Non-pharmacological management of antipsychotic-induced weight gain: systematic review and meta-analysis of randomised controlled trials

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Background
Antipsychotic-induced weight gain is a major concern in the treatment of psychosis. The efficacy of non-pharmacological interventions as well as the optimal intervention approach for this side-effect remain unclear.

Aims
To determine the effectiveness of non-pharmacological interventions and specific treatment approaches to control antipsychotic-induced weight gain in patients with first-episode or chronic schizophrenia.

Method
Systematic review and meta-analysis of randomised controlled trials.

Results
Ten trials were included in the meta-analysis. Adjunctive non-pharmacological interventions, either individual or group interventions, or cognitive-behavioural therapy as well as nutritional counselling were effective in reducing or attenuating antipsychotic-induced weight gain compared with treatment as usual, with treatment effects maintained over follow-up.

Conclusions
Non-pharmacological weight-management interventions should be a priority, particularly during the early stages of antipsychotic treatment. Preventive approaches have the potential to be more effective, acceptable, cost-efficient and beneficial.

Declaration of interest
None. Funding detailed in Acknowledgments.

Method

Search strategy
Systematic bibliographic searches were performed to find relevant English and non-English language trials from the following databases: the Cochrane Central Register of Controlled Trials (CENTRAL), Medline, EMBASE, PsychINFO, CINAHL, UMI Proquest Digital Dissertations, Information Science Citation Index Expanded (SCI-EXPANDED), Information Social Sciences Citation Index (SSCI), Information Arts and Humanities Citation Index (A&HCI) and registers of ongoing clinical trials, with each database being searched from inception to May 2007. We additionally searched conference abstracts from ISI Science and Technology proceedings, and ISI Information Social Science and Humanities proceedings. The abstracts, titles and index terms of studies were searched using the following keywords: ‘weight gain’, ‘weight loss’, ‘weight change’ and ‘body weight’ in conjunction with ‘exercise’, ‘psychoeducation’, ‘intervention’, ‘diet’, ‘behavioural therapy’, ‘cognitive therapy’, ‘physical therapy’, ‘group intervention’, ‘management’, and ‘schizophrenia’ or ‘psychosis’. Further papers were found by hand-searching the references of all retrieved articles and previous reviews. We also screened hand-searched copies of the following journals (from January 2000): British Journal of Psychiatry, Journal of Clinical Psychiatry, American Journal of Psychiatry, Schizophrenia Bulletin, Schizophrenia Research and Journal of Clinical Psychopharmacology.

Study selection
Considered for inclusion were RCTs of a specific non-pharmacological adjunctive intervention aimed at preventing or controlling antipsychotic-induced weight gain, with at least 75% of participants diagnosed with schizophrenia-spectrum disorders using either DSM or ICD criteria. Comparison interventions could include either standard care or an active comparator intervention. Participants could be both young adults with recent-onset psychosis and adults with chronic schizophrenia, hospitalised or out-patients, during treatment with first- or second-generation antipsychotics. The primary outcome was considered to be mean change in body weight and body mass index (BMI) by the end of intervention, with secondary outcome measures including mean change in both body weight and BMI by follow-up. Additional
secondary outcome measures comprised mean change on ratings of quality of life, medication adherence and relapse rates.

Two reviewers (M.A.-J. and C.G.-B.) independently assessed the proportion of total variation in study estimates that fitted. Random effects are, in general, more conservative than fixed-effects approaches. If there was evidence of inconsistency across trials, a random-effects meta-analysis was conducted. With decreasing heterogeneity the random-effects approach moves asymptotically towards a fixed-effects model.

Data extraction

Two reviewers (S.H. and M.A.-J.) independently extracted relevant data from included trials, including treatment approach (prevention of weight gain v. weight loss), the nature of the intervention (cognitive–behavioural therapy, CBT v. nutritional counselling (psychoeducation, diet and exercise), treatment format (group v. individual), intervention provider, length of intervention, participants' characteristics, comparison intervention, antipsychotic type and dosage. Additional extracted information included measures of quality of life, medication adherence and relapse rates. Any discrepancies were resolved by consensus. Authors were contacted for the provision of missing data if necessary for the meta-analysis and to determine the eligibility of several studies.

Assessment of methodological quality

Trials were assessed against the following quality criteria: random sequence generation, allocation concealment, masked assessment of outcomes, number of withdrawals, intention-to-treat analysis and manual-based intervention. A maximum credit of five points was given if random allocation and allocation concealment were adequate, outcome was assessed by masked raters, data were assessed according to the intention-to-treat principle and the intervention was manualised.

