Cost-effectiveness of the mobile application TCApp combined with face-to-face CBT treatment compared to face-to-face CBT treatment alone for patients with an eating disorder: study protocol of a multi-centre randomised controlled trial

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Abstract

Background: The clinical utility of the existing apps for people with eating disorders (EDs) is not clear. The TCApp has been specifically developed for people with EDs, is based on the principles of Cognitive Behavioural Treatment (CBT) and allows a bidirectional link between the patient and the therapist. The objectives of the study are, first, to assess the clinical efficacy of a combined intervention for Eating Disorders (EDs) that includes an online intervention through the TCApp plus standard face-to-face CBT in comparison to standard face-to-face CBT alone, and second, to examine the cost-effectiveness of the TCApp and identify potential predicting, moderating and mediating variables that promote or hinder the implementation of the TCApp in ED units in Spain.

Methods: The study methodology is that of a randomised controlled trial combining qualitative and quantitative methods, with a 6-month follow-up. Approximately 250 patients over 12 years old with a diagnosis of an ED from several ED units in Spain will be randomised to one of two different conditions. Participants, their caregivers, healthcare professionals and technical staff involved in the development and maintenance of the application will be assessed at baseline (T0), post-intervention (T1) and at 6 months follow-up (T2). Primary outcome measures will include ED symptomatology while secondary measures will include general psychopathology and quality of life for patients, quality of life and caregiving experience for family caregivers and adoption-related variables for all participants involved, such as perceived usability, user’s satisfaction and technology acceptance. For the cost-effectiveness analysis, we will assess quality-adjusted life years (QALYs); total societal cost will be estimated using costs to patients and the health plan, and other related costs.

Discussion: The study will provide an important advance in the treatment of EDs; in the long term, it is expected to improve the quality of patient care and the treatment efficacy and to reduce waiting lists as well as direct and indirect costs associated with the treatment of EDs in Spain.

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Background
In the last three decades, Eating Disorders (EDs) have become a particularly relevant public health concern due to their growing incidence, the severity of the associated physical and psychiatric comorbidities [1], the high suicide and mortality rates [2] and the resistance of patients to treatment [3], among other reasons. In addition, they constitute today the third most prevalent chronic illness among the adolescent female population in developed and westernized societies and a public health priority around the world [4, 5]. Specifically in Spain, 75,000 adolescents are affected by one of the different manifestations of the disorder, among which 90% are female [6].

Although we do not have specific data related to the economic burden of treatment in Spain, various Cost Of Illness (COI) studies conducted in other European countries indicate that the direct costs for diagnosis and treatment of patients (medical expenditures, nonmedical costs, out-of-pocket expenses of the patient and his/her family), together with the indirect costs (loss of productivity due to sickness, reduced productivity or death) imply a high economic impact [7] and a significant loss of quality of life for people affected [8, 9] and their families [10]. However, according to the systematic review conducted by Stuhldreher et al. (2012), COI studies are sparse, the costs are often underestimated, the majority target only outpatient settings, and due to their heterogeneous nature, it is impossible to carry out comparisons between studies.

The widespread use of new technologies offers a promising and innovative way to improve the quality of care for patients with an ED and to reduce its economic burden. Systematic reviews of the literature demonstrate the effectiveness of eHealth as a complementary therapeutic tool in the treatment of EDs [11–16]. For example, Internet-based interventions have proved their efficacy in reducing ED-related symptoms in a number of Randomised Controlled Trials (RCTs) [17–19]. However, to date, there is a complete absence of RCTs evaluating the effectiveness of mHealth interventions for EDs and existing studies only offer preliminary results about acceptability and usability of such tools [20–22].

In particular, mHealth interventions for EDs may help patients to increase their treatment adherence and to better deal with feelings of stigma associated with face-to-face psychological treatment. Additionally, considering the shortage of ED specialists and the long waiting lists in mental health outpatient settings, mHealth interventions may offer improved accessibility and availability of treatment, at lower costs [20].

