Baseline Predictors of Attrition in a Randomised Controlled Trial of a New Group Psychological Intervention for People with Recurrent Binge Eating Episodes Associated to Overweight or Obesity

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Short report

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Abstract

Purpose: Understanding the high rate of treatment attrition in trials of people with eating disorders is important as it can compromise the quality of the trials. In clinical practice, it may also contribute to illness chronicity, relapse, and costs. Thus, we investigated factors associated with treatment attrition to a new manualised psychotherapy HAPIFED compared to CBT-E, for individuals with Bulimia Nervosa or Binge Eating Disorder comorbid with overweight or obesity.

Methods: In total, 98 participants were recruited with 50 randomised to HAPIFED and 48 to the control intervention CBT-E, all administered in groups of up to 10 participants. An investigator external to the site conducted the random allocation, which was concealed from the statistician involved in the analysis, and known only to the therapists until the finalization of the 12-month follow-up after the end of active treatment. Three scenarios in the timeline treatment of a total of 30 sessions were assessed: 33% or 60% or 75% of presence. Logistic regression analysis was performed to find the correlates of attrition.

Results: None of the six variables - frequency of binge eating episodes, purging, eating disorder symptom severity, weight, illness duration and mental health-related quality of life - significantly predicted attrition at 33%, but longer illness duration predicted lower treatment attrition at both 60% and 75% presence of the interventions. Also for 75% presence, lower body weight predicted lower treatment attrition.

Conclusions: Lower attrition due to late treatment completion was associated with longer binge eating illness length and a lower body weight. More research is needed to recognize factors that may interfere with engagement in treatments aiming to avoid early dropout.

Trial registration: US National Institutes of Health clinical trial registration number NCT02464345, date of registration 1 June 2015.

Background

A challenge in the care of people with eating disorders is the high rate of attrition of treatments, varying between 29% and 73% in outpatient studies [1]. Fassino et al. [1] argued that such high dropout can compromise the quality of the trials, reducing power and increasing the likelihood of Type I and II errors. In clinical practice, it may also contribute to illness chronicity, relapse, and costs [1]. A recent systematic review, evaluating predictors in following manualised Cognitive Behavioural Therapy (CBT), found poor consistency for the tested predicted variables in samples with Bulimia Nervosa (BN), Binge Eating Disorder (BED) and mixed samples including normal weight eating disorders [2]. Similar results for sociodemographic aspects and eating disorder severity were found in a comprehensive review, but the authors noted that general psychopathology favours the non-sequence of treatment, and that binge-purging subtype of anorexia nervosa, two borderline personality disorder traits (high maturity fear and impulsivity) and two psychological traits (high maturity fear and impulsivity) are predictors for dropouts in eating disorders treatment [1]. In another systematic review and meta-analyses [3], weight suppression, more frequent binge eating and purging behaviours, less motivation for the treatment and avoidant attachment associated with binge/purge subtype were more likely to predict dropout. There are many factors that relate to treatment engagement, factors in the treatment alliance and type of therapy offered [4]. This present paper examines the pre-treatment characteristics of participants, knowledge of which may aid clinicians in tailoring or targeting therapy efforts cognizant that people with these characteristics may be more likely to disengage with the therapy. It is also clear that more studies need to be done to determine risk factors for attrition or treatment outcome in internet-based interventions for BED [5]. Jensen et al. [5] found minor differences between completers and non-completers on depression and no differences in BED-symptoms, BMI, and sociodemographic variables. Participants who completed treatment showed large reductions in eating disorder pathology.

