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A randomized controlled trial of the efficacy and cost-effectiveness of a brief intensified cognitive behavioral therapy and/or pharmacotherapy for mood and anxiety disorders: Design and methods

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Publication Details

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**Methods:** Presented are the rationale, design, and methods of a two-armed randomized controlled trial comparing 'treatment as usual' (TAU) with a brief intensified cognitive behavioral therapy (CBT) and/or pharmacotherapy. Eligible participants (N=500) of five Dutch outpatient Mental Healthcare Centers are randomly assigned to either TAU or to the experimental condition (brief CBT and/or pharmacotherapy). Data on patients' progress and clinical effectiveness of treatment are assessed at baseline, post-treatment (3 months after baseline), and at 6 and 12 months post-treatment by Routine Outcome Monitoring (ROM). Cost analysis is performed on the obtained data.

**Discussion:** Since few studies have investigated both the clinical and cost effectiveness of a stepped-care approach intervention and a shortened diagnostic ROM method in both anxiety and/or mood disorders within secondary mental health care, the results of this study might contribute to the improvement of (cost)-effective treatment options and diagnostic methods for these disorders.

**Keywords**
controlled, trial, efficacy, cost, effectiveness, brief, randomized, intensified, mood, cognitive, behavioral, therapy, pharmacotherapy, methods, design, disorders, anxiety

**Disciplines**
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**Keywords**
Cognitive behavioral therapy; Brief therapy; Cost-effectiveness; Anxiety and mood disorders; Routine Outcome Monitoring
1. Introduction

Anxiety and mood disorders belong to the most common mental disorders. Several epidemiological surveys found anxiety and mood disorders to be the most prevalent class of mental disorders in the general population [1,2]. The World Health Organization surveys estimated their global lifetime prevalence to be 14.3% and 10.6%, respectively [1,3]. These disorders frequently start early in life and tend to have a chronic or relapsing course [4]; moreover, their presence contributes to a high disease burden for both the patient and their family [3]. They also have a substantial impact on daily functioning at home/work and on quality of life [5-8], comparable to the impact and effects of major chronic illnesses [5,9,10]. Consequently, the economic costs of these disorders for healthcare systems and society are high [11]. It is estimated that in 2010 the direct and indirect costs of anxiety and mood disorders in Europe were 74.4 + 113.4 billion euro, respectively [12]. Earlier studies in the USA and UK reported even higher estimates [13].

In the last decades, evidence-based treatments for anxiety and mood disorders have become available, i.e. cognitive behavioral therapy (CBT) and interpersonal therapy specifically developed for these disorders, as well as pharmacotherapy (mainly antidepressants) [14,15]. As important differences in effectiveness between the treatments are absent [13] guidelines were developed advocating a stepped-care model [16,17]. Moreover, to enhance effectiveness, for each of these treatments protocols and guidelines became available based on
those used in randomized controlled trials (RCT) [4,18-20]. Recently, we added Routine Outcome Monitoring (ROM) to the stepped-care approach to help the diagnostic process and treatment evaluation [19,20].

It is clear that guidelines, protocols and ROM have the potential to improve treatment efficacy. However, these improvements may not yet be fully realized as adherence to the guidelines and protocols remains questionable [17,21-23], even when ROM is added [19,20]. This implies that treatments last longer, consist of too many sessions and, thus, unnecessarily prolong suffering and increase related costs [17,21,22]. Moreover, the current economic situation offers a strong incentive to make treatments as cost-effective as possible. This applies not only to the treatments as such, but also to ROM; from an economic point of view ROM should be as concise as possible.

As most patients are treated in the first phase of the stepped-care model, it is in this phase that cost reduction is most profitable. Brief therapy is suggested to be suitable as a first step in a stepped-care model [22,24].

This paper describes an RCT designed to evaluate the effects and costs of a shortened first treatment in the stepped-care protocol for anxiety and mood disorders in secondary care, an area where there is a paucity of research.

It consists of time-limited (brief) CBT and/or medication treatment using a protocol following the multidisciplinary guidelines, but confined to a 7-week period and a maximum of 7 sessions. Also examined is the feasibility of a shorter, less labor-intensive ROM. In addition, the cost-effectiveness of the treatment and the adapted ROM are compared to ‘treatment as usual’ (TAU).
2. Design and Method

2.1. Study goals

Primary aim of this study is to evaluate the efficacy and effectiveness of a newly developed time-limited (brief) therapy intervention compared with TAU. Secondary aim is to evaluate the cost-effectiveness of the experimental intervention as compared with TAU. Additionally, the feasibility of a shortened, less work intensive and time-consuming ROM is evaluated. Patient and therapist satisfaction with the new intervention is also explored.