Statistical analyses

Outcomes were pooled using MetaView, meta-analytic standard software used by the Cochrane Collaboration (RevMan 4.2.9 (PC version), Cochrane Collaboration, Oxford, England). Given that weight and BMI are continuous outcome measures, the weighted mean difference (WMD) was estimated using a fixed-effect meta-analysis with 95% confidence intervals for both end-of-treatment and follow-up time points. We conducted one primary comparison (non-pharmacological interventions v. treatment as usual) and three subgroup comparisons (preventive v. weight loss interventions; individual v. group therapy; CBT v. nutritional counselling). We further examined treatment effects according to sample characteristics (recent-onset psychosis v. chronic schizophrenia). To investigate treatment effects in different subgroups the overlap of the confidence intervals of the summary estimates was considered. In addition, the significant differences between subgroups were explored following the method of Deeks et al.7 This method is based on the chi-squared statistic for heterogeneity. The statistic estimated is compared with a chi-squared distribution to test the significant difference between subgroups.

We assessed heterogeneity of intervention estimates by visually inspecting the overlap of confidence intervals on the forest plots and by the I-squared statistic. The I^2-test of heterogeneity describes the proportion of total variation in study estimates that is due to heterogeneity. If there was evidence of inconsistency of estimates across trials, a random-effects meta-analysis was fitted. Random effects are, in general, more conservative than fixed-effects models because they take heterogeneity among studies into account. With decreasing heterogeneity the random-effects approach moves asymptotically towards a fixed-effects model. Additionally, data from included trials were entered into a funnel graph (trial effect v. trial size) in order to investigate the likelihood of overt publication bias. In the absence of bias, the plot should resemble a symmetrical inverted funnel. If publication bias exists it is expected that, of published studies, the largest ones will report the smallest effects.

Sensitivity analyses were performed to further assess the robustness of the findings to the choice of statistical method (fixed- or random-effects model), the exclusion of the lowest-quality trials (trials with a quality score lower than 1) and the exclusion of the smallest trials (trials with a sample size of less than 40 participants).

Results

Of 28 studies retrieved, 10 were eligible for inclusion. We excluded 5 studies that did not include comparison groups,6–10 6 studies that were non-randomised,17–22 2 RCTs that did not fully describe the sample characteristics and further information could not be obtained,23,24 1 RCT after the authors confirmed that less than 75% of the sample had a diagnosis of schizophrenia-spectrum disorders;25 1 RCT that reported 90% withdrawal rates and did not provide comparison group data;26 1 RCT that only measured eating habits and did not provide body weight or BMI changes;27 and 1 which did not provide data in a usable format and we were unable to obtain further information.28

Six of the included trials investigated cognitive–behavioural intervention strategies,29–34 three nutritional counselling interventions,35–37 and one combined nutritional and exercise interventions.38 Five trials tested group intervention formats,30,31,33,34,36 and five examined individual interventions.29,32,35,37,38 Four studies aimed to prevent antipsychotic-induced weight gain,29,35–37 and six aimed to reduce body weight in those who had already experienced weight increase.30–34,38 Data could be extracted and pooled in meta-analyses from seven of the ten eligible studies. In three studies we were able to pool relevant data with the help of the authors.31,33,37

Interventions lasted between 8 weeks and 6 months with efficacy measures taken at the completion of the trial intervention. Three studies reported follow-up periods ranging from 2 to 3 months after the end of the intervention.31,33,36 With one exception, all trials were carried out in out-patient settings. Only one trial utilised a sample of patients with recent-onset psychosis.29 Trials were conducted in Europe, Asia, the USA and Australia. Study medications included a broad range of first- and second-generation antipsychotics. Other characteristics of the included trials are outlined in the online Table DS1.

Results for all non-pharmacological interventions

Ten trials involving 482 patients compared non-pharmacological interventions with treatment as usual. There was a statistically significant reduction in mean body weight for those in the non-pharmacological intervention groups compared with those on treatment as usual (WMD: 2.56 kg, 95% CI: 2.20 to 2.56 kg, P < 0.001) (Fig. 1). There was no evidence of statistical heterogeneity (I^2=28.9%).