In addition, it is worth noting that traditional Cognitive Behavioural Therapy (CBT) is considered the “gold standard” treatment for EDs. CBT uses real-time self-monitoring of patients’ eating habits and behaviours to help them gain a clearer picture of their problem and understand what triggers their behaviour and what are the consequences of it. Self-monitoring is the key driver of behavioural change as it helps patients intervene in the moment [12]. By doing so, all the relevant information regarding the eating problem is entered in real time and can be recovered repeatedly when the therapists or the patients want. Although the efficacy of online CBT interventions as alternatives to face-to-face treatment for patients with EDs is rather controversial [13, 23], such interventions may have a lot of advantages when used as therapeutic tools. Within this context, smartphone apps can offer an attractive and personalised treatment option and can reach a wider range of patients with low motivation for change or others who desire anonymity. Self-monitoring could be enhanced through smartphone-based interventions, as data entry is simpler than traditional pen-and-paper methods. In addition, reminders and motivational messages as well as bidirectional interactions between patient and therapist can improve adherence to treatment [24].

Nevertheless, while growth in mHealth interventions has been rapid, advances in the existing evaluation frameworks have not been seen [25–27]. As a result, a huge number of health apps are widely available to the general healthcare public without possessing best-practice guidelines and certifications or a standardised validation process to assess their long-term cost benefits [28].

The TCApp1 represents a tool for connecting patients and therapists in the time between medical consultations. It is currently available on Google Play and Apple Store markets, there are more than 412 patients who are currently using it and has been developed in collaboration with different public and private mental health institutions in the Barcelona area (Althaia, Hospital de Sant Rafael, CST, ITA and Hospital Sant Joan de Déu). The TCApp was designed from the very beginning with therapists’ and patients’ needs and interests in mind. By using the TCApp, patients and therapists are in continuous contact, allowing for a quicker reaction from the therapist according to the patients’ needs.
Through the TCApp, patients can record their thoughts, actions, emotions and whatever the therapists consider relevant for the therapy, since the app can be customized according to the therapy requirements of each specific patient. It involves algorithms based on artificial intelligence that can generate alarms when strategic words (i.e., suicide, death, etc.) are written. It also introduces technologies to allow real-time online contact with therapists and gamification aesthetics with prizes, rewards and reminders to improve patients’ engagement.

The back-office tool for therapists is a web-based platform where therapists can see in real time what their patients have registered (i.e., generation of graphs in a period of time to see parameter comparison and patient evolution) and they can interact in real time with them, using PUSH notifications. The tool is integrated in Azure server in order to ensure accordance with the most restrictive data protection laws and it is prepared for the integration in local management systems of hospitals and clinics. Lastly, there is currently no application available to provide the same services and benefits as the TCApp as most of the available applications contain self-help functionalities, rather than allowing for a bidirectional link between the patient and the therapist.

The purpose of this trial is to conduct a multicentre, randomized controlled trial with 250 patients diagnosed with an ED. In this experiment, the patients from the experimental group will test a mHealth application (TCApp developed by HealthApp) and then, a clinical efficacy analysis, nursing. The total sample will be approximately 250 patients with a diagnosis of EDs who are currently under treatment, from different public and private mental health services in Spain (Parc Taulí Hospital, Sant Rafael Hospital, Servei de Salut de les Illes Balears, Sant Joan de Déu Hospital, Niño Jesús University Children’s Hospital, San Carlos Clinic Hospital, Dexeus University Hospital of the Quirónsalud group and Eating Disorders Institute ITA).

Methods
Design
We will follow a mixed-methods approach, combining qualitative and quantitative methods, through a multicentre RCT with two parallel groups (an intensive intervention group with TAU and TCApp and a TAU control condition) with a 1:1 allocation.

Sample
The total sample will be approximately 250 patients with a diagnosis of EDs who are currently under treatment, from different public and private mental health services in Spain (Parc Taulí Hospital, Sant Rafael Hospital, Servei de Salut de les Illes Balears, Sant Joan de Déu Hospital, Niño Jesús University Children’s Hospital, San Carlos Clinic Hospital, Dexeus University Hospital of the Quirónsalud group and Eating Disorders Institute ITA).

