BMJ Open  Does weight management research for adults with severe obesity represent them? Analysis of systematic review data

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ABSTRACT
Objective  Our objective was to determine the extent to which current evidence from long-term randomised controlled trials (RCTs) of weight management is generalisable and applicable to underserved adult groups with obesity (body mass index (BMI) ≥35 kg/m²).

Methods  Descriptive analysis of 131 RCTs, published after 1990–May 2017 with ≥1 year of follow-up, included in a systematic review of long-term weight management interventions for adults with BMI ≥35 kg/m² (the REBALANCE Project). Studies were identified from MEDLINE, EMBASE, PsychINFO, SCI, CENTRAL and from hand searching. Reporting of trial inclusion and exclusion criteria, trial recruitment strategies, baseline characteristics and outcomes were analysed using a predefined list of characteristics informed by the PROGRESS (Place of residence, Race/ethnicity/culture/language, Occupation, Gender/sex, Religion, Education, Socioeconomic status, Social capital)—Plus framework and the UK Equality Act 2010.

Results  Few (6.1%) trials reported adapting recruitment to appeal to underserved groups. 10.0% reported culturally adapting their trial materials. Only 6.1% of trials gave any justification for their exclusion criteria, yet over half excluded participation for age or mental health reasons. Just over half (58%) of the trials reported participants’ race or ethnicity, and one-fifth reported socioeconomic status.

Where outcomes were reported for underserved groups, the most common analysis was by sex (47.3%), followed by race or ethnicity (16.8%). 3.1% of trials reported outcomes according to socioeconomic status.

Discussion  Although we were limited by poor trial reporting, our results indicate inadequate representation of people most at risk of obesity. Guidance for considering underserved groups may improve the appropriateness of research and inform greater engagement with health and social care services.

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BACKGROUND
In high-income countries, and increasingly in low/middle-income countries, lower incomes, less education, lower socioeconomic status (SES) and disability are associated with greater risk of obesity for adults.1-5 While the underlying causes of obesity are varied, there is an increasing association between obesity and deprivation that is driving poorer health outcomes and increasing health inequalities.6 For example, most of these risk factors are stronger for women than men,1-3,7 although men with obesity may be less likely than women to undertake weight management programmes.8 Being an adult with obesity is associated with a lower health-related quality of life than a healthy adult of the same SES.9 More severe obesity, such as body mass index (BMI) ≥35 kg/m² or more, with its associated greater risks for comorbidities, reduced quality of life and premature mortality,10 is particularly related to lower SES1,11 and intellectual and physical disabilities.4 Poorer outcomes from COVID-19 are also strongly

STRENGTHS AND LIMITATIONS OF THIS STUDY
⇒ A unique analysis exploring whether randomised trials evaluating interventions for weight management for people with severe obesity consider the needs of underserved groups.
⇒ Data set includes up-to-date, best available randomised controlled trial evidence (considered to be the highest level of evidence to inform guidelines and clinical practice) identified using robust and exhaustive literature search strategies.
⇒ Data set includes wide-ranging weight management interventions delivered in different settings, from an international perspective.
⇒ Our analysis was limited by poor reporting of whether subgroup reporting by underserved groups was planned and whether trials were adequately powered to detect subgroup differences.
⇒ Non-randomised study designs, interventions with shorter follow-up and unpublished studies may have been more inclusive in their designs and reporting.
related to obesity, particularly severe obesity. For countries such as the USA, UK, Australia and New Zealand, some racial or ethnic groups may also be at much greater risk of obesity, especially severe obesity. Preventing obesity and providing effective interventions, particularly for people with more severe obesity, are therefore, a major public health challenge and vital in terms of addressing health inequalities. While organisations such as the US Food and Drug Administration, the National Institutes of Health (NIH) and the National Institute for Health Research (NIHR) have produced guidelines on the inclusion of individuals of all ages, sexes/genders, races and ethnicities, and other physical, sensory/perceptual, cognitive and emotional characteristics, there is a lack of accessible policy-ready evidence on what works in terms of interventions to reduce inequalities in obesity. It is also recognised that some groups (for example, socially disadvantaged, less educated, and minority race or ethnic groups) may be less likely to be recruited into randomised controlled trials (RCTs) for lifestyle interventions. Similarly, religion and sexual orientation have been linked to weight and body image. It is, therefore, important to understand the extent to which the current evidence base is applicable to those who are most at risk of experiencing poorer obesity-related health outcomes and have more severe obesity.

