A single-arm pilot study of guided self-help treatment based cognitive behavioral therapy for bulimia nervosa in Japanese clinical settings

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Abstract
Objective: Guided self-help treatments based on cognitive behavioral therapy (CBT-GSH) are regarded as a first-line effective treatment for bulimia nervosa (BN). With limited application for CBT-GSH in Japanese clinical settings, we conducted a single arm pilot study in order to confirm the acceptability and availability of CBT-GSH in Japan.

Results: 25 women with BN received 16–20 sessions of face-to-face CBT-GSH. Primary outcomes were the completion rate of intervention and abstinence rates from objective bingeing and purging as assessed by the Eating Disorder Examination. Secondary outcomes were other self-report measurements of the frequency of bingeing and purging, and characteristic psychopathologies of eating disorders. Assessments were conducted before CBT as baseline as well as after CBT. 92% (23/25) of the participants completed the CBT sessions. After CBT-GSH, 40% (10/25) of the participants (intention-to-treat) achieved symptom abstinence. The mean binge and purge episodes during the previous 28 days improved from 21.88 to 10.96 (50% reduction) and from 22.44 to 10.88 (52% reduction), each (before CBT-GSH to after CBT-GSH), and the within-group effect sizes were medium (Cohen’s d = 0.67, 0.65, each). Our study provided a preliminary evidence about the feasibility of CBT-GSH in Japanese clinical settings for the future.

Trial registration This study was registered retrospectively in the national UMIN Clinical Trials Registry on July 10, 2013 (registration ID: UMIN000011120)

Keywords: Eating disorders, Bulimia nervosa, Guided self-help treatments based on cognitive behavioral therapy (CBT-GSH)

Introduction
Bulimia nervosa (BN) is a serious psychiatric illness characterized by repetitive overeating and purging behaviors to prevent weight gain, and is commonly seen in young women. The prevalence rate of BN in Japan is estimated to be between 1.9 and 2.9% [1], and it is reported that the prevalence of BN among Japanese females has been increasing over time [2].

The treatment of BN includes psychotherapy as well as medication based mainly on antidepressants [3]. Especially, cognitive behavioral therapy (CBT), which is focused on BN, has demonstrated high efficacy [4, 5], and self-help approaches that rely on highly structured CBT treatment manuals are considered promising [5]. In fact, previous reviews have shown that CBT-based guided self-help (CBT-GSH) can be effective in treating patients with BN [6–9]. However, in Japan the penetration of both highly structured CBT and CBT-GSH is lagging behind, so patients with eating disorders (EDs) are facing the difficulty of gaining access to the best suited treatments [10].
“Getting Better Bite by Bite (GBBB)” [11] is one of the well confirmed bibliographies for a self-help program that has been evaluated by controlled trial. In previous studies, it was revealed that 7.5–30% of the patients using this manual achieved full remission from BN in both adults [12–15] and adolescents [16] at the end of treatment.

The purpose of this study was to investigate our previous study [17] with a larger number of participants and to report the acceptability and preliminary outcomes of CBT-GSH with the Japanese versions of GBBB for patients with BN in Japanese clinical settings.

Main text
Materials and methods
Study design
This study was a prospective clinical pilot study of a single-group open-label trial. The Research Ethics Committee of Chiba University Graduate School of Medicine approved all procedures. This study was registered in the national UMIN Clinical Trials Registry (ID: UMIN000011120). Written informed consent was obtained from all participants. All procedures were performed in accordance with the Helsinki Declaration. We estimated that a sample size of 18 would allow us to detect significant differences in the frequency of binging or purging before and after CBT-GSH with 80% power (α = 0.05) based on the mean and standard deviation of the frequency of binging or purging reported in previous studies with similar intervention on GBBB [14, 15]. Since 25–29% dropout rates were suggested in their studies [14, 15], we determined our sample size to be 25.

Participants
Inclusion criteria into this study were a primary diagnosis of BN according to DSM-IV, and age 18–39 years. Exclusion criteria were psychosis, mental retardation, pervasive developmental disorders, current high risk of suicide, substance abuse or dependence, personality disorder and unstable medical condition. We recruited 32 patients with bulimia nervosa by clinical referrals from Chiba University Hospital, as well as other local psychiatric hospitals and clinics between October 2012 and November 2015. As 6 participants met the exclusion criteria and one participant declined treatment, 25 patients were eligible for participation in the intervention. Patients were assessed by senior psychiatrists at Chiba University Hospital using the Structured Clinical Interview for DSM-IV Axis I Disorders, Research Version, Patient Edition (SCID-I/P), and all patients met the current DSM-IV criteria for BN.