Inclusion criteria

- Diagnosis of an Eating or Feeding Disorder, based on: a) the Schedule for Affective Disorders and Schizophrenia-Present and Lifetime version (K-SADS-S-PL) (DSM-V criteria) [29] for minor patients or b) the Structural Clinical Interview for DSM-IV (SCID-1) [30] for adult patients. The diagnosis should be one of the following types: Anorexia Nervosa (AN); Bulimia nervosa (BN); Binge Eating Disorder (BED); Other Specified Feeding or Eating Disorder: Atypical Anorexia Nervosa, Bulimia nervosa (of low frequency and/or limited duration), Binge-eating disorder (of low frequency and/or limited duration), purging disorder, night eating syndrome.
• Treatment regimen: Day Hospital or Outpatient treatment, regardless of the illness duration or the severity of the disorder
• Treatment received by ED unit of reference: Standard Cognitive Behavioural Therapy
• Understanding of Spanish, Catalan or English language, depending on the language option chosen by the participant for the TCApp
• Minimal digital skills and availability of proper mobile phone for patients

Exclusion criteria

• Age less than 12 years
• Treatment regimen: Inpatient treatment
• Diagnosis of psychosis
• Intellectual disability
• Have a mobile phone with a Windows Phone operating system

Procedure and randomization

First, all material with information related to the study (research protocol, informed consent, patient information sheet, Data Collection Logbook, safety- and privacy-related issues concerning the TCApp) have been submitted for approval to each one of the Ethical Committees of the participating hospitals. It should be mentioned that the approval of the Ethical Committee of the University leading the study (Open University of Catalonia, UOC) was obtained on February 21st, 2017.

Participants will be recruited after previous recommendation by one of the ED specialists working at each centre. Specialists will do a preliminary screening following the inclusion and exclusion criteria in order to identify potential candidates for the study. Interested individuals will be able to confirm their participation by notifying the ED specialist who will be responsible for their treatment. Then, an information letter and an informed consent form will be delivered to them.

After completing and signing these documents (for underage patients, their parents will be required to sign the informed consent and for patients aged more than 18 years, they will have to sign informed consent themselves), initial clinical interviews will be conducted by psychologists or other collaborators working in the ED unit. All interviewers will be previously trained in administering the K-SADS-PL or SCID interview, depending on the participant’s age. The objective of these interviews is: a) to determine whether participants are definitely eligible for the study following the inclusion criteria, b) to establish the diagnosis for each patient and c) to evaluate them for possible comorbidities. At this time, sociodemographic and clinical data of each patient will also be collected through a brief interview. Then, those who meet the inclusion criteria will be invited to complete the baseline questionnaires for the study. During this baseline evaluation (T0), questionnaires will be administered to patients, their informal caregivers and the ED specialist responsible for the online monitoring of each patient. In addition, telephone interviews will be conducted with the technical staff and the ED specialists.

After completion of the baseline questionnaires, participants will be randomized to one of the two study conditions (experimental or control group). Randomization will be carried out by an independent researcher in blocks of 10 participants within each ED unit (50% of patients from each block will be assigned to the experimental group and the other 50% to the control group), using a random allocation program. Allocation concealment will be also ensured.

After this, patients will be notified about the group they belong to during their next visit to the ED unit. At this time, patients from the experimental group will be given oral and written instructions about how to download and use the TCApp. ED specialists responsible for the online monitoring will be also taught the basic principles for using the application. In turn, patients from the TAU control group will be told that access to the TCApp will be offered to them after a waiting period of 6 months (T2 evaluation completed).

Then, each group of patients will receive the treatment that corresponds to them during a period of 12 weeks. At the end of the 12-week treatment, patients from the experimental group will stop using the TCApp and the evaluation post-treatment (T1, 12 weeks later) will be carried out, including: a) a brief clinical interview (patients), b) questionnaires (patients, informal caregivers, ED specialists); c) telephone interviews (technical staff, ED specialists) and d) focus groups with ED specialists of each institution who are interested in participating as well as with patients of the experimental group.

In the follow-up evaluation T2, 6 months after the beginning of treatment, patients, their informal caregivers and ED specialists responsible for the online monitoring will complete some questionnaires. In addition, telephone interviews will be conducted with ED specialists and a brief clinical interview will be carried out with patients. At the end of this phase, patients from the control group will be given access to the TCApp after being contacted by the ED specialist who will be responsible for their online monitoring.

Figure 1 presents the procedure and the timeline of the study and Table 1 provides a detailed description of the methodology with a definition of the study variables and their assessment tools.