2.2. Study design

The study is a pragmatic, two-armed RCT using a parallel group design. Five Dutch mental health clinics are projected to enroll a total of 500 participants over an 18-month period of active recruitment. Eligible participants who provide informed consent are randomly assigned to one of two groups: the control group (TAU), or the experimental group. Patients in both groups are assessed by ROM at baseline and after 3 months (post-treatment). Follow-up assessments are conducted in all patients at 6 and 12 months post-treatment.

Primary and secondary outcomes are assessed by ROM. Primary outcomes are the scores on the Web Screening Questionnaire (WSQ) and Brief
Symptom Inventory (BSI). Secondary outcomes are the scores on the other instruments assessed by ROM (see section 2.8.1 and Table 1). The design and methodology of this study allows to assess analysis of equivalency (non-inferiority), since we do not expect to find the introduced intervention to be superior to TAU.

2.2.1. Control Group (TAU)

Individuals assigned to the control group receive standard psychiatric treatment called; Treatment As Usual (TAU). TAU varies across centers depending on the current activities at the participating MHCs. In MHCs, TAU is not strictly formalized; a multidisciplinary team is free to assign a therapy from a wide range of evidence-based therapeutic options (including: pharmacotherapy and psychological treatment, psychosocial interventions, contact with a psychiatric test nurse) according to the stepped-care approach. The treatment decision is based on professional experience, taking into account the specific problems and characteristics of the individual patient. The number of sessions depends on the therapy that is offered and can be weekly or (almost always) at a lower frequency of sessions, and are not confined to a maximum of sessions.

2.2.2. Experimental Group

The experimental group receives a brief, intensified cognitive CBT and/or pharmacotherapy confined to a fixed time period (7 weeks) and a limited number of weekly sessions (maximum 7 sessions within 7 weeks). The offered CBT and
pharmacotherapy are described in more detail in section 2.9. An intake and outtake session are also involved when in the experimental group (described in section 2.8.2).

2.3. Study setting

The study is conducted at five outpatient mental health clinics from the Dutch Regional Mental Health Provider (RMHP) Rivierduinen (RD). RD provides secondary mental health care for an area with over one million inhabitants. In the Netherlands access to mental health care is easy and is not limited by insurance or the financial means of the individual patient. Health insurance is compulsory for all citizens and regulated by the government [25,26]. The Dutch mental healthcare system is organized in a stepped-care manner and uses evidence-based treatment guidelines. According to a stepped-care approach a brief but intensive first step is offered and patients who are insufficiently helped by the initial intervention are allowed to ‘step up’ to subsequent treatment [16]. The therapeutic principles within the treatment protocols of the intervention are referred to as a ‘first step’ of a stepped-care approach.

2.4. Participants

Eligible participants are males and females aged 18 to 65 years, currently diagnosed with an anxiety and/or depressive disorder as main diagnosis. Patients with current psychotic or bipolar traits, homicidal or suicidal risk or
severe social dysfunction, as diagnosed by their general practitioner (GP), psychiatrist, or as assessed in a diagnostic interview, are excluded. All eligible subjects need to have adequate understanding of the Dutch language.

Patients with the following DSM IV [27] diagnoses are therefore included: minor or major depressive disorder (single episode or recurrent), depressive disorder NAO, dysthemia, panic disorder (with or without agoraphobia), panic disorder NAO, social phobia, simple phobia, generalized anxiety disorder, obsessive compulsive disorder, posttraumatic stress disorder (type I or single trauma), adjustment disorder (with anxiety and/or depressive mood). Co-morbidity associated with other psychiatric diagnoses (with the exception of psychotic or bipolar disorder) is allowed in order to establish a clinically relevant, broadly representative sample.

2.5. Sample size

The sample size was calculated using the method of Cohen [28] and based on review of the available literature of earlier comparable studies. We aimed at detecting an equivalence with an acceptable difference of 5% on the primary outcome measures WSQ and BSI (see section 2.8.1 and Table 1) and a 15% maximal difference in outcome scores between TAU and the intervention under the usual assumptions of an $\alpha = 0.05$ and power of 80%. This results in an intended total sample size of 500 participants.
2.6. Recruitment, screening and enrolment procedures

All patients referred by their GP to one of the participating MHCs for the treatment of anxiety and/or mood disorders are, initially, eligible to participate in the study.

