated treatment effect from statistical regression model, adjusted for sex, age, comorbidities, number of medications at admission and the baseline DAI score.

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response to pharmacotherapy, however a clinical outcome that would confirm this suggestion was not assessed.

Though the MARS [12] and DAI [13] are available as valid and reliable instruments, controlled studies in psychiatry that employ them are rare. De Las Cuevas et al used the DAI in a sample of depressed patients [41]. Firstly, they used the Morisky questionnaire to divide the study population into adherers and non-adherers. Then, they assessed a mean DAI score of 4.92 for adherent patients, which is comparable to the score measured in our study (4.79 at follow-up). An observational study by Medina et al reported a mean DAI score of 2.1 at discharge in patients with schizophrenia and bipolar patients [42], which corresponds to the DAI score of control patients (2.12) in our trial. Even though a high percentage achieved a DAI score > 0, the absolute scores of (non-) adherent patients remain rather low, between 2 and 5. This might indicate, in concordance with Brain et al, that the optimal DAI cut-off value to distinguish between adherers and non-adherers could be around 4 [43].

Canales et al conducted a controlled study with a similar interventional program in an American psychiatric inpatient setting, but did not follow patients after discharge [44]. They provided pharmacy services, including weekly medication reviews, pharmacotherapy recommendations and medication education. They did not assess adherence, but focused on several outcome parameters. Clinical response in the intervention patients was better, and there were less adverse effects. Even though the transferability of the results to the German health care setting and vice versa is not entirely possible, it can be assumed that the program’s benefits would extend beyond adherence to include clinical outcomes.

There are several limitations concerning this study. First of all, the chosen method for measuring adherence was a self-report questionnaire. A patient’s self-report is an indirect and subjective approach, and thus a less valid way of measuring adherence compared to direct methods [10]. Nonetheless, self-report is widely used due to its cost-effectiveness and practicality [9]. Velligan et al even asserted that data from subjective measurements were best correlated with the clinical state, thereby justifying the use of the MARS [45]. Jonsdottir et al demonstrated that the adherence measured by MARS correlated significantly to serum concentration of medication, but also ascertained a poor specificity [46]. The DAI is also reported to be a predictor of adherence measured objectively by MEMS [43].

Another principal limitation of the study is that it was designed as a non-randomized trial. Seasonal fluctuations and rotating physicians influence the patients’ behavior and lead to a possible over- as well as under-estimation of the intervention’s effect size.

Another limitation concerns the choice of the interventional measures. Although psychoeducation on disease and drugs and simplification of drug regimens before discharge are no standard procedures in the study setting, many other hospitals have already included them in their routine. To enhance comparability and to avoid retesting of approved methods, it would have been favorable to shift the psychoeducation and simplification of drug regimens to usual care.

The study time was too short for measuring relapse and subsequent readmission to the hospital [9]. However, we hypothesize reduced relapse and readmission rates following the intervention because more adherent behavior has been shown to relate to improved clinical outcome [47], including reduced rates of relapses and re-hospitalizations [48,49]. It was beyond the scope of our trial to determine the impact of the intervention on these outcomes. Because numerous interventions fail to change the adherence due to narrow or non-individualized interventions [4], the results of this trial support a successful method that should be further investigated with respect to clinical and economic outcomes.

It would have been favorable to divide the study population into subgroups to determine the impact of the single interventions in comparison to the whole program. Since this was not
carried out, the identification of the most efficacious component of the program is not possible. However, it has already been established that standardized psychoeducation alone is not effective for promoting adherence [14]. We speculate that the promising element of individualization in our psychoeducational program is a main factor underlying the observed effect.

In conclusion, it can be determined that the implementation of a multi-dimensional and inter-sectoral program enhances the patients’ adherence significantly up to three months after discharge. Considering the strong need for successful interventions to improve adherence among psychiatric patients, further studies are needed to explore the impact on clinical and economic outcomes.

Supporting Information
S1 File. Trial protocol as accepted by ethics committee_German. (PDF)
S2 File. Trial protocol as accepted by ethics committee_English. (PDF)
S3 File. CONSORT Statement. (PDF)

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Author Contributions
Conceived and designed the experiments: AP CW JK KF. Performed the experiments: AP CW. Analyzed the data: AP CW KF AM BL. Wrote the paper: AP CW AM BL JK KF.

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