Measurements
After enrollment, we carried out the following measures in pre-intervention and post-intervention. Current symptoms were assessed by the Eating Disorder Examination Edition 16.0D (EDE 16D) [18], the Eating Disorders Examination Questionnaire (EDE-Q) [19], the Bulimic Investigatory Test, Edinburgh (BITE) [20], the Patient Health Questionnaire (PHQ-9) [21], and the Generalized Anxiety Disorder scale (GAD-7) [22]. In addition, their height and weight were measured to calculate their body-mass index (BMI) before each CBT session.

The primary outcomes were the completion rate of intervention and abstinence rates from objective bingeing and purging as assessed by EDE 16D, a structured clinical interview. Drop-outs were defined as participants who discontinued therapy despite recommendations of the therapists and those who never returned phone calls at the end of the intervention. Remission was confirmed as no longer meeting the criteria for BN diagnosis using SCID-I/P interviews conducted by senior psychiatrists at post-intervention. EDE-Q is a well-known 36-item self-report questionnaire that assesses the frequency of abnormal eating behaviors and eating disorder-related symptom profile. EDE-Q includes four subscale scores—restricting (EDE-Qr), eating concern (EDE-Qe), shape concern (EDE-Qs), and weight concern (EDE-Qw). Bingeing BITE is a 33-item self-report questionnaire designed to assess symptoms of bulimia or binge eating. BITE is composed of two subscales: the symptom scale (BITE-sas) and the severity scale (BITE-ss). The former measures the degree of symptoms present, and the latter provides an index of the severity of binging and purging behaviors as defined by their frequency. PHQ-9 is an easy-to-use screening tool for assessing current depressive symptoms and GAD-7 is a simple assessment tool to measure a participant’s current anxiety.

Interventions
Our CBT-GSH intervention was conducted in 16 weekly 50-min face-to-face sessions, which could be extended to as many as 20 sessions depending on the therapeutic need or participant’s request. Termination of intervention within at least 8 sessions was also permitted in case of symptom abstinence.

As for bibliotherapy, the self-help manual “Getting better bite by bite” [11] was used. The contents of the sessions were tailored for each participant according to their specific difficulties. Structured diagnostic interviews were conducted and self-report questionnaire were completed following the completion of the CBT intervention.

CBT-GSH was conducted by 7 therapists (3 clinical psychologists, 2 psychiatrists, 1 nurse, and 1 psychiatric
social worker) who were experienced in the use of CBT-GSH for EDs and had completed the 2 year CBT training program at Chiba University [17]. All therapists attended the workshop for CBT-GSH for BN presented by Prof. Ulrike Schmidt at Chiba University in 2013. All therapists attended weekly group supervision sessions with other therapists as well as supervision sessions with a senior supervisor. Concomitant treatment with medications of equable dosage was permitted throughout the study.

**Statistical analyses**
Data analyses were carried out with SPSS version 21 (IBM Corp., Armonk, NY, USA). We used within-group t-tests to compare pre- and post-CBT-GSH. Effect size was reported by Cohen’s d. Statistical significance was designated as P < 0.01. For the participants who dropped out, the data gathered at their final visits was used for analysis.

**Results**

**Characteristics of participants and completion rate**
32 patients with BN were enrolled, 26 proved eligible for the study, and 25 agreed to embark upon the treatment. Two participants discontinued therapy while remaining symptomatic (4th session and 5th session, respectively), and thus 23 participants (92%) completed the CBT sessions.

The demographic and clinical characteristics of 25 participants are summarized in Table 1. The participants were all female native Japanese speakers, between age 18 and 36 (mean = 25.60, SD = 5.52 years). All participants met the principal DSM-IV diagnostic criteria for BN. 21 (84%) of the patients practiced self-induced vomiting; eight (32%) also took laxatives to influence their shape and weight. Four patients (16%) neither vomited nor took laxatives but restricted their eating or exercised sufficiently to meet the criteria (purging type, 21; non-purging type, 4). Additional Axis I diagnoses for the patients included major depressive disorder (36%), other mood disorders (e.g., dysthymic disorder, 4%; bipolar II disorder, 12%) and other anxiety disorders (e.g., obsessive–compulsive Disorder, 4%; panic disorder, 4%). Seven patients had been on medication, and 10 patients had a history of anorexia nervosa. Other demographic and clinical variables of the participants are shown in Table 1.

**Treatment outcomes**
The average number of sessions per participant (completer) was 18.83 (SD = 3.38). Table 2 shows the abstinence rates for patients with BN on EDE 16D pre- and post-CBT-GSH. 10 participants (40%) achieved all-symptom abstinence and 13 participants (52%) met the criteria for remission by DSM-IV at the end of treatment.