A double-blinded randomised controlled trial (RCT) evaluated the effectiveness of a manualised psychotherapy for people with BN or BED co-morbid with high body mass index (BMI) compared to the Cognitive Behavioural Therapy – Enhanced (CBT-E) [6]. The main aspect of this new intervention named “Healthy Approach to Weight Management and Food in Eating Disorders” (HAPIFED) is the integration of features of CBT-E and Behavioural Weight Loss Therapy (BWLT) with the purpose of promoting weight loss and binge eating remission. To our knowledge, there has been no previous trial of an integrated CBT-E with BWLT for disorders of binge eating in people with high BMI. In this trial, participants were randomly offered 30 sessions of one of the interventions in a group format. A special feature of HAPIFED is a higher number of sessions when compared to CBT-E and other behavioural weight loss programs commonly offered. The reason for extending the treatments in this trial to over six months was to provide sufficient time for implementing the cognitive work as well as the necessary behavioural changes. To avoid bias related to treatment intensity and duration, CBT-E broad version was used which was extended to 30 sessions incorporating additional modules addressing psychological-maintaining factors, namely, clinical perfectionism, core low self-esteem and interpersonal problems. All participants were evaluated in three stages comprising of 22 sessions out of 30 sessions. So it was a meaningful extension of CBT-E.
In the main study for HAPIFED [7], we investigated the efficacy and safety of introducing a weight loss intervention to the treatment of people with disorders of recurrent binge eating and a high body mass index. The results showed that HAPIFED was not superior to CBT-E in promoting clinically significant weight loss and was not significantly different in reducing most ED symptoms. No harm was observed with HAPIFED, in that no worsening of ED symptoms was observed.

The aim of this study was to evaluate features that can predict adherence to psychological treatments (combined or not to multidisciplinary interventions as in HAPIFED) for individuals with BN or BED associated with high BMI. This study investigated the results of the active phase of treatments only (blinded investigation). Three points in the timeline of the 30 sessions were considered for examination being which were 33%, 60% and 75% of treatment sessions completed (the rationale is discussed below in the first sub-section under Methods). We have used logistic regression to determine the correlates of eating disorder treatment completers versus non-completers which is a dichotomous variable.

As exploratory hypotheses, we anticipated that frequency of binge eating episodes, presence of purging behaviour, and eating disorder symptom severity would be associated with drop out. Further, we investigated the following putative predictors of dropout: mental health-related quality of life, illness duration and weight. The term "drop out" is used to describe both the unilateral ending of regular treatment by a patient and the decision for administrative discharge made by a treatment team.

Methods

Definitions

1. For the purpose of this study, we considered three scenarios in the timeline treatment of a total of 30 sessions (1 individual + 29 group sessions). The aim was not to investigate "percentage of sessions attended" but rather the timing of attrition (early/middle/late). So, we present 3 time points in a way that there is a continuum:

   a. 33% of presence, corresponding to 10 sessions with no other requirement.

   b. 60% of presence, corresponding to 18 sessions with no other requirement.

   c. 75% of presence, i.e., (i) completed the treatment until the final session and missed less than 4 sessions in a sequence, and/or (ii) completed at least 22 out of 30 sessions.

The 33% treatment completion is clinically relevant as it is of interest who will engage in treatment. Early dropouts are of concern as they benefit little from the intervention. The 75% treatment completion is also clinically relevant as reflects those who will complete most of treatment, and thus may benefit most from the intervention. So, both 33% and 75% treatment completions have clinical utilities. However, we also chose 60% treatment completion on the ground that there is no consensus on what is treatment completion or number of sessions for treatment to be completed [1, 8]. However late attrition is usually 75% or more of therapy sessions and has been applied in previous studies of the authors [8].

2. Frequency of binge eating episodes was considered when the participant reported at least one episode per week during the last 3 months.

3. Purging was defined as any current use of self-induced vomiting, laxatives, and/or diuretics as a method of weight and/or shape control within the past 3 months. It could be compensatory or non-compensatory.

Participants

We aimed to include 100 participants in the protocol, but had to stop at 98 people, when two participants were randomised but failed to engage in treatment due to time to finish the trial allowed by the research funding agency. Thus, this secondary analysis study comprised a sample of 98 participants, adults, both genders, with BMI ≥ 27 and < 40 kg/m² (either overweight or obese), recruited from clinical and community sources, with threshold or subthreshold BN or BED diagnoses, according to the Diagnostic and Statistical Manual of Mental Disorders, fifth edition (DSM-5) [9]. Sixty-six (67.3%) participants met the DSM-5 diagnostic criteria for BED and 13 (13.3%) for BN. Of the remaining 19 participants, 5 (5.1%) had other specified feeding or eating disorder (OSFED) BED-type, 7 (7.1%) had OSFED BN type and 7 (7.1%) unspecified feeding or eating disorder (UFED). Of those receiving UFED diagnosis, all reported regular recurrent binge eating, but five did not fulfil Criterion B or C for BED, one with recurrent binge eating and self-induced vomiting episodes did not meet Criterion D for BN, and one met all criteria for BN, but only reported subjective binge eating.
The group with threshold or subthreshold BN or BED diagnoses is not a very heterogeneous group as there are some overlaps between the two conditions. We were interested in studying effects of HAPIFED on eating disorders symptoms and weight, so we expected that cases with related features would benefit from treatment. Additionally, the observation that it is common to include subthreshold cases in RCTs has led proposals for revisions of classificatory systems to widen the diagnosis of the main categories [10]. The condition BN includes recurrent binge-eating episodes or the consumption of abnormally large amounts of food in a short period of time which is followed by self-induced vomiting, strict dieting, over-exercising and/or the misuse of laxatives, enemas or diuretics. On the contrary, if one consistently eats large amounts of food, and those eating episodes cause shame, regret, guilt, or sadness, one may have BED.