Study conditions

The experimental group will receive the standard treatment based on CBT principles that is offered by the different ED
units in Spain, plus an online intervention using the TCApp for a period of 12 weeks. Only one ED specialist will be responsible for the online monitoring of each patient. For the specific purposes of our study, this role has been assigned to the nursing staff for most of the centres.

The TCApp application proposes patients a series of different functionalities, including daily self-records of their thoughts, emotions and actions, a chat with their therapists and motivational exercises. An online platform of the TCApp (BackOffice) is also available for therapists for the online monitoring of each patient. There, therapists have the possibility to follow the patient’s daily self-records, generate personalized reports and graphs and communicate with him/her via chat, based on the information that the patient has provided online.

During these 12 weeks, the patient should use the TCApp at least once a day, completing at least one self-record per day and/or contacting his/her therapist via chat when needed. The therapist responsible for the online monitoring should connect to the online platform and perform the following actions at least once a week: follow the patient’s daily self-records, generate personalized reports or graphs and communicate with him/her via chat. After a 12-week period, patients from the experimental group and their therapists will stop using the TCApp (they will be discharged).

The TAU control group will receive the standard face-to-face CBT, offered by the different ED units in Spain. Patients from the control group will be offered access to the TCApp after a 6-month period.

**Measurements**

**Primary outcome measures (for patients)**

*Eating Disorder Examination Questionnaire* (EDE-Q) [31, 32]. The EDE-Q is derived from the Eating Disorder Examination Interview (EDE) (Fairburn & Cooper, 1993) and is commonly used to assess the frequency of core ED behaviors and the attitudinal features related to ED pathology over the past 28 days. Twenty-two of the 36 items assess attitudes related to ED pathology and are divided into four subscales (concerns about weight, shape and eating, and restraint). Items are rated on a 7-point Likert scale ranging from 0 = not 1 day/not at all, to 6 = every day/markedly. The core ED behaviors (remaining 14 items) are assessed in terms of their presence and frequency. A global score of eating psychopathology can be calculated by summing and averaging all of the items. Higher scores indicate higher ED psychopathology. The EDE-Q has demonstrated good reliability and validity both in its original version [33] and in Spanish [32].

*The Short Evaluation of Eating Disorders* (SEED) [8, 34]. The SEED was developed for the assessment of
key ED symptoms. It includes 3 items related to AN symptoms (degree of underweight, fear of weight gain and distortion of body perception) and 3 items related to BN symptoms (amount of binge eating episodes, amount of compensatory behaviors and over concern with body shape and weight). The questionnaire allows the calculation of two severity indices for AN and BN (range from 0 to 3). The construct validity and the criterion-related validity of the SEED yielded positive results and the sensitivity to change of the instrument was also satisfactory [34].

**Secondary outcome measures**

**For patients Beck Depression Inventory (BDI-II) [35].** The BDI-II is an instrument used for assessing the severity of depression in adult and adolescent patients over a period of 2 weeks. It includes 21 items, which are answered on a 4-point Likert scale ranging from 0 (not at all) to 3 (extreme form of each symptom). The total score is obtained by summing the severity ratings of each depression item. The questionnaire showed strong internal consistency and good test-retest reliability in both English and Spanish.

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**Table 1** Instruments for measuring clinical efficacy and cost-effectiveness