This study aimed to determine the extent to which the findings from intervention studies of weight management, as exemplified by long-term RCTs, are generalisable and applicable to those most at risk, particularly underserved groups with severe obesity. To examine these questions, we set out:

1. To describe inclusion and exclusion criteria for RCTs of adult weight management interventions, and in those trials:
   2. To describe efforts to tailor recruitment strategies to improve recruitment of people from underserved groups.
   3. To describe efforts to culturally adapt interventions to increase the accessibility or appeal to shared characteristics of an underserved group.
   4. To describe reported baseline characteristics and outcomes for these groups.

**METHODS**

Our data set comprised 131 RCTs included in a systematic review of weight management interventions for adults with BMI ≥ 35 kg/m², as part of the REBALANCE Project (REview of Behaviour And Lifestyle interventions for severe obesity: AN evidenCE synthesis; NIHR HTA Project (REview of Behaviour And Lifestyle interventions)). RCTs were restricted to publications after 1990 up to May 2017 to reflect more recent clinical practice. Literature searching was conducted in June 2016 and updated in April/May 2017. Details of the literature search method and search strategy are available in the online supplemental files. Trials had to report long-term data on weight change (≥1 year of follow-up) and include trial populations with a baseline mean or median BMI ≥ 35 kg/m². The decision to focus on long-term RCTs for this study was informed by the preference for high-quality, long-term evidence of lasting effectiveness in guideline documents and are, therefore, most likely to influence treatment policy decisions. Reports published as abstracts or conference proceedings only were excluded. Three reviewers screened titles, abstracts and full-text reports with a 10% quality assessment check. We attempted to contact the first, second and last authors of the main publications to identify all additional materials (ie, protocols, trial materials and diet books) to inform our data extraction for the main REBALANCE report. Full details of the completed REBALANCE Project, including the protocol, have been published.

In the absence of definitions of underserved groups, we identified underserved groups by using protected characteristics informed by the PROGRESS (Place of residence, Race/ethnicity/culture/language, Occupation, Gender/sex, Religion, Education, Socioeconomic status, Social capital)-Plus framework and the UK Equality Act 2010. Four reviewers (MA-M, MC, MI and CR) conducted double data coding of each RCT for their reporting of whether trials reported details of their inclusion and exclusion criteria, trial recruitment strategies, baseline characteristics and outcome reporting for the following characteristic groups, with disagreements resolved by consensus:

- Older age
- Physical health
- Mental health (including, but not limited to, depression, psychosis, schizophrenia, substance abuse and eating disorders)
- Comorbidities (eg, types 1 and 2 diabetes mellitus)
- Gender/sex (including RCTs recruiting only men or women)
- Sexual orientation
- Gender reassignment
- Marriage or civil partnership status
- Pregnancy
- Religion or belief
- Place of residence/housing (including residents of supported accommodation and homeowner status)
- Race or ethnicity
- Language
- Occupational status
- Education/literacy
- SES, including individual SES and participants recruited from rural or disadvantaged geographical locations
- Social capital (including social support networks and/or social isolation)
PROGRESS-Plus (personal characteristics associated with discrimination (eg, age, disability), features of relationships (eg, smoking parents, excluded from school), time-dependent relationships (eg, leaving the hospital, respite care, other instances where a person may be temporarily at a disadvantage))