The exclusion criteria were: use of weight loss medication; clinical conditions that could interfere with appetite regulation; history of bariatric surgery; current diagnosis of psychosis or bipolar disorder; high level of suicide risk; and, current participation in psychotherapy for eating disorders. This RCT was conducted by specialists at a university centre for treatment of eating disorders (PROATA) in the Universidade Federal de São Paulo, Brazil. Participants were recruited from July 2015 to November 2017, via waiting list, advertisements in the internet, printed and oral media.

All participants were evaluated in three stages (see trial protocol for details) [4]. The third stage comprised the assessment with a semi-structured interview that confirmed the eating disorder diagnosis and detailed eating disorders symptoms and behaviour. In total, ten groups were organized, with 5 groups receiving the experimental intervention (HAPIFED) and the other 5 groups receiving the control intervention (CBT-E). An investigator (PH) external to the site conducted the allocation through a website www.sealedenvelope.com. The randomisation process was concealed from the statistician involved in the analysis, and only the therapists of the Brazilian research team knew it until the finalization of the 12-month follow-up after the end of the active treatment. See the flow chart (Figure 1).

Four female therapists were guided in agreement with CBT-E and HAPIFED manuals. They were trained by experienced therapists (PH and Jessica Swinbourne) receiving monthly telephone supervision during the 2 pilot groups, and monthly telephone supervision during the trial by PH who visited Brazil 6 times in 3 years for face-to-face supervisions with the Brazilian therapists and other members of the HAPIFED project. Each pair of therapists conducted both CBT and HAPIFED groups to administrate the non-specific therapists’ effect.

For both interventions were offered an initial individual session and more 29-group sessions, being twice weekly for the first four weeks and weekly after that until the end of active treatment comprising a total of 6 months. HAPIFED is a multidisciplinary program including four sessions with dietician and/or occupational therapist accompanied by the psychological therapists. Differently from CBT-E, HAPIFED emphasizes a nutritional counselling gave by the nutritionist, the behavioural monitoring including appetite cues directed to the hunger and satiety perception, the behavioural activation, e.g. stimulate the remission of body avoidance, healthy exercise encouragement, and emotion regulation skills mainly focusing in mood intolerance. Despite CBT-E originally be offered in 20 sessions, in this protocol the number of sessions were extended in number of sessions and duration to equate to HAPIFED. Besides that, both interventions received four group follow-up sessions during the first 6 months’ follow-up after the end of the active treatment, and a final assessment was conducted in 12-month follow-up.

**Measures**

For the purpose of this study, the following instruments were used:

1. For the evaluation of frequency of binge eating episodes, purging behaviour and eating disorder symptom severity: the semi-structured Eating Disorder Examination Edition 17.0D (EDE) [11] interview was used. The EDE generates eating disorder diagnoses and assesses the symptom severity using four subscales, which are averaged for a global score. The version 16.0 was translated to Brazilian/Portuguese with a satisfactory reliability (80% inter-interviewer agreement and 0.69 Kappa were evaluated with considering the diagnosis using the EDE interview) and concurrent validity (77.3% agreement and 0.68 kappa). For a consistency with the most recent edition - EDE 17.0D – small modifications were made in the previous Portuguese version, in order to derive DSM-5 diagnoses.

2. After a first screening by telephone, the eligible participants were invited for a first presencial interview when the informed written consent was signed, the inclusion and exclusion criteria were rechecked and a medical history and physical examination were conducted by a clinical physician. Weight and height were measured using a calibrated, electronic digital scale and a stadiometer, from which BMI (kg/m²) was calculated.