| Dimension                        | Methodology                | Assessment measures                                                                 | Stakeholder              | T0 | T1   | T2 |
|----------------------------------|----------------------------|-------------------------------------------------------------------------------------|--------------------------|----|------|----|
| Clinical efficacy                | Clinical interview         | Minors (6-18 years old): Kiddie Schedule for Affective Disorders and Schizophrenia-Present and Lifetime version (K-SADS-PL) (DSM-5) Adults: Structured Clinical Interview for DSM Disorders (SCID-I) | Patient                  |    |      |    |
| Physical and clinical variables  | Brief clinical interview   | Comorbidities, medication, illness duration, Body Mass Index, among others          | Patient                  |    | x    | x  |
| Eating Disorder psychopathology  | Questionnaire              | - Eating Disorders Examination Questionnaire (EDE-Q)                                 | Patient                  |    | x    | x  |
|                                  |                            | - Short Evaluation of Eating Disorders (SEED)                                        |                          |    |      |    |
| Depression                       | Questionnaire              | Beck Depression Inventory-II (BDI-II)                                               | Patient                  |    |      |    |
| Anxiety                          | Questionnaire              | State-Trait Anxiety Inventory (STAI)                                                 | Patient                  |    |      |    |
| Motivation to change             | Questionnaire              | Patients with Anorexia Nervosa: Anorexia Nervosa Stages of Change Questionnaire (ANSOCCQ) Patients with Bulimia Nervosa or Binge Eating Disorder: Bulimia Nervosa Stages of Change Questionnaire (BNOSOCQ) | Patient                  |    |      |    |
| Dropout                          | Telephone interview        | Reasons for dropout                                                                   | Clinician responsible for the online monitoring |    |    | x  |
| Risk of suicide                  | Questionnaire              | Beck Hopelessness Scale (BHS) / Suicide Intent Scale (SIS)                           | Patient / Clinician responsible for the online monitoring |    |    | x  |
| Quality of life                  | Questionnaire              | 12-14 years old: EQ-SD-Y (children version) > 14 years old: EQ-SD-5 L                | Patient                  |    |      |    |
| Caregiving experience            | Questionnaire              | Experience of Caregiving Inventory (ECI)                                              | Family caregiver         |    |      | x  |
| Cost-utility and cost-effectiveness analysis | | | | |
| Cost related to the development and maintenance of the online platform | Telephone interview | Staff involved, number of extra hours x pay rate/hour | Technical staff |    |    | x  |
| Costs related to healthcare utilization, medication and school and / or work absenteeism | Questionnaire | iMTA questionnaire for Costs associated with Psychiatric Illness (TIC-P) / iMTA questionnaire for Costs associated with Psychiatric Illness, Parent-Form (TIC-P Children) | Adult patient / family caregiver (referring to minor patient) |    |    | x  |
| Patient’s healthcare utilization | Telephone interview | Number of visits to ED specialists, number of visits to the emergency department, medication consumption and its costs | Clinician responsible for the online monitoring |    |    | x  |
| Cost of implementing the online intervention | Telephone interview | Clinicians involved, number of extra hours for online monitoring x pay rate/hour | Clinician responsible for the online monitoring |    |    | x  |
| Usability and satisfaction       | Questionnaire              | - Client Satisfaction Questionnaire (CSQ-8) - System Usability Scale (SUS)           | ED specialists and patients from the experimental group |    |    | x  |
languages, among clinical and non-clinical samples [36, 37].

State-Trait Anxiety Inventory (STAI) [38]. The STAI is composed of 20 items for assessing trait anxiety and 20 items for state anxiety. Responses are on a 4-point Likert scale, ranging from “Almost never” to “Almost always”. Higher scores indicate higher levels of anxiety. The instrument showed satisfactory reliability with Cronbach’s α ranging from 0.86 to 0.95 and also test-retest reliability coefficients ranged from 0.65 to 0.75 over a 2-month interval [38]. The Spanish validation by Spielberger, Gorshuch and Lushene [39] showed satisfactory reliability for State Anxiety (0.90-0.93) and for Trait Anxiety (0.84-0.87). Test-retest reliability varied 0.73 and 0.86 on STAI-T [40].

Anorexia Nervosa Stages of Change Questionnaire (ANSOCQ) [41] and the Bulimia Nervosa Stages of Change Questionnaire [42] will be used to measure patients’ readiness to recover behaviours and attitudes related to their eating disorder. In particular, the ANSOCQ is used for AN patients and the BNSOCQ for patients with BN or Binge Eating Disorder. Both questionnaires are composed of 20 items and scores on each item range from 1 to 5, following stages of change model by Prochaska and colleagues (1998) [43], according to which “1” signifies the Precontemplation stage, “2” is for Contemplation, “3” is for Preparation, “4” is for Action and “5” is for Maintenance. Total scores are obtained by summing the individual items and range from 20 to 100. The ANSOCQ showed good internal consistency and one-week test-retest reliability, both in the English and Spanish version [44]. In turn, the validation study of the BNSOCQ, which was conducted in Spain, demonstrated good internal consistency with Cronbach’s alpha of 0.94 and one-week test-retest reliability of 0.93 [42].

Beck Hopelessness Scale (BHS) [45]. The BHS is a 20-item questionnaire designed to measure negative attitudes about the future in clinical and research settings. More specifically, it measures three major aspects of hopelessness: feelings about the future, loss of motivation and expectations. The Spanish validation of the instrument showed adequate reliability (Cronbach’s α ranging from 0.82 to 0.84) and moderate construct validity [46].