We adopted the following steps in recruitment: first, all referred patients are globally screened by an experienced psychiatrist for the presence of depression and/or anxiety disorders as current, main problem. This global screening is based on written information provided by the GP containing an interpretation of the patient's current health status and referral for further mental health care; this step does not require face-to-face contact with the patient. Subsequently, the potentially eligible patients are invited for a first ROM assessment. Prior to this first ROM assessment, the psychiatric research nurse conducting the ROM assessment invites the patients to participate in the study. Those who agree to participate are asked to provide written informed consent.

When informed consent is given, the baseline ROM assessment (T₁) according to the study design is conducted. After completion of this assessment, participants’ randomization by the research team takes place (see section 2.7). Depending on the randomized treatment condition, final eligibility is assessed during the subsequent intake session by means of the inclusion and exclusion criteria of the study.

Patient enrollment began March 1, 2010 and will end December 31, 2011. Follow-up assessment is ongoing and is projected to continue until December 31, 2012.
2.7. Randomization and blinding

After successful screening, provision of written informed consent and completion of the baseline measurement (T1); (see section 2.6), all eligible participants in this RCT are randomly assigned to one of two groups: the experimental group or the control group (TAU). Random allocation was generated by using a variable blocked design developed by an independent researcher from the Department of Medical Statistics & BioInformatics, LUMC and derived by computer. Randomization takes place on the individual level by clinical center (n = 5) and gender. This procedure is used to increase the likelihood that the distribution between groups is balanced on the two potentially important confounding variables and to conceal random allocation sequence. Participants and clinicians are informed about the outcome of the randomization; the psychiatric test nurses (assessors) involved in the ROM assessment in the study, are kept blinded to the randomization condition throughout the entire study.

Randomization and the subsequent assignment of participants to the intervention will be performed by the researcher (D.M.), whom is not an assessor.

2.8. Assessment

For both treatment conditions assessment information is obtained in two-fold, as shown in Figure 1.
First, patients participating in this study are assessed by ROM (see 2.8.1.) at four time intervals: 1) $T_1$ at baseline (start of study), 2) $T_2$ immediately post-treatment (3 months after baseline measurement), 3) $T_3$ 6 months post-treatment (first follow-up measurement), and 4) $T_4$ 12 months post-treatment (second follow-up measurement). The second assessment method is provided by an intake and outtake evaluation (see also 2.8.2).

The timetable of assessments is shown in Figure 1.

2.8.1. Routine Outcome Measures and feedback

This study is conducted on data collected using ROM [19]. ROM is a monitoring system for patient care, implemented in 2002 in the outpatient clinics of RD and the Department of Psychiatry of the LUMC. All outpatients referred to these clinics by their GP for treatment of a mood, anxiety and/or somatoform disorder, are assessed by ROM.

ROM periodically measures the presence and severity of psychiatric symptoms in patients and thereby monitors patients' progress/changes during treatment by conducting an extensive battery of psychometric instruments. An overview of the instruments used in ROM is available at http://www.lumc.nl/psychiatry/ROM-instruments.

These instruments are routinely assessed at baseline and during treatment at several time points [19,29]. The baseline assessment also comprises a standardized diagnostic interview (Dutch version of the Mini-
International Neuropsychiatric Interview Plus, version 5.0.0) [30,31], the collection of socio-demographic and socio-economic data, and the administration of general measures of health and disease-specific severity scales.

Instruments are both self-report and interviewer based. All interviewer-based instruments are carried out by a psychiatric research nurse and the self-report questionnaires are completed using a touch-screen computer. A web-based software QuestManager (www.psyquestmanager.nl) was developed to assist the ROM method and is also used in the current study.

Data collected by ROM are provided to the clinician and patient as written feedback (a brief report) by the psychiatric research nurse. This written feedback consists of an overview of the main measurement results. Furthermore, a summary of the diagnostic interview and a summary of the main questionnaires is given (one or two pages). The clinician shares and discusses these results with the patient. The assessment outcomes are used to support decision-making for the future course of the treatment [19,29].