3. For the evaluation of mental health-related quality of life: the 12-Item Short Form Survey (SF-12) [12] was applied. The SF-12 is a self-report instrument that measures physical and mental health-related quality of life.

4. For the measurement of eating disorder illness duration: the participants fulfilled a self-report questionnaire where they were asked about the illness duration. In the first interview, a self-reported questionnaire was completed by the participants with sociodemographic
information (age, sex, occupation, marital status, etc) and the illness duration where they needed to complete a space with years and months, e.g., Illness duration: ..... years and ..... months.

Participant Flow

As we aimed to study early, middle and late treatment attrition three time points, 33%, 60% and 75%, were analysed. Figure 1 is a flowchart showing how the eligible number (n=98) of participants was arrived at and the number of participants completing at least 10 or 33% sessions, at least 18 or 60% sessions and at least 22 or 75% sessions. This figure shows that the number of participants for these treatment attrition three time points were 71, 51 and 45 respectively.

Statistical analyses

Data were cleaned including correcting for coding errors. Descriptive statistics such as mean for a continuous covariate and its standard error, as well as proportion for a categorical covariate and its standard deviation were estimated for completers and non-completers of sessions. It was more clinically useful to know if sufficient sessions were attended that a therapist would be satisfied the person had ‘completed’ therapy. Fewer sessions than this the therapy may have been less effective because it was incomplete. So the dependent variable identified a decision issue and hence was clearly dichotomous and so treating it as continuous would not be informative to clinicians. Logistic regression analysis was thus performed to determine the significant predictors of treatment completion. The maximum likelihood estimation was not used to fit the logistic models for predicting the odds of completing at least 18 and 22 sessions as they did not satisfy the widely used criterion of having at least 10 events per predictor for using maximum likelihood in logistic regression. For these models the penalised likelihood with Firth's correction [13] was used as it should perform much better in estimating unbiased regression coefficients. The missing data were estimated in the analysis by multiple imputation using multivariate normal imputation. All analyses were performed using SAS version 9.4 [14].

Results

A total sample of 98 participants were included in this study of which 49 were randomly allocated to the treatment group and the remaining 49 to the control group. Out of 98 participants, 66(67.3%) participants met the DSM-5 diagnostic criteria for BED, 5 for other specified feeding or eating disorder (OSFED) BED-type, 5 for regular BE. Of the remaining participants, 13 (13.3%) and 9 had BN (threshold) and recurrent BN (Other specified feeding or ED) respectively.

The majority were women (n = 94, 96%), Caucasian (n = 73, 74.5%), and were employed (n = 59, 60.2%). Forty-five percent were married (n = 44) and 43% (n = 42) completed tertiary education. The mean age was 40.55 years (SE 1.18), mean weight measured at baseline was 89.26 kg (SE 1.27), mean BMI at baseline was 33.68 (SE 0.33), and the baseline weight range was 61.9-114.8.

For the whole sample, the mean values for illness duration, eating disorder symptom severity score, mental health-related quality of life, binge eating frequency and baseline weight were 14.98 (SE 1.15), 2.56 (SE 0.08), 35.59 (SE 1.10), 39.40(SE 3.00), 88.27 kg (SE 1.27) while 18.37% (SE 0.039) performed purging. These results are presented in Table 1.

Table 1. The mean/percentage distribution of the predictors
Six baseline (pre-randomisation) features with a putative association with treatment completion were analysed – frequency of binge eating episodes, presence of purging behaviour, eating disorder symptom severity, mental health-related quality of life, illness duration and baseline weight. No significant statistical differences were found for these variables between participants who completed 33% of the sessions. For those who completed 60% session there was a significant difference for illness duration. No significant differences for frequency of binge eating episodes, purging behaviour, eating disorder symptom severity and mental health-related quality of life were found between participants who had 33%, 60% and 75% or the most stringent of treatment completions, respectively. Participants who achieved the most stringent treatment completion criterion had significantly lower baseline weight ($p = 0.04$) and longer illness duration ($p = 0.03$) than those who did not complete 75% of sessions. (See Table 2).