EQ-5D-5 L and EQ-5D-Y [47]. The EuroQoL-EQ-5 L is a standardized instrument used as a health outcome measure for economic evaluation studies. It consists of five dimensions: mobility, self-care, usual activities, pain/discomfort and anxiety/depression. Each dimension has 5 levels: no problems, slight problems, moderate problems, severe problems and extreme problems. The questionnaire also includes an EQ Visual Analogue scale (EQ VAS) about how good or bad the individual’s health is today. The EQ-5D-5 L has been validated in 6 countries, including different groups of patients with chronic conditions and a student cohort. The psychometric properties of EQ-5D-5 L were superior to the previous version of the instrument (EQ-5D-3 L) in terms of validity, although reliability remains to be assessed for the EQ-5D-5 L. In addition, the child-friendly EQ-5D version (EQ-5D-Y) [48] will be used for children and adolescents following and same structure as the EQ-5D (descriptive system and EQ VAS).

iMTA questionnaire for Costs associated with Psychiatric Illness (TiC-P) [49]. The TiC-P will be used for adult patients only, to evaluate the healthcare utilization due to the psychiatric illness, such as number of patient visits to their general practitioner, medical specialists and paramedics, during the last 3 months. The questionnaire will also assess productivity losses due to absence from work or due to reduced efficiency during paid or unpaid work, as well as medication consumed due to the patient’s psychiatric disorder. The instrument establishes direct medical costs and productivity costs and is widely used among different countries for economic evaluations in mental health. The Spanish version of the questionnaire has been obtained upon permission from the original authors. The feasibility and reliability of the instrument was satisfactory and the construct validity of the questions related to healthcare utilisation and long-term work absenteeism was also good [50].

The System Usability Scale (SUS) [51, 52]. The SUS is a 10-item questionnaire with five response options that range from 4 = “Strongly agree” to 0 = “Strongly disagree”. The SUS provides a “quick and dirty” reliable tool for measuring the usability of a range of systems. The participant’s scores for each question are converted to a new number, added together and then multiplied by 2.5 to convert the original scores of 0-40 to 0-100.

Client Satisfaction Scale (CSQ-8) [53]. The CSQ-8 is an 8-item questionnaire that is designed to measure client satisfaction with services. Scores are on a 4-point Likert scale and range from 8 to 32, with higher scores indicating greater satisfaction. The instrument showed satisfactory reliability with Cronbach’s α ranging from 0.83 to 0.93 and correlations of the instrument with other measures of general satisfaction ranged from 0.6 to 0.8. The CSQ-8 is also available in Spanish by Martínez and Beiti [54].

For clinicians responsible for the online monitoring Suicide Intent Scale (SIS) [55]. The SIS is a questionnaire that measures the severity of the suicide attempt and should be fulfilled by the clinician responsible for the treatment of the patient. The scale consists of 20 questions, which are scaled from 0 to 2 and cover three aspects: 1) objective circumstances surrounding the attempt, 2) perceptions of potential lethality, expectations of rescue, purpose of the attempt, impulsivity and
reaction to the attempt and 3) other aspects. Both the English and the Spanish version of the instrument showed satisfactory reliability and validity [55, 56]. The System Usability Scale (SUS) [51, 52]. Client Satisfaction Scale (CSQ-8) [53].

For caregivers Experience of Caregiving Inventory (ECI) [57]. The ECI is composed of 66-items and is designed to measure how a person caring for someone with a serious mental illness appraises his/her experience. It is composed of ten subscales, eight of which measure negative aspects of caregiving (difficult behaviors, negative symptoms, stigma, problems with services, effects on family, need to backup, dependency and loss) and two measure positive aspects (rewarding personal experiences and good aspects of relationship with the patient), which can be grouped into two subscales: the ECI-negative and the ECI-positive. Responses are on a 5-point Likert scale (ranging from 0 = “never” to 4 = “nearly always”). Higher scores indicate a more positive or negative appraisal of the caregiving experience. The ten scales showed satisfactory reliability with Cronbach’s α ranging from 0.74 to 0.91. The internal reliability coefficients of the caregivers were 0.81 for the ECI-positive and 0.92 for the ECI-negative.