Pooling treatment effects of mean BMI change across all interventions yielded similar significant results in favour of the non-pharmacological interventions (WMD: −0.91 kg/m^2, 95% CI: −1.13 to −0.68 kg/m^2, P < 0.001), with no evidence of statistical heterogeneity (I^2=13.8%).

Follow-up outcomes

Three trials incorporated follow-up measures ranging from 2 months36 to 3 months.31,35 Pooling treatment effects of mean
change in body weight and in BMI demonstrated that the statistically significant advantages of non-pharmacological interventions were maintained at follow-up (WMD = −4.14 kg, 95% CI = −5.80 to −2.49 kg, P < 0.001). Although one trial with high discontinuation rates at follow-up (n = 31; 61%) reported results only for those who completed follow-up assessment, exclusion of this trial resulted in equivalent treatment effects.

**Subgroup analyses**

Prevention of antipsychotic-induced weight gain v. weight loss

Trials were analysed according to whether they aimed to prevent antipsychotic-induced weight gain or whether they were designed to reduce weight in patients who were already overweight or obese (Fig. 1). Although there was evidence of some statistical heterogeneity among trials that intended to reduce weight gain (I² = 51.0% v. I² = 0.0% among those aimed to prevent weight gain) treatment effects were similar. Furthermore, when a random-effects model was fitted there was little change on the subgroup overall estimates (WMD = 2.80 kg, 95% CI = 2.52 to 0.72 kg, P = 0.41). When a random-effects meta-analysis was fitted there was little change on the subgroup overall estimates (WMD = 2.49 kg, 95% CI = 2.37 to 0.72 kg, 95% CI = 7.91 to 4.19 kg, P = 0.18).

The only trial that evaluated an early intervention in young patients with recent-onset psychosis found that weight gain could be significantly attenuated (WMD = −2.80 kg, 95% CI = −4.93 to −0.67 kg, P < 0.01). Similar treatment effects were obtained in trials with participants with chronic schizophrenia (WMD = −2.54 kg, 95% CI = −3.20 to −1.87 kg, P < 0.001).

**Cognitive–behavioural therapy v. nutritional counselling interventions**

Trials were analysed by type of non-pharmacological intervention: CBT v. nutritional counselling (Fig. 3). Although CBT trials appeared to show a smaller effect compared with nutritional counselling intervention trials (WMD = −2.14 kg, 95% CI = −2.98 to −1.30 kg, P < 0.001 v. WMD = −2.37 kg, 95% CI = −3.54 to −1.21 kg, P < 0.001 using a random-effects model), Trials that aimed to prevent weight gain appeared to show a slightly larger effect on mean body weight change than those designed to reduce weight (Fig. 1). However, the confidence intervals of the summary treatment estimates overlapped to an important degree. Subsequently, the approach described by Deeks et al. showed that there was no statistically significant difference between both subgroups (χ² = 1.67, P = 0.20).

**Recent-onset psychosis v. chronic schizophrenia**

Finally, trials were examined according to the characteristics of the sample: recent-onset psychosis v. chronic schizophrenia (Fig. 4). The only trial that evaluated an early intervention in young patients with recent-onset psychosis found that weight gain could be significantly attenuated (WMD = −2.80 kg, 95% CI = −4.93 to −0.67 kg, P < 0.01). Similar treatment effects were obtained in trials with participants with chronic schizophrenia (WMD = −2.54 kg, 95% CI = −3.20 to −1.87 kg, P < 0.001).

**Additional outcome measures**

Only two trials provided data regarding the impact on quality of life of these interventions. Know et al. did not find differences between the groups in terms of quality of life (only a trend towards statistical difference in the physical score changes), but Evans et al. reported significant differences in favour of the treatment group in subjective improvement in quality of life.

Finally, no trials reported data regarding the influence of weight-management interventions on medication adherence.
Assessment of risk of bias

A description of the conduct of the trials included in the meta-analysis and assessment of the risk of bias is presented in the online Table DS2. Few trials gave explicit reports of trial conduct; one described the generation of random sequences, and only one fully disclosed allocation concealment, and a few provided explicit description of who was masked. The attrition rate for the 10 trials varied between 0 and 50% in the control groups, and 0 and 20.7% in the intervention groups. Only two trials appeared to include all randomised patients in their analysis. Four trials were conducted using manual-based interventions.

To determine the influence of study quality on the overall estimates, we performed stratified analysis according to methodological quality. The four low-quality trials (0 points) showed more benefit than the higher-quality trials (WMD=−2.96 kg, 95% CI −3.90 to −2.03 kg). Exclusion of these studies, however, affected the overall effect and the confidence intervals only marginally (WMD=−2.21 kg, 95% CI −3.08 to −1.33 kg).