For inclusion and exclusion criteria, trials were coded by predefined categories indicating whether any of the characteristic groups were clearly reported in the inclusion/exclusion criteria, or, where details were not reported, whether the setting of the trial encouraged/discouraged inclusion of individuals from a particular characteristic group (eg, recruitment was set in a health centre predominantly serving people from a characteristic group), or by whether it was unclear that the trial included/excluded people from any of the characteristic groups. For baseline and outcome reporting, we coded whether the protected characteristic was reported and, if reported, whether it was reported for individual treatment groups or the trial population as a whole. Where subgroup analyses were reported, we coded these according to whether it was clear/unclear from the study report that analyses were preplanned. For a trial to be coded as having adapted their recruitment strategy for an underserved group, additional efforts to employ strategies that would appeal to that particular group (eg, held recruitment days in particular settings or developed recruitment materials in multiple languages) had to be demonstrated. Trials that solely recruited from one characteristic group using conventional recruitment methods (eg, newspaper or radio advertisements) were not coded as having an adapted recruitment strategy. Similarly, trials had to demonstrate that their interventions were designed with an underserved group in mind to be coded as having delivered a culturally adapted intervention. The focus of this study was to provide a description of trial methods and trial reporting to answer each of our research questions in relation to underserved groups; therefore, no formal statistical analysis was conducted.

Patient and public involvement

Although we did not consult patient representatives for this particular analysis, three patient representatives were members of the REBALANCE Project Advisory Group, who contributed to developing the research questions, data interpretation and reporting of the research findings.

RESULTS

From the total of 131 included trials, 19 were identified from database searching and 112 were identified from autoalert searching. The Preferred Reporting Items for Systematic Reviews and Meta-Analyses flow chart and list of included studies are presented in the online supplemental files. Of the 131 trials, 41 (31.3%) provided us with additional materials for their publications for the main REBALANCE report, although most of the information for the current analysis was obtained from the primary publications. The majority (81 of 131, 61.8%) of included studies were set in North America (80 in the USA and 1 in the USA and Canada), 41 out of 131 (31.3%) were in European countries (including 8 in the UK), 8 (6.1%) were in the Southern Hemisphere (6 in Australia, 1 in New Zealand and 1 in Australia and New Zealand) and 1 in Brazil. None of the trials were set in low-income countries. Just under half (62 of 131, 47.3%) of the studies were published between 2011 and 2017. Five (3.8%) trials were linked to publications reporting qualitative data. Few trials had follow-up duration longer than 12 months, with the exception of the US Look AHEAD trial (median duration of 9.6 years), and four trials with follow-up times of 5 years. The Look AHEAD trial was the largest trial, including over 5000 participants. Interventions were wide ranging, including very low calorie (19 of 131, 14.5%), orlistat (12 of 131, 9.2%), bariatric surgery (11 of 131, 8.4%) and other lifestyle weight management programmes incorporating diet and physical activity advice (89 of 131, 67.9%). Details of the characteristics of the included studies can be found in the online supplemental files of the REBALANCE report.

Trial recruitment

More than half of the trials (71 of 131, 54%) recruited participants either solely or partially through a health service provider, for example, either solely from outpatient clinics and general practices, or by physician referral and targeted mailing. Recruitment methods were unclear or not reported in 14 trials (10.7%). Recruitment methods for the other trials were mainly advertisements in local newspapers or other media. Based on their reporting, only three (2.3%) trials were judged to have adapted their recruitment strategies to appeal to underserved groups. These preplanned strategies included holding pre-recruitment presentations in US schools, recruitment events at football stadiums of Scottish Premier League football clubs, and having bilingual staff take informed consent and provide written consent forms in both English and Spanish languages. It was unclear in a further five (3.8%) trials whether recruitment strategies had been adapted beyond conventional methods.