EQ-5D-5 L [48].
iMTA questionnaire for Costs associated with Psychiatric Illness, Parent-Form (TiC-P Children). The Dutch version of the TiC-P Children has been used to measure medical and non-medical costs in children with mental health disorders. The questionnaire was obtained upon permission from the original authors and was translated from Dutch into Spanish by our research group.

Qualitative assessment
In order to perform the economic evaluation, all relevant costs due to healthcare utilization (number of visits to ED specialists, number of visits to the emergency department, medication consumption and its costs) as well as other costs related to the implementation of the online intervention with the TCApp for both clinicians responsible for the online treatment and technicians (amount of extra working hours) will be evaluated through a telephone interview with each stakeholder.

Sample size calculation
The a priori sample size calculation was based on results from previous studies that implemented Internet-based programs in the treatment of EDs [58–60]. A small between-group effect size (Cohen’s d = 0.40) is expected. The calculation was conducted by the software program G*POWER. The primary analysis will concern the hypothesis that the average level of eating pathology at post-intervention in the control group, based on the EDE-Q scores, will be higher than the average levels of eating pathology in the experimental group. Assuming an alpha of 0.05 and a power of 0.80 (β – 1) in an independent samples one-way t-test study, a minimum of 100 participants would be required per study arm. Allowing for a dropout rate of 25% of study participants from baseline, a total of 250 participants need to be recruited.

Data management
All data related to the trial, including clinician’s paper notes of the diagnostic interviews (K-SADS, SCID), self-report questionnaires, informed consent forms, and audio recordings of the telephone interviews, will be securely stored in the workplace of our research group located in the UOC. Only authorised researchers directly involved in the study will have access to this information. According to the UOC’s approval report, all obtained data will be kept for a minimum of 5 years. After obtaining the signed informed consent from participants, a unique code will be allocated to each one of them. The file that links participants to their codes and the databases will be also securely stored on a secure server, password-accessible only to the research team.

Statistical analysis
Both intent-to-treat and completers analyses will be carried out. Intent-to-treat analysis will include every participant who will be randomly allocated to one of the study conditions, that is to say, 250 patients. Whenever possible, we will try to collect follow-up data from participants who have dropped out, in order to keep our dataset as complete as possible. Baseline differences between completers and dropouts will be analysed (Pearson’s chi-square test Student’s t test) using data from the clinical interviews and the baseline questionnaires and possible reasons for dropout will be examined through interviews with the ED specialists (T1, T2).

A participant will be considered a completer if he/she has completed the initial clinical interview as well as T0 and T1 evaluations. For participants from experimental group, to be considered completers they will have to have used the TCApp at least 70% of the time that was initially agreed upon before the start of the experiment (at least once a day during a period of 12 weeks). Only data from completers will be used to determine the treatment effect on the main outcome variable.

All analyses will be carried out using SPSS (Statistical Package for the Social Sciences). First, we will examine the data using descriptive statistics. Between-group analysis will be conducted using Pearson’s chi-square test or Fisher’s exact test for categorical data and Student’s t test or the Mann-Whitney U test for continuous variables, depending on the normality of the distribution. The
normality of the distribution of the variable will be assessed using the Kolmogorov-Smirnov test. The effect of the intervention in terms of reduction in the primary and secondary outcome measures over time will be computed by using a mixed model linear regression analysis with time, experimental condition and their interaction as independent variables. Since there may be some heterogeneity in the implementation of the intensive online intervention across the different participating institutions, between-centre heterogeneity will be explored in subgroup analyses.

Economic evaluation
A cost-effectiveness analysis of the TCApp will be performed, which will compare the effects and costs of the intensive online intervention (TAU + TCApp) with the ones for TAU. The effects will be measured with quality-adjusted life years (QALYs) obtained from the EQ-5D-5 L and EQ-5D-Y questionnaires. Furthermore, the effects regarding the primary outcome variables (EDE-Q, SEED) will also be taken into account. On the other hand, costs will cover the development of the App, the costs of using the product, and the medical costs (visits and medication) for both groups. Moreover, societal costs such as productivity losses and the caregiving burden will also be assessed. Information on costs will be obtained from the iMTA questionnaire and from interviews with the clinicians and technicians. This evaluation will determine the incremental costs and effects of the intensive intervention compared to the control condition, and will produce an incremental cost-effectiveness ratio (ICER).