Publication bias

The funnel plot showed evidence of mild asymmetry (Fig. 5). The smallest studies (fewer than 40 participants included in the analysis) showed slightly larger effects (WMD=−3.00 kg, 95% CI −4.53 to −1.46 kg). However, exclusion of the smallest studies had little effect on the overall estimate WMD=−2.47 kg, 95% CI −3.17 to −1.77 kg).

Discussion

Adjunctive non-pharmacological interventions are effective in reducing or attenuating antipsychotic-induced weight gain when...
compared with treatment as usual in patients with schizophrenia-spectrum disorders. These findings with regard to reduction in mean body weight were confirmed by similar reductions in BMI, which is considered to be a better indicator of obesity and being overweight. Furthermore, treatment effects may be maintained at follow-up.

Effects of intervention modality

Results from this study showed no statistically significant or practically important differences between therapeutic approaches, either individual compared with group interventions, or CBT compared with nutritional counselling. Conversely, there is evidence that suggests that adherence to weight-management programmes is positively correlated with further weight loss.39 The choice of therapeutic approach will depend, then, on those factors that are likely to engage patients in a therapeutic alliance. Thus, the tailored combination of weight-management techniques in a flexible and innovative manner which addresses individual needs and promotes therapeutic alliance is likely to produce best outcomes.

Weight gain induced by antipsychotics and first-episode psychosis

To date, only one RCT has shown the effectiveness of preventive strategies in attenuating antipsychotic-induced weight gain in a young cohort with recent-onset psychosis.29 Although there are few studies, it seems apparent that there is great potential for interventions aimed at early stages, before weight gain takes place. Weight gain is arguably a greater problem for young people experiencing a first episode of psychosis. This group is considered to be especially susceptible to substantial weight gain, which could interfere with the early recovery process. First, younger populations are already less disposed to adhering to medication regimes and potential weight gain may exacerbate non-adherence. Second, the physical changes produced by weight gain may result in social discrimination and stigma as young patients are more sensitive to issues of body image and self-esteem than their older counterparts. Early interventions could prevent or attenuate this medication side-effect as well as the adverse consequences derived from weight gain.

This is consistent with a clinical staging model where treatment effects are thought to be the greatest when delivered as early as possible.43 Two fundamental assumptions underlie this model. First, patients in the earliest stages of schizophrenia have a better response to treatment and a better prognosis than those in later stages. Second, treatments offered in the early stages should have a better response to treatment and a better prognosis than those in later stages. Second, treatments offered in the early stages should be more effective and as well as more effective. Given this background, preventive weight-management interventions have the potential to be more effective, acceptable, cost-efficient and beneficial.

Clinical implications

How clinically meaningful is a weight loss of 2.6 kg? Several authoritative bodies, such as the Institute of Medicine, have...
implied that weight losses of as little as 5% in individuals at risk of metabolic syndromes can result in clinically meaningful reductions in morbidity and risk of early mortality. The majority of individuals with schizophrenia experience clinically significant weight gain, which is associated with greater risk of developing several diseases, including diabetes, hypertension and coronary heart disease. As a result, people with schizophrenia have a 20% shorter life expectancy than the population at large. In this review, the average baseline weight was approximately 80 kg (ranging from 66.5 to 101.3 kg). Therefore, even a weight loss of 1.9–3.2 kg represents a reduction of 2.5–4.0% of initial body weight in a significant number of patients. It may be plausible, then, to expect that these reductions in body weight could result in corresponding reductions in morbidity and early mortality.

**Limitations of the study**

This study has some limitations. First, most of the trials included short-term follow-up periods. As a result we could not draw conclusions on the long-term effectiveness of these interventions. Second, reporting on generation of random sequence, allocation concealment, intention-to-treat analyses and masking was poor, making assessment of the potential for biased estimates of treatment effect difficult. Given the relationship between poor reporting and larger treatment effects, findings reported by these trials may have overestimated summary treatment effects. Third, some of the included trials were limited by the sample size. Therefore, differences between treatment modalities need to be explored in adequately designed RCTs. Furthermore, there was evidence of skew in the data provided by several trials included in the present review. Meta-analytic techniques frequently face the problem of managing non-parametric data. Although there is not a clear consensus regarding the resolution of this statistical issue, we note the limitations of our analysis in accounting for skewed data. Another limitation relates to the generalisability of the findings to clinical practice. Therapists in clinical trials are highly motivated and skilled in the implementation of the intervention being tested, which may affect the generalisability of the results to the population of therapists. As a result, these findings need to be evaluated in pragmatic trials of intervention effectiveness in a range of clinical settings. Finally, as with all systematic reviews, publication bias is a potential source of error. Although there was some evidence of such bias, exclusion of the smallest studies only marginally affected the overall effect.