For the present study, all eligible patients are routinely assessed by ROM. Table 1 lists the instruments used to assess the disorders of interest for the current study. To further test the hypotheses of the present study, five additional questionnaires are added to the regular ROM and are indicated (in bold print) in Table 1.

- Insert Table 1 here.
Similar to the regular ROM, the baseline assessment comprises the standardized diagnostic interview, MINI-Plus 5.0.0, and the collection of socio-demographic and socio-economic data.

For the present study, the written ROM feedback depends on the randomized treatment condition. Clinicians in the experimental group are provided with minimal information about the results of the questionnaires, i.e. an overview of the results of the patient’s performance on the two primary outcome measures: the Web Screening Questionnaire (WSQ) [32] and the Brief Symptom Inventory (BSI) [33]. The summary of the diagnostic interview and information about the other measures is left out. Clinicians in the TAU group are provided with the regular, extensive, feedback about all measurement results.

2.8.2. Intake and outtake

In addition, to examine the progress of patients’ wellbeing and the effectiveness of the treatment, an intake and outtake session is included in the study design (Figure 1). Both intake and outtake session are semi-structured and conducted by the same experienced psychiatrist (see 2.8.5), during one 45 minute (approximately) session.

The intake takes place before the start of the treatment, after the ROM baseline measurement. A semi-structured clinical interview, especially designed for this study, is administered. Compared to a regular intake, this intake session is protocolized, shorter and more structured. Moreover, the intake aims to ensure that the patient is eligible to participate in this study by means of the in- and
exclusion criteria (section 2.4). Personal data, including demographics and the current clinical picture, are also obtained during this intake session.

The outtake evaluation takes place within (maximally) 2 weeks post-treatment (Figure 1) and is part of a stepped-care approach. The aim of this semi-structured outtake session is to evaluate if patients are sufficiently helped by the offered initial intervention or ‘stepping up’ to subsequent (additional) treatment is necessary. The ‘stepping-up’ is according to clinical experience and the local and national guidelines as handled by the involved MHC. During the outtake the progress of the patient’s symptoms and his/her current clinical status will be assessed. Furthermore, it evaluates the patient’s wellbeing and an ‘end diagnosis’ is formulated. A possible subsequent treatment plan can be discussed by patient and clinician.

When the outtake session demonstrates that the achieved treatment effect is insufficient, the patient is offered to ‘step up’ to additional treatment according to the treatment guidelines in the same MHC, or elsewhere.

2.9. Treatment

For the current study the following treatment protocols are formulated by the research team:

1. Pharmacotherapy protocol for mood and/or anxiety disorders (maximum 4 sessions within 7 weeks)

2. Brief Cognitive Behavioral Therapy protocol for depression (minimal 5, maximum 7 weekly sessions)
3. Brief Cognitive Behavioral Therapy protocol for anxiety (minimal 5, maximum 7 weekly sessions)

4. Eye Movement Desensitization and Reprocessing (EMDR) therapy protocol for post-traumatic stress disorder (PTSD) (maximum 6 sessions within 7 weeks)

The treatment decision was made by the patient and therapist, based on the professional experience of the therapist and taking into account the specific disorder and characteristics of the individual patient, thereby acknowledging the patient’s preferences as is good practice and according to the common accepted principles of shared-decision making [34,35].

The treatment protocols are based on the existing treatment guidelines described in the (multidisciplinary) guidelines in Dutch mental healthcare [4,36,37] and on acknowledged evidence-based literature [4,38-41]. All treatment protocols are evaluated during an outtake session (see section 2.8.2.).

Delivering treatment by combining the described protocols is also possible.

2.9.1. Pharmacotherapy for mood and/or anxiety disorders

In the current study the pharmacotherapy protocol for mood disorders and/or anxiety is characterized by a quick onset and aims to reach an optimal clinical effect of the used medication, involving rapid stepping up to the most optimal dose and minimizing the side-effects for patients as much as possible. Patients are treated with a selective serotonin reuptake inhibitor (SSRI) in 4 sessions
within a 7-week period. Medication use is evaluated after this fixed period and continued when necessary. The protocol provides a scenario for patients when not using an SSRI (SSRI 1 condition) or, on the other hand, a scenario for patients with a history of (sufficient and adequate) SSRI use and now starting with a new different SSRI (SSRI 2 condition). When patients are currently using an SSRI (as prescribed by their GP), medication is continued and when necessary adapted according to the pharmacotherapy protocol.