Table 2. Results of descriptive statistics at baseline and logistic regression for presence of 33% sessions, 60% sessions and 75% sessions, respectively.
<table>
<thead>
<tr>
<th>Baseline feature</th>
<th>33% sessions of presence</th>
<th>60% sessions of presence</th>
<th>75% sessions of presence</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>Mean (SE)</td>
<td>Odds Ratio</td>
</tr>
<tr>
<td>MHRQoL</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Completers</td>
<td>71</td>
<td>36.31 (1.33)</td>
<td>1.02</td>
</tr>
<tr>
<td>Non-completers</td>
<td>27</td>
<td>33.85 (1.97)</td>
<td>0.74</td>
</tr>
<tr>
<td>EDE global score</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Completers</td>
<td>71</td>
<td>2.52 (0.10)</td>
<td>0.80</td>
</tr>
<tr>
<td>Non-completers</td>
<td>27</td>
<td>2.68 (0.13)</td>
<td>0.97</td>
</tr>
<tr>
<td>Weight/kg</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Completers</td>
<td>71</td>
<td>87.98 (1.44)</td>
<td>0.97</td>
</tr>
<tr>
<td>Non-completers</td>
<td>27</td>
<td>92.65 (2.57)</td>
<td>1.02</td>
</tr>
<tr>
<td>Illness duration</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Completers</td>
<td>71</td>
<td>15.86 (1.26)</td>
<td>1.02</td>
</tr>
<tr>
<td>Non-completers</td>
<td>27</td>
<td>13.66 (2.22)</td>
<td>1.01</td>
</tr>
<tr>
<td>Binge eating frequency</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Completers</td>
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<td>41.38 (3.84)</td>
<td>1.01</td>
</tr>
<tr>
<td>Non-completers</td>
<td>27</td>
<td>33.89 (4.07)</td>
<td>0.94</td>
</tr>
<tr>
<td>Purging behaviour</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Completers</td>
<td>71</td>
<td>15.5 (0.04)</td>
<td>0.74</td>
</tr>
<tr>
<td>Non-completers</td>
<td>27</td>
<td>18.5 (0.08)</td>
<td>0.97</td>
</tr>
</tbody>
</table>

*Effect size calculated from odds ratio and regression coefficients are adjusted for confounding. EDE = Eating Disorder Examination; MHRQoL = Mental Health-Related Quality of Life. In the logistic regression models, for purging the reference category is 'not purging' while the remaining

**Discussion**

This study investigated putative predictor variables of treatment completion in a sample of 98 participants of a RCT testing the efficacy of a multidisciplinary intervention (HAPIFED) for people with BN or BED co-morbid with high BMI against a control therapy (CBT-E). Six potential predictor variables measured at baseline were considered in this study - frequency of binge eating episodes, presence of purging...
Conclusion

Purging was included as a predictor because it is one of the most consistent predictors of outcome in the literature. Also, people with BN do not always purge because there are two types, purging BN and non-purging BN. People with non-purging type of BN fast or perform compulsive exercise. It differs from BED by the fact that people with non-purging type BN have regular extreme weight control behaviours while people with BED don’t. People with BED can also purge like those with purging BN but they do it less frequently (less than weekly).

There were also some participants in our study having non-purging type of BN, e.g., in our sample, out of 20 people with BN (13 threshold, 7 subthreshold), 4 were non-purging. In the regression analysis we didn’t focus on differential diagnosis because it is likely to be confounded with binge eating frequency, binge eating duration, binge eating symptom severity, purging etc. which were included as predictors in the models. SF-12 was used because it’s an indicator of mental health related quality of life while the primary aim of HAPIFED was to improve overall health and wellbeing rather than just reduction of body weight. The latter was only a secondary aim. Because of this SF-12 was considered as a predictor of attrition in an RCT to investigate the effect of HAPIFED in comparison to BWLT.

None of the six variables investigated significantly predicted attrition at 10 sessions (33%) of the interventions. For the period of 18 sessions (60%), longer illness duration predicted lower treatment attrition while for 22 sessions (75%) both longer illness duration and lower baseline weight predicted lower treatment attrition. That the majority of predictor variables were not statistically significant in this study is consistent with findings from other studies [1].

Some studies investigated the impact of illness duration and weight on outcomes in treatment, but not on prediction of treatment completion [2,3]. In the present study, attrition increased over the course of therapy, and particularly after 33% of sessions. The duration of the intervention in this RCT was longer than the usually reported – six months of active treatment - what may have affected the increasing dropout rates along the treatment. Another possible explanation for the non-completion of treatments in our study may be related to difficulties in dealing with the cognitive and behavioural changes proposed in therapies within a sample of individuals with recurrent binge eating associated with high BMI.