Regarding adaptions to interventions, seven trials (5.3%, six from the USA and one from New Zealand) recruited participants from diverse racial or ethnic groups and reported cultural adaptations to their interventions. Five (3.8%) of these trials included advice on regional or culturally adapted recipes and foods for specific ethnic groups. Two trials (1.5%) had interventions that were delivered by bilingual staff. One trial also reported that the intervention was designed for delivery in populations with limited literacy and numeracy and impaired access to health-promoting resources. While two trials recruited participants from workplace settings (an automobile manufacturer and a university; both were not considered to meet the PROGRESS-Plus occupation definition), and
one trial recruited married participants,\textsuperscript{70} the trials did not report any attempts to alter their recruitment strategies or interventions to appeal to underserved occupation or social capital groups. The trials did not report recruitment strategies or adaptations to interventions for sexual orientation.

A further five trials (all from the USA)\textsuperscript{71–75} sought recruitment from specific racial or ethnic groups and reported intervention adaptations to increase cultural salience. Four of these trials\textsuperscript{71–74} included culturally specific dietary advice and recipes. One trial described including bilingual interventions\textsuperscript{21} and three trials described including interventionists from specific racial or ethnic groups.\textsuperscript{72–74} Two trials\textsuperscript{73 74} described using logos and programme identification ‘for African-Americans’, with one of these trials\textsuperscript{73} including a video greeting from an African-American principal investigator.

The number of trials reporting inclusion and exclusion criteria by protected characteristic groups is presented in table 1. Four older trials (3.1\%)\textsuperscript{47 52 76 77} did not report any inclusion criteria, and inclusion criteria were unclear in one further study.\textsuperscript{42} Seven (5.3\%)\textsuperscript{52 70 78–82} trials did not report any explicit exclusion criteria and did not report that they had no exclusion criteria. Eight (6.1\%) trials reported either full\textsuperscript{36 45 72 83 84} or partial\textsuperscript{60 85} justification for their exclusion criteria. Justification for exclusion criteria included prevention of poor adherence and losses to follow-up,\textsuperscript{38 45 84 85} such as substance abuse, mental health problems or cognitive impairment (that might, in the opinion of the investigators, hinder participation), lower BMI cut-offs for Asian people,\textsuperscript{60} non-English-language speakers where the intervention required English language comprehension,\textsuperscript{84} influence of pregnancy and breast feeding on weight,\textsuperscript{84} taking medications that influence weight,\textsuperscript{72} and contraindications or safety concerns associated with participating in the intervention (eg, risk of participants with cardiovascular disease participating in exercise programmes).\textsuperscript{36 45 84 85} Over half (58.0\%) of the trials reported excluding people with mental health conditions and 44.2\% excluded people with substance abuse or addiction issues. The majority of trials also excluded

| Table 1 | The number (and per cent) of trials (n=131) reporting inclusion and exclusion criteria by protected characteristics included in the REBALANCE systematic review of RCTs |
| Protected characteristic is reported in the inclusion criteria, n (%) | Protected characteristic is not reported in inclusion criteria, but an effort was made to recruit from the protected characteristic group, n (%) | Unclear if the protected characteristic was targeted for inclusion or if the trial unintentionally recruited solely/mainly from the protected characteristic group, n (%) | Protected characteristic is reported in the exclusion criteria, or the reported exclusion criteria clearly excluded the protected characteristic, n (%) |
| Place of residence/housing | 5 (3.8) | 0 | 0 | 0 |
| Race/ethnicity | 7 (5.3) | 5 (3.8) | 2 (1.5) | 1 (0.8) |
| Occupation status | 0 | 0 | 0 | 0 |
| Women only | 19 (14.5) | 0 | 4 (3.1) | 0 |
| Pregnancy | 0 | 0 | 0 | 72 (54.9) |
| Men only | 4 (3.0) | 0 | 1 (0.76) | 5 (3.8) |
| Religion/belief | 1 (0.8) | 0 | 0 | 0 |
| Education/literacy