2.9.2. Brief CBT for depression

The brief CBT for depression protocol is focusing on decreasing the depressive mood of the client and maintain this improvement and based on the cognitive behavioral therapy manuals/protocols for depression [38,42].

For the current study this protocol is confined to a maximum of 7 weekly sessions (minimal 5 sessions) within a 7 week time-period. The first 3 sessions of this brief CBT protocol are dedicated to activation-enhancement and training of social skills. The last 4 sessions emphasizes tracing and altering irrational cognitions by challenging them [38,42]. Each treatment session consists of a 45-min face-to-face contact and is characterized by a quick onset (no waiting list).

2.9.3. Brief CBT for anxiety

The brief CBT protocol for anxiety disorder is also characterized by a quick onset (no waiting list) and a maximum number of sessions (minimal 5, maximum 7
weekly sessions within a 7-week period). Each treatment session consists of a
45-min face-to-face contact.

The main focus of the brief CBT for anxiety protocol is on the core CBT
techniques for the treatment of anxiety disorders: anxiety/tension-reducing
techniques, cognitive techniques and exposure techniques (used when
adequate) [40]. If dysfunctional worrying is interfering, anti-worrying techniques
are offered.

2.9.4. Eye Movement Desensitization and Reprocessing Therapy

The EMDR therapy for the PTSD protocol is characterized by a quick onset (no
waiting list) and a limited number of sessions (a maximum of 6 sessions within 7
weeks). Furthermore, all EMDR sessions comprise a 45-min face-to-face contact.
The EMDR protocol of this study is based on the Dutch Manual EMDR [43] and
approved principles described in the literature [39], or the treatment of PTSD for
patients suffering from a Type I trauma or single trauma.

2.10. Therapist selection, training and supervision

The treatment protocols are performed by experienced clinicians: psychiatrists
and psychologists working at the participating MHCs.

Clinicians working at RD provide treatment in accordance with the
multidisciplinary guidelines of the National Steering Committee describing
evidenced-based treatments for mood and anxiety disorders. The participating
clinicians are all professionally educated and trained in CBT. Years of experience
ranged between 1-7 years. They were especially trained to work with the brief treatment protocols within this study.

The participating clinicians were initially instructed by the research team. Two hours of instructions were provided for participating clinicians on two consecutive days. On the first day, a 2-hour instruction was provided for clinicians involved in the CBT condition. During this instruction, the EMDR protocol was also introduced. On the subsequent day, a 2-hour instruction was provided for clinicians involved in providing the pharmacotherapy within this study.

During these instruction meetings, consensus was achieved on a number of core elements of the treatments. Furthermore, the content of future supervision session for all involved clinicians was discussed.

Clinicians in the TAU group received no specific training from the research team. They provide the usual treatment to the patients; globally following the available multidisciplinary treatment guidelines, since adherence is questionable.

Furthermore, six certified psychiatric test nurses were trained by the research team to assess the ROM measurements according to the guidelines of the study. The research nurses are all trained and certified to assess ROM measurements.

Instruction was given on the administration and reporting of the ROM measurements designed for this study. The psychiatric test nurses were informed about the logistics of the study and how to apply the blinding method used in the present study.
Once every 3 months supervision sessions were organized by the research team for the clinicians and the psychiatric test nurses. Interview techniques, how to avoid protocol violations and other challenges were discussed and evaluated.

In addition, unrestricted support is provided to all study clinicians and psychiatric test nurses via email and through visits made by the research team.

2.11. Fidelity monitoring

All treatment sessions within the brief CBT protocols will be audio-taped, using a digital voice recorder to ensure treatment protocol fidelity. The taped treatment sessions will be randomly checked and scored on protocol consistency and reliability by the research team. Furthermore, therapist adherence and satisfaction will be monitored using an evaluation questionnaire. This evaluation questionnaire furthermore monitors the delivery and compliance of the different treatment protocols since it questions how many sessions were involved and if treatment was successfully delivered. This evaluation questionnaire, and the list of the criteria for protocol consistency and reliability, can be obtained from the corresponding author.

2.12. Statistical analyses

Descriptive statistics will be used to describe the characteristics of the two groups, and the outcome variables, at the four measurement points. To evaluate potential group differences at baseline, post-treatment and at the 6 and 12-month
follow-up measurements, repeated measures analysis will be conducted to analyze the short and long-term effects between the experimental and control group.