Nackers et al. [15] highlighted the importance of addressing the commonly observed individuals’ desire to lose a high proportion of weight in a short period of time. We hypothesized that, in our study, those with lower baseline weight were more motivated to accept the increased focus of treatments on the critical examination of unrealistic expectations of achieving an “ideal weight” versus “real weight” and on the perception of the internal cues that regulate appetite regulation (HAPIFED).

It is possible that a longer eating disorder illness or chronicity may be associated with poorer health and less motivation to change and stay in therapy - at least up to 60% of sessions. Furthermore, this effect may not reduce over time as it was found at 75% of sessions that a longer duration was still positively associated with treatment completion.

Strengths of this study included using an RCT having longer duration of the intervention than usual as it allowed examination of attrition and its correlates over three time points. When number of events (attrition) per predictor was inadequate to perform maximum likelihood estimation using logistic regression, the state-of-the art penalised likelihood estimation with Firth’s correction was applied. Limitations of this study included the inability to investigate associations between attrition and substance use or personality disorder, as these were too infrequent in the sample. Other limitations include having only 4 men out of 98 participants, thus limiting generalizability and precluding subgroup analysis. It may be harder to predict earlier attrition compared to later attrition. The heterogeneous sample and the different treatments may have contributed to the difficulty in predicting attrition. Because of using a heterogeneous sample, the severity of ED symptoms was likely to have differed between participants. So we included the severity of ED symptoms as one of the predictors of dropouts for the treatments. We suggest to use a less heterogeneous and larger sample for a future study.
Like any study assessing the effect of an intervention on outcome, high attrition may also impair the quality of studies evaluating interventions for disorders of recurrent binge eating associated with high BMI. However, predictors of treatment completion have not been consistently established. This study found that a longer binge eating illness duration and lower baseline body weight predicted completion of 75% of treatment sessions, and longer binge eating illness duration predicted completion of 60% of sessions of psychological interventions for these disorders. More research is needed to recognize factors that may interfere with engagement in treatments aiming to avoid early dropout, relapses and chronicity.

**Declarations**

**Author's contributions:**

All authors contributed to the study conception and design. Material preparation, data collection were performed by Marly A. Palavras, and analysis were performed by Haider Mannan. The first draft of the manuscript was written by Haider Mannan, Marly A. Palavras, Phillipa Hay, and Angelica Claudino commented on previous versions of the manuscript. All authors read and approved the final manuscript.

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**Compliance with Ethical Standards**

Ethical approval: The Human Research Ethics Committee of the Universidade Federal de São Paulo, Brazil approved the RCT study (CAAE 43874315.4.0000.5505). The trial was formally registered in the U.S. National Institute of Health with Clinical registration number NCT02464345, on 1 June 2015. Access to the data was limited to investigators and authorized researchers only.

Informed consent: All participants agreed and signed the written informed consent.

**Competing interests:**

PH receives/has received sessional fees and lecture fees from the Australian Medical Council, Therapeutic Guidelines publication, and New South Wales Institute of Psychiatry and royalties/honoraria from Hogrefe and Huber, McGraw Hill Education, and Blackwell Scientific Publications, Biomed Central and Plos Medicine and she has received research grants from the NHMRC and ARC. She is Chair of the National Eating Disorders Collaboration Steering Committee in Australia (2019-) and was Member of the ICD-11 Working Group for Eating Disorders (2012-2018) and was Chair Clinical Practice Guidelines Project Working Group (Eating Disorders) of RANZCP (2012-2015). She has prepared a report under contract for Shire Pharmaceuticals in regards to Binge Eating Disorder (BED; July 2017) and Honoraria for training Psychiatrist in BED assessment. HM, MP and AC declare no conflicts.

**Consent for publication**

Not applicable

**Availability of data and materials**

The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

**References**


Figures
Figure 1

Flow chart diagram

Assessed for eligibility
(n = 250)

Excluded (n = 47)
- Not meeting inclusion criteria (n = 26)
- Nonavailable (n = 15)
- Declined to participate (n = 6)
- Other reasons (n = 10)

Randomized
(n = 25)

Completed 10 sessions
(36%)
(n = 7, 28%)

Completed 18 sessions
(60%)
(n = 15, 52%)

Completed 22 sessions (73%)
(n = 15, 48%)

(except for the last session and session < 6 sequential sessions)