**Table 1 Demographic and clinical data of the patients with bulimia nervosa**

| Bulimia nervosa (n = 25) | Range          |
|--------------------------|----------------|
| Age (years)              | 25.60 – 36.9    |
| Duration of ED (years)   | 0.3 – 15.4      |
| Education (years)        | 9 – 18          |
| BMI (kg/m²)              | 17.9 – 24.6     |
| Gender (female, n, %)    | 25 (100.0)      |
| Comorbidity (n, %)       |                |
| Major depressive disorder| 9 (36.0)        |
| Dysthymic disorder       | 1 (4.0)         |
| Bipolar II disorder      | 3 (12.0)        |
| Obsessive–compulsive disorder | 1 (4.0)    |
| Panic disorder           | 1 (4.0)         |
| Medication (n, %)        |                |
| Medication-free          | 18 (72.0)       |
| SSRI monotherapy         | 4 (24.0)        |
| SSRI and topiramate      | 1 (4.0)         |
| SSRI and ethyl loflazepate| 1 (4.0)       |
| Topiramate and ethyl loflazepate | 1 (4.0) |
| Past history (n, %)      |                |
| Anorexia nervosa         | 10 (40.0)       |
| Major depressive disorder| 3 (12.0)        |
and purge episodes improved significantly together with a decline in other eating and general psychopathology. Since previous studies using GBBB revealed that 7.5–30% of the patients achieved full remission from BN [12–16], our result is at least comparable to those studies.

The recent questionnaire survey of children and youth in schools revealed high rates of suspected cases of BN and binge eating disorders (BED) among adolescents, but there were few chances to access special treatment services due to the hesitation of the patients as well as the lack of available delivery services in EDs in the community [23]. A way of providing support in the community should be required, as the proportion of teens at the estimated onset age of EDs has increased year by year in Japan [24]. Improving delivery of best evidence-based intervention for patients with EDs is one of the priority issues to be solved in Japan [2]. One of the advantages of self-help interventions is that they can be implemented by a wide range of health care providers [25]. Since the therapists from various professional backgrounds demonstrated the high acceptability and good therapeutic outcomes of CBT-GSH, our study indicated favorable accessibility of the CBT-GSH in Japanese clinical settings.

### Table 2 Abstinence rates from objective bingeing, purging assessed by EDE 16D between pre and post CBT-GSH

|                      | Pre CBT-GSH (n = 25) | Post CBT-GSH (n = 25) |
|----------------------|----------------------|-----------------------|
|                      | N   | %   | N   | %   |
| Objective bingeing   |     |     |     |     |
| Abstinence           | 0   | 0.0 | 10  | 40.0|
| Subclinical          | 1   | 4.0 | 4   | 16.0|
| Clinical             | 24  | 96.0| 9   | 36.0|
| Drop-outs            | –   | –   | 2   | 8.0|
| Vomiting             |     |     |     |     |
| Abstinence           | 4   | 16.0| 12  | 48.0|
| Subclinical          | 1   | 4.0 | 4   | 16.0|
| Clinical             | 20  | 80.0| 7   | 28.0|
| Drop-outs            | –   | –   | 2   | 8.0|
| Bingeing or purging  |     |     |     |     |
| (at least one symptom)|     |     |     |     |
| Abstinence           | 0   | 0.0 | 10  | 40.0|
| Subclinical          | 0   | 0.0 | 3   | 12.0|
| Clinical             | 25  | 100.0| 10  | 40.0|
| Drop-outs            | –   | –   | 2   | 8.0|

CBT-GSH guided self-help treatments based on cognitive behavioral therapy, EDE 16D Eating Disorder Examination Edition 16.0D, “abstinence”, behavior absent during the previous 28 days; “subclinical”, behavior present during previous 28 days less than twice per week; “clinical”, behavior present during previous 28 days two or more times per week.

### Table 3 Changes in the frequencies of bingeing and purging, psychological tests between pre and post CBT-GSH (n = 25)