All analyses will be conducted according to the intention-to-treat (ITT) approach. Additionally, analysis per protocol will be conducted.

Chi-square analyses and t-tests for independent samples will be used for data measured on one occasion (e.g. patient satisfaction, therapist satisfaction, baseline demographic features) to detect possible differences between the two groups. Differences are considered statistically significant at $p < 0.05$.

Missing values will be imputed with regression imputation techniques. Generalized Estimating Equation (GEE) analyses will additionally be performed on the dataset with missing data and when missing values are imputed. Analyses adjusting for cluster effects will be performed when analyzing the data.

All analysis will be done using SPSS (version 17, Windows).

2.12.1 Health economics analysis

Cost-effectiveness will be also calculated. An economic evaluation based on the TIC-P questionnaire (see Table 1) examines the costs and other aspects of the study protocol. Direct costs per patient in the experimental condition versus the costs per patient in the control condition will be compared. Costs are validated by reference; if these references are not available, costs will be estimated by costs research.
Cost or product losses are verified by the ‘friction cost method’ [44]. The friction cost method estimates the indirect costs of disease, which explicitly considers economic circumstances that limit production loss due to disease. According to this method, these indirect costs mainly occur during the time it takes to replace a worker, i.e. the friction period [45].

2.13. Approvals and data/safety monitoring

The study protocol was approved by the Medical Ethical Committee (MEC) of the Leiden University Medical Center (LUMC). After full verbal and written information about the study, written informed consent was obtained from all participants at the start of baseline assessment. To safeguard the anonymity of the patients and to ensure proper handling of the data, processing of all data is in accordance with a comprehensive protocol: the Psychiatric University Network REgistration Leiden (PAREL-regulations). The MEC of the LUMC approved the regulations of this protocol and agreed with this policy [19]. Confidentiality of data is maintained by using a unique research identification (ID) number for each participant, which enables to identify individuals without using names. Only a limited number of persons (researcher) have access to the record that links the ID number to identifiable information.
3. Current status and demographics of sample

Data will be collected until at least December 31, 2012. At the time of completion of this paper, the inclusion period of the study is still ongoing. A total of 161 patients have completed the baseline measurement. Their preliminary socio-demographic and clinical characteristics are presented in Table 2.

> Insert Table 2 here.

These 161 participants are evenly distributed across both groups; 79 in the TAU group and 82 in the experimental group. Demographic data (educational and employment status, and ethnic background) are missing for 4 patients. Moreover, for 12 patients the MINI-Plus diagnostic interview did not lead to a DMS-IV classification in ROM; therefore, only a clinical diagnosis is available for these patients.

At baseline there were no significant differences in demographic data between the two groups. The mean age of the patients is 37.2 (SD 11.5; range 18-65) years, and there is a similar distribution for gender (61.5% female and 38.5% male) in both groups.

Furthermore, the participants comprise a relatively homogeneous group with common mental disorders, i.e. mainly mood and anxiety disorders. The majority of the included patients have more than one clinical diagnosis.

At baseline, of all patients 45.3% had an anxiety disorder only, and 37.9% were diagnosed with depression only. In total, 15.5% of the patients were
diagnosed with both an anxiety and mood disorder. This distribution was similar in both treatment arms (Table 2).

4. Discussion

Evidence-based clinical guidelines advocating a stepped-care approach are available in mental healthcare and have demonstrated success in the treatment of mood and anxiety disorders [46]. Progress within these stepped-care approaches is carefully monitored and patients are able to ‘step up’ when no subsequent improvement occurs [16].

However, initiation of and adherence to these recommended and effective treatments within these guidelines is usually poor [18]. The optimal content and organization of how to provide this stepped care is unclear, and implementation and acceptability of stepped care as a method of delivering psychological/psychiatric services has not yet been adopted [21,22]. Little information is available about how stepped care should be effectively implemented [22] and only a few randomized trials present convincing evidence and evaluations of both the cost and clinical effectiveness of this stepped-care model. Most studies investigated either the cost effectiveness [47,48] or the clinical effectiveness [49-52] of stepped care. When simultaneously investigated, the studies examined the effectiveness of a stepped-care model for patients with either a mood disorder [53] or an anxiety disorder [54], and only in primary care.

The present study aims to examine the clinical effectiveness and cost effectiveness of an innovative stepped-care intervention for patients (aged 18-65
years) with anxiety and/or mood disorders in secondary care. To carefully monitor the patient's progress, the ROM method is added.