**Strengths of the study**

Although it is plausible that some studies assessing non-pharmacological interventions to manage antipsychotic-induced weight gain were not discovered by our literature search, our procedures kept this to a minimum. We conducted a thorough search of the electronic literature, including databases that contain unpublished literature, undertook hand-searches and made efforts to access grey literature. Another common problem in meta-analysis is incomplete reporting of consistent outcome data in primary articles. We minimised the impact of such incomplete reporting by contacting authors when feasible. This review includes several trials not included in previous meta-analysis of weight-management interventions, a focus on non-pharmacological approaches with careful evaluation of different treatment strategies and an assessment of trial conduct and potential risk of bias. Although previous systematic reviews have also suggested the effectiveness of healthy living interventions in patients with schizophrenia, they included a limited number of RCTs as well as quasi-experimental studies and did not perform meta-analytic techniques. Furthermore, we found a notable consistency across all study estimates, which was reflected in the robustness of the findings across analytic methods and when the smallest and lowest-quality studies were excluded.

**Implications for future research**

Although the results from this study suggest that non-pharmacological interventions may be effective in reducing antipsychotic-induced weight gain, further research needs to address several salient issues. Given the adverse impact of weight gain on medication adherence and relapse rates, quality of life, social stigma and discrimination as well as self-esteem, interventions to prevent weight gain have the potential to reduce these negative effects. Even though these outcomes were not consistently reported or measured, there is some evidence that nutritional counselling improves quality of life, overall health and body image. Further, CBT may promote client satisfaction and physical well-being. Moreover, we are aware of no data that would allow precise quantification of the impact of weight-management interventions on adherence to medication regimens, subsequent relapse rates and other salient aspects such as perception of social stigma and social isolation. Further research should investigate these issues in order to fully elucidate all the potential benefits of these interventions.

Well-designed trials are required, including further comparison studies of different types of intervention against another. These trials should also address fundamental questions such as the effects of longer interventions and booster sessions, long-term maintenance of outcomes, intervention effects on clinical morbidity and physical health, as well as their cost-effectiveness. In addition, the development and evolution of preventive treatment strategies is critical. Future interventions should be innovative and encourage engagement with therapy by promoting well-being and global recovery.


### Table DS1: Characteristics of randomised controlled trials of non-pharmacological interventions for antipsychotic-induced weight gain