|                      | Pre CBT-GSH | Post CBT-GSH | P     | Effect size |
|----------------------|-------------|--------------|-------|-------------|
|                      | Mean  | SD   | Mean  | SD   |       | d    | (95% CI) |
| Objective binge daysa (EDE 16D) | 17.56 | 9.93 | 8.48  | 10.73 | <0.001* | 0.88 (0.30, 1.46) |
| Purge daysa (EDE 16D) | 17.24 | 11.31 | 9.44  | 11.81 | 0.001* | 0.67 (0.10, 1.25) |
| Frequency of binge episodesb (EDE-Q item 17) | 21.88 | 15.69 | 10.96 | 17.12 | 0.001* | 0.67 (0.09, 1.24) |
| Frequency of purge episodesb (EDE-Q item 22) | 22.44 | 19.32 | 10.88 | 16.35 | 0.001* | 0.65 (0.08, 1.22) |
| BMI                  | 20.34 | 2.34 | 20.63 | 2.10 | 0.280 | – 0.13 | (−0.68, 0.42) |
| EDE-Qg               | 4.10  | 1.27 | 2.49  | 1.57 | <0.001* | 1.13 (0.53, 1.73) |
| EDE-Qr               | 3.54  | 1.70 | 1.86  | 1.85 | <0.001* | 0.95 (0.36, 1.54) |
| EDE-Qe               | 3.93  | 1.42 | 2.03  | 1.69 | <0.001* | 1.22 (0.61, 1.83) |
| EDE-Qw               | 4.38  | 1.27 | 2.91  | 1.59 | <0.001* | 1.02 (0.43, 1.61) |
| EDE-Qs               | 4.54  | 1.28 | 3.16  | 1.64 | <0.001* | 0.94 (0.35, 1.53) |
| BITEs                | 11.84 | 3.09 | 5.40  | 4.72 | <0.001* | 1.61 (0.97, 2.25) |
| BITEsas              | 23.84 | 4.95 | 14.48 | 8.40 | <0.001* | 1.36 (0.74, 1.98) |
| PHQ-9                | 13.92 | 6.75 | 6.28  | 4.90 | <0.001* | 1.30 (0.68, 1.91) |
| GAD-7                | 8.68  | 5.19 | 4.16  | 3.79 | <0.001* | 0.99 (0.41, 1.58) |

CBT-GSH guided self-help treatments based on cognitive behavioral therapy, SD standard deviations, d Cohen’s d, EDE 16D Eating Disorder Examination Edition 16.0D, EDE-Q Eating Disorder Examination Questionnaire, BMI body mass index, BITE Bulimic Investigatory Test; Edinburgh, PHQ-9, 9-item patient health questionnaire, GAD-7 7-item generalized anxiety disorder scale.

* P < 0.05, within-group t-tests

a Number of symptomatic days during the previous 28 days
b Number of symptomatic episodes during the previous 28 days
c The presented date includes the data gathered from 2 dropped out participants at the final visit
Conclusions
Our study provided a preliminary evidence about the feasibility of CBT-GSH in Japanese clinical settings for the future.

Limitations
The limitations arising from the design of the present study include the small sample size, a single-arm design with no control group, a single-site study, and a lack of follow-up data. Further study will be required using a randomized controlled trial across longer follow-up.

Abbreviations
CBT-GSH: guided self-help treatments based on cognitive behavioral therapy; BN: bulimia nervosa; EDs: eating disorders; CBT: cognitive behavioral therapy; DSM-IV: Diagnostic and Statistical Manual of Mental Disorders, 4th Edition; SCIP-IV: Structured Clinical Interview for DSM-IV Axis I Disorders, Research Version, Patient Edition; GBBB: ‘Getting Better Bite by Bite’; EDE: Eating Disorder Examination Questionnaire; EDE-Q: Eating Disorders Examination Questionnaire global score; EDE-Qr: Eating Disorders Examination Questionnaire restricting score; EDE-Qe: Eating Disorders Examination Questionnaire eating concern score; EDE-Qs: Eating Disorders Examination Questionnaire shape concern score; EDE-Qw: Eating Disorders Examination Questionnaire weight concern score; BITE: Bulimic Investigatory Test, Edinburgh; BITE-sas: Bulimic Investigatory Test, Edinburgh, the symptom scale; BITE-ts: Bulimic Investigatory Test, Edinburgh, severity scale; PHQ-9: 9-item patient health questionnaire; GAD-7: 7-item generalized anxiety disorder scale; BMI: body mass index.

Authors' contributions
RS wrote the draft of this manuscript. KA, MT, NN, RS, HI, RU and TY performed the clinical treatment and management. MN, KA, YH and JM contributed to the conception and design of this study. MN critically reviewed the manuscript and supervised the whole study process. KA and ES critically reviewed the manuscript. MN administered supervision sessions as the senior supervisor, RS and MN performed clinical investigation (diagnosis). MI and ES supervised the overall conduct of the study. Especially, KA and RS contributed equally to this study as the first authors. All authors read and approved the final manuscript.

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Competing interests
The authors declare that they have no competing interests.

Availability of data and materials
The datasets analyzed during the current study are available from the corresponding author on reasonable request.

Consent to publish
Not applicable.

Ethics approval and consent to participate
All procedures were performed in accordance with the Helsinki Declaration. This study was approved by the Research Ethics Committee of Chiba University Graduate School of Medicine (Reference No. 1333). This study was registered in the national UMIN Clinical Trials Registry (ID: UMIN000011220). Written informed consent was obtained from all participants.

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