To our knowledge this is the first long-term study to simultaneously analyze the clinical and cost effectiveness of a stepped-care approach in the treatment of both mood and anxiety disorders in a secondary care setting. The ROM method, monitoring patients’ progress at fixed time intervals with a follow-up period of 1 year, allows to establish the long-term efficiency expectations of treatment success and effects on health status and economic costs. Moreover, for the current study, ROM is shortened and evaluated in terms of feasibility. Since the baseline characteristics of the participants in both the experimental and the control group show no significant differences, the study outcomes are expected to be highly generalizable.

The findings of the present study will have potential implications for provision of the most convenient form of mental healthcare in the Netherlands. The offered time-limited, brief intervention could provide valuable information to help the development of an optimal treatment protocol for patients with anxiety and/or mood disorders. The limited number of sessions reduces the average amount of therapist input per patient and, moreover, is an adequate response to patient preferences for brief psychological interventions [55]. Since the intervention has not only fewer sessions but also an earlier start after intake, it is expected to reduce waiting lists; this is a common problem in the provision of mental healthcare.
Results of the analyses will allow to compare both the clinical (patient improvement) and cost effectiveness (economic costs of the intervention) between the experimental intervention and TAU. Besides answering our primary question (‘is the experimental intervention at least as effective as providing regular care, i.e. treatment as usual’) these analyses are expected to provide insight into how to increase the quality and efficiency of care. The experimental intervention is expected to reduce costs and increase efficiency on (at least) the short term. The study design will also provide insight into long-term effects, possibly encouraging the implementation of an effective stepped-care model in mental healthcare.

Apart from evaluation of the effectiveness of the intervention, the study also examines what works best for the individual patient. The broad inclusion criteria allow the recruitment of patients with a wide range of mood and/or anxiety disorders. The resulting sample is then broadly representative of the patient population commonly referred to outpatient mental healthcare centers in the Netherlands. No indication of a selection bias is expected and, since the ROM method outlines patient characteristics, the advantages of the experimental intervention for the individual patient can be clearly assessed.

A major strength of this study is that it is a pragmatic randomized trial. In such trials, patients and therapist are the same as those encountered in daily practice. Care is provided by healthcare professionals from the field and, since the sample of patients is the same as those seen in daily practice, this enhances the external validity of the study. Moreover, since the stepped-care algorithm
used covers the whole continuum from enrollment, diagnostics, assessments and
treatment, this study reflects the ‘real’ effects of daily practice thereby allowing
generalization and implementation of any beneficial logistical and organizational
effects in (clinical) practice.

However, this advantage also carries some risks. Besides the logistic
difficulties of implementing a pragmatic randomized trial, it is difficult to maintain
treatment integrity when conducting a study in daily practice. We aim to minimize
this limitation by means of our instruction meetings and supervision sessions for
all clinicians and psychiatric test nurses involved. Moreover, since all treatment
sessions within the brief CBT protocols are audio-taped, treatment protocol
fidelity is closely examined and, hopefully, achieved.

Moreover, since conducting a study in daily practice involves the
possibility of combining CBT and medication therapy and enhances the
possibilities of non-specific effects in therapy, the potential therapeutic benefit of
the intervention can only be formulated with caution.

While the design of the current study addresses many of the limitations of
previous research, a preliminary reflection on further limitations and strengths of
our design is required. Although homogeneous patient groups are expected in
both treatment arms, the control group may have some nonspecific effects on the
expected outcomes. For example, no control is made regarding the number of
visits made by persons in the TAU group to their physicians. Therefore, it is
possible that patients in the control group make fewer visits to their physician or,
in some cases, no visits at all if they are still on a waiting list. Although this does
reflect daily practice, we cannot rule out the nonspecific effects of an increased number of visits and/or attention from physicians as an explanation for the (possible) better outcomes in the experimental group.

Another limitation concerns the expected dropout rates at the four ROM measurement assessments, which may affect the results of the study. Although compliance with the ROM procedure is relatively successful, a 20% dropout rate is expected at reassessments. In response to these high attrition rates, the aggregated data are also analyzed according to the intention-to-treat analysis. This might yield a more valid reflection of the results and conclusions about the effectiveness of the experimental intervention.

We have described the rationale and design of an RCT examining the clinical and cost effectiveness of a time-limited, stepped-care based intervention in the treatment of mood and anxiety disorders. This study is collecting a substantial amount of data which will improve our understanding of how to develop effective strategies to adequately diagnose and treat patients with mood and/or anxiety disorders in secondary care.