<table>
<thead>
<tr>
<th>Study</th>
<th>Participants/setting</th>
<th>Sample size, n</th>
<th>Age of participants, years: mean (s.d.)</th>
<th>Intervention, length and provider</th>
<th>Comparison group</th>
<th>Follow-up</th>
<th>Antipsychotic medication</th>
<th>Medication dose: mean (s.d.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Álvarez-Jiménez et al</td>
<td>Out-patients with first-episode schizophrenia-spectrum disorders who had not been previously treated with antipsychotic medication</td>
<td>(a) 28 (b) 33</td>
<td>(a) 26.0 (15.5) (b) 27.5 (8.5)</td>
<td>3-month individual early behavioural intervention that incorporated behavioural interventions, nutrition and exercise. Intervention provided by clinical psychologists</td>
<td>Non-structured information on weight gain</td>
<td>12 wk</td>
<td>(1) Olanzapine (2) Risperidone</td>
<td>(3) Haloperidol</td>
</tr>
<tr>
<td>Brar et al</td>
<td>Out-patients or stable long-term in-patients with schizophrenia (n=38) or schizoaffective disorder (n=33) with a BMI &gt; 26 kg/m²</td>
<td>(a) 34 (b) 37</td>
<td>(a) 40.0 (10.1) (b) 40.5 (10.6)</td>
<td>14-week group-based cognitive-behavioural treatment. Treatment providers not reported</td>
<td>Non-structured information on weight gain</td>
<td>14 wk</td>
<td>Risperidone</td>
<td>(a) 4.7 (1.7) (b) 4.2 (1.8)</td>
</tr>
<tr>
<td>Evans et al</td>
<td>Out-patients with schizophrenia (n=16), schizoaffective (n=12), schizoaffectiform (n=10) or bipolar disorder (n=8)</td>
<td>(a) 29 (b) 22</td>
<td>(a) 34.6 (9.6) (b) 33.6 (11.6)</td>
<td>6 individual nutritional counselling sessions over a 3-month period carried out by dieticians</td>
<td>Passive nutritional education by using a booklet</td>
<td>24 wk</td>
<td>Olanzapine</td>
<td>(a) 15.5 (6.9) (b) 14.3 (6.9)</td>
</tr>
<tr>
<td>Khazaal et al</td>
<td>Out-patients with schizophrenia-spectrum disorders (n=49), bipolar disorder (n=5) or other mental disorders (n=7) with a reported weight gain &gt; 2 kg during antipsychotic treatment</td>
<td>(a) 31 (b) 30</td>
<td>(a) 43.0 (9.8) (b) 38.3 (10.4)</td>
<td>12-week group-based treatment sessions that incorporated cognitive-behavioural interventions, moderate physical activity and food intake moderation. Intervention provided by senior psychologists</td>
<td>Informative 2 h group session including nutritional recommendations</td>
<td>24 wk</td>
<td>Atypical and typical antipsychotics</td>
<td>N/R</td>
</tr>
<tr>
<td>Kwon et al</td>
<td>Out-patients with schizophrenia or schizoaffective disorder who had experienced weight gains of more than 7% of body weight</td>
<td>(a) 29 (b) 14</td>
<td>(a) 32.0 (9.2) (b) 29.8 (8.1)</td>
<td>12-week individual weight-management programme that included diet and exercise management based on CBT. Intervention provided by a dietician and an exercise coordinator</td>
<td>Non-structured information on weight gain</td>
<td>12 wk</td>
<td>Olanzapine</td>
<td>(a) 10.4 (5.9) (b) 11.3 (4.8)</td>
</tr>
<tr>
<td>Littrell et al</td>
<td>Out-patients with schizophrenia (n=54) or schizoaffective disorder (n=26)</td>
<td>(a) 35 (b) 35</td>
<td>(a) 33.7 (9.2) (b) 43.5 (10.0)</td>
<td>4-month intervention group that consisted of weekly psychoeducation classes focused on nutrition and exercise. Intervention provided by master’s-level trained clinician</td>
<td>No specific intervention</td>
<td>24 wk</td>
<td>Olanzapine</td>
<td>(a) 16.6 (4.2) (b) 16.3 (4.1)</td>
</tr>
<tr>
<td>McKibbin et al</td>
<td>Out-patients with schizophrenia (n=48) or schizoaffective disorder (n=9) and type-2 diabetes mellitus</td>
<td>(a) 28 (b) 29</td>
<td>(a) 54.8 (8.2) (b) 53.1 (10.4)</td>
<td>24-week group-based educational lifestyle intervention that included basic education about diabetes treatment, nutrition, behavioural change strategies and exercise. Treatment providers not reported</td>
<td>Usual care by the participants’ physicians and three brochures relevant to diabetes management</td>
<td>24 wk</td>
<td>Atypical and typical antipsychotics</td>
<td>N/R</td>
</tr>
<tr>
<td>Scocco et al</td>
<td>Out-patients with schizophrenia (n=15) or schizoaffective disorder (n=5)</td>
<td>(a) 9 (b) 8</td>
<td>(a) 57.1 (12.4) (b) 39.2 (9.9)</td>
<td>8-week individual dietary intervention. Intervention provided by a nutritionist</td>
<td>No specific intervention</td>
<td>8 wk</td>
<td>Olanzapine</td>
<td>N/R</td>
</tr>
<tr>
<td>Weber &amp; Wyne</td>
<td>Out-patients with schizophrenia or schizoaffective disorder with a BMI &gt; 26 kg/m²</td>
<td>N/R</td>
<td>N/R</td>
<td>16-week cognitive-behavioural group intervention including presentations on low-fat diets and plans to increase exercise provided by a psychiatric nurse</td>
<td>No specific intervention</td>
<td>16 wk</td>
<td>Atypical antipsychotics</td>
<td>N/R</td>
</tr>
<tr>
<td>Wu et al</td>
<td>In-patients with schizophrenia with a BMI &gt; 27 kg/m²</td>
<td>(a) 28 (b) 25</td>
<td>(a) 42.2 (7.5) (b) 39.0 (6.7)</td>
<td>6-month diet intervention and regular physical activity implemented by a dietician</td>
<td>No specific intervention</td>
<td>24 wk</td>
<td>Clozapine</td>
<td>N/R</td>
</tr>
</tbody>
</table>

(a), intervention group; (b), control group; BMI, body mass index; N/R, not reported.

a. Weight change is reported at intervention end-point.
<table>
<thead>
<tr>
<th>Study</th>
<th>Randomisation procedure</th>
<th>Allocation concealment</th>
<th>Masking (outcome assessor)</th>
<th>Intention-to-treat analysis</th>
<th>Withdrawals, n (%)</th>
<th>Manual-based treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Álvarez-Jiménez et al²⁹</td>
<td>'Randomly assigned by computer-generated blocks of four random numbers' p.1254</td>
<td>'Randomisation was performed by a member of the team not involved with either the assessments or the treatments' p.1254</td>
<td>'Research assessors and patients intended to be blind to the intervention status' p.1254</td>
<td>Yes</td>
<td>None</td>
<td>'EBI was conducted according to a manual developed by our group' p.1255</td>
</tr>
<tr>
<td>Brar et al³⁰</td>
<td>'Randomly assigned' p.206. No other statement</td>
<td>Not stated</td>
<td>'Rater-blinded' p.205</td>
<td>No</td>
<td>(a) 13 (38.2) (b) 9 (24.3)</td>
<td>No statement</td>
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<tr>
<td>Evans et al³⁵</td>
<td>'Randomly assigned' p.479. No other statement</td>
<td>Not stated</td>
<td>No statement</td>
<td>No</td>
<td>(a) 6* (20.6) (b) 11* (50)</td>
<td>No statement</td>
</tr>
<tr>
<td>Khazaal et al³¹</td>
<td>No statement</td>
<td>Open label</td>
<td>No</td>
<td>No</td>
<td>(a) 6 (19.3) (b) 2 (6.6)</td>
<td>'Handbook for a CBT treatment' p.171</td>
</tr>
<tr>
<td>Kwon et al³²</td>
<td>'Randomised' p.547. No other statement</td>
<td>Not stated</td>
<td>No statement</td>
<td>No</td>
<td>(a) 11 (37.9) (b) 1 (7.1)</td>
<td>No statement</td>
</tr>
<tr>
<td>Littrell et al³⁶</td>
<td>'Randomly assigned' p.238. No other statement</td>
<td>Not stated</td>
<td>'Open-label' p.238</td>
<td>Yes</td>
<td>None</td>
<td>'Using the &quot;solution of wellness&quot; modules' p.239</td>
</tr>
<tr>
<td>McKibbin et al³³</td>
<td>'Randomly assigned' p.38. No other statement</td>
<td>Not stated</td>
<td>No statement</td>
<td>No</td>
<td>(a) 3 (10.7) (b) 3 (10.3)</td>
<td>'Manualised-intervention' p.37</td>
</tr>
<tr>
<td>Scocco et al³⁷</td>
<td>'Randomly assigned' p.117. No other statement</td>
<td>Not stated</td>
<td>No statement</td>
<td>No</td>
<td>(a) 0 (0.0) (b) 2 (6.8)</td>
<td>No statement</td>
</tr>
<tr>
<td>Weber &amp; Wyne³⁴</td>
<td>'Randomised' p.96. No other statement</td>
<td>Not stated</td>
<td>Measurements were completed by one research assistant who was 'blind' p.97</td>
<td>No</td>
<td>(a) 0 (0.0) (b) 2 (28.5)</td>
<td>No statement</td>
</tr>
<tr>
<td>Wu et al³⁸</td>
<td>No statement</td>
<td>Unclear</td>
<td>Unclear</td>
<td>No</td>
<td>(a) 0 (0.0) (b) 3 (12)</td>
<td>No statement</td>
</tr>
</tbody>
</table>

(a), intervention group; (b), control group; CBT, cognitive–behavioural therapy; EBI, early behavioural intervention.

* At intervention end-point.
Non-pharmacological management of antipsychotic-induced weight gain: systematic review and meta-analysis of randomised controlled trials

Mario Álvarez-Jiménez, Sarah E. Hetrick, César González-Blanch, John F. Gleeson and Patrick D. McGorry

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Supplementary Material
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