Discussion
In this article we present the protocol of a study aimed to assess the clinical efficacy of an intensive intervention program using the TCApp. In addition to the assessment of changes in ED pathology and other secondary outcomes, such as anxiety, depression and quality of life of both patients and their caregivers, we will assess the differential cost-effectiveness of an intensive treatment (TCApp + TAU) compared to that of TAU. This is an important strength of this study because to our knowledge, there are only a few apps for mental disorders with supporting evidence from RCTs and none of them was specifically designed for people with EDs or ED professionals [11, 61]. Once the TCApp has proven to be an efficient and cost-effective tool for use in ED units in Spain, the long-term contributions of the current study are as follows: 1) to promote the clinical use of the TCApp in ED units not only in Spain but also on an international level, 2) to improve the quality of patient care using the TCApp as a complementary tool alongside face-to-face psychological treatment, while reducing direct and indirect costs associated with the treatment of the illness and 3) to explore the future use of the application in other mental disorders whose treatment is based on CBT, such as depression, addictions and anxiety.

Among the strengths of the study is its large sample size, which is clinically relevant, as it will be recruited from different public and private ED units in Spain. Another strength is the use of face-to-face assessment and a diagnostic interview to establish patient’s clinical diagnosis and confirm his/her final inclusion in the study, according to the inclusion and exclusion criteria. Furthermore, the assessment of the experience of caregivers is another strength worth mentioning.

Regarding the limitations of the study, the short follow-up time as well as the delayed access to the TCApp by the control group, which may influence our results, should be mentioned. Finally, a limitation related to the nature of the study is the fact that it is a multi-centre study that includes several private and public ED units in Spain. To overcome this limitation, when performing the statistical analyses, between-centre heterogeneity will be explored in subgroup analyses.

Endnotes
1http://www.bcnhealthapp.com

Abbreviations
AN: Anorexia Nervosa; ANSOCQ: Anorexia Nervosa Stages of Change Questionnaire; BDI: Beck Depression Inventory; BED: Binge Eating Disorder; BHS: Beck Hopelessness Scale; BN: Bulimia Nervosa; BNSOCQ: Bulimia Nervosa Stages of Change Questionnaire; CBT: Cognitive Behavioural Treatment; COI: Cost Of Illness; CSQ-8: Client Satisfaction Scale; DSM: Diagnostic and Statistical Manual of Mental Disorders; ECI: Experience of Caregiving Inventory; EDE: Eating Disorder Examination Interview; EDE-Q: Eating Disorder Examination Questionnaire; EDs: Eating Disorders; ICER: Incremental Cost-Effectiveness Ratio; K-SADS-S-PL: Schedule for Affective Disorders and Schizophrenia-Present and Lifetime version; QALYS: Quality-Adjusted Life Years; RCT: Randomised Controlled Trial; SCID-1: Structural Clinical Interview for Axis I; SEED: Short Evaluation of Eating Disorders; SIS: Suicide Intent Questionnaire; SPSS: Statistical Package for the Social Sciences; STAT: State-Trait Anxiety Inventory; SUS: System Usability Scale; TAU: Treatment As Usual; UOC: Open University of Catalonia; VAS: Virtual Analogue Scale

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Authors’ contributions
DA and FL conceived the study. DA and FL initiated the study design; ES reviewed it and provided us with insights from a clinical perspective and CF helped with implementation. JA provided us with a list of ED units in Spain willing to participate in the trial. DA and CF provided statistical expertise in clinical trial design. All authors contributed to refinement of the study protocol and approved the final manuscript.

Ethics approval and consent to participate
Ethical approval of the study protocol and the informed consent forms was received from the Open University of Catalonia review board (UOC review board, 21/02/2017) with respect to the scientific content of the protocol and its compliance with applicable research and human subjects regulations. In addition, the trial also received ethical approval from the committees of each participating hospital (Parc Taulí Hospital, Sant Rafael Hospital, Servei de Salut de les Illes Balears, Sant Joan de Déu Hospital, Niño Jesús University Children’s Hospital, San Carlos Clinic Hospital, Dexeus University Hospital of the Quirónsalud group and Eating Disorders Institute ITA).

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Competing interests
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