If the experimental intervention proves to be as effective as regular care, this type of intervention could facilitate the growing need of providing the most optimal and (cost)-efficient mental healthcare. This study will hopefully elucidate how we can provide the most convenient and adequate mental healthcare for our patients.
Acknowledgments

This is a collaborative study between Leiden University Medical Center and Rivierduinen (a regional mental healthcare provider) and is funded entirely by Rivierduinen. The authors thank the staff at Rivierduinen for their contribution to this research, as well as all the patients participating in this study.
References


Fig. 1. Timetable of assessments during the study.

Note: T₁: baseline assessment; T₂: post-treatment assessment; T₃: first follow-up assessment; T₄: second follow-up assessment.
Table 1. ROM study measures by time interval

<table>
<thead>
<tr>
<th>Instrument</th>
<th>Full Name</th>
<th>Domain</th>
<th>Cluster</th>
<th>Typ</th>
<th>Time- interval</th>
<th>Reference</th>
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<tbody>
<tr>
<td>AGO</td>
<td>Agoraphobic Cognitions Questionnaire</td>
<td>Psychopath</td>
<td>Spec</td>
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<td>Spec</td>
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<td>BSI</td>
<td>Brief Symptom Inventory</td>
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<td>Gen</td>
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<td>Dutch Mental Healthcare Thermometer of Appreciation by Clients</td>
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<td>Gen</td>
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<td>Spec</td>
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<td>Gen</td>
<td>SR X X X X X</td>
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<td>Mini- International Neuropsychiatric Interview Plus 5.0.0.</td>
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<td>Gen</td>
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<td>Padua Inventory revised</td>
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<td>PSWQ</td>
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<td>Psychosoc Func</td>
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<td>Web Screening Questionnaire for common mental disorders</td>
<td>Psychopath Gen SR X X X X [32]</td>
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aSR=Self Report; OS=Observer Scale; bT1: baseline assessment; T2: post treatment assessment; T3: first follow-up assessment; T4: second follow-up assessment; cHealth Questionnaire H_Q (in Dutch: Gezondheidsvragenlijst); dMental Healthcare Thermometer (in Dutch: GGZ Thermometer)

Note: A list of all ROM measures is available at [http://www.lumc.nl/psychiatry/ROM-instruments](http://www.lumc.nl/psychiatry/ROM-instruments)
Table 2. Socio-demographic and clinical characteristics of the eligible patients at baseline.

<table>
<thead>
<tr>
<th></th>
<th>Control group (N=79)</th>
<th>Randomized condition Experimental group (N=82)</th>
<th>Total (N=161)</th>
<th>P-value</th>
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<td><strong>Age in years: mean (SD)</strong></td>
<td>N 37.2 (11.5)</td>
<td>N 36.0 (12.8)</td>
<td>N 36.6 (12.1)</td>
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<td><strong>Gender</strong></td>
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<tr>
<td>Male</td>
<td>32 40.5</td>
<td>30 36.6</td>
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<tr>
<td>Female</td>
<td>47 59.5</td>
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<td>Dutch</td>
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<td>5 6.4</td>
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<td><strong>Educational status</strong></td>
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<td>Lower education</td>
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<td>Higher education</td>
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<td>Employed</td>
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<tr>
<td>Other</td>
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<td>7 9.1</td>
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<td><strong>Marital status</strong></td>
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<td>Married/Cohabitating</td>
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<tr>
<td>Divorced/separated/widow</td>
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<td>Single</td>
<td>30 38.0</td>
<td>27 35.1</td>
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<tr>
<td><strong>DSM IV(^e) diagnosis (n, %)</strong></td>
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<tr>
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<td>2 1.2</td>
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<td>Post Traumatic Stress Disorder</td>
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<td>21 13.0</td>
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<td>.308</td>
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</tbody>
</table>

\(^a\)TAU=Treatment As Usual; \(^b\)CBT=Cognitive Behavioral Therapy; \(^c\)Demographic data; ethnic background, educational status and employment status are missing for 4 participants; \(^d\)Lower education= basic education, primary education, no education at all. Higher education= Higher education, university; \(^e\)DSM IV=Diagnostic Statistical Manual (of mental disorders) 4th version.

Note: patients may have more than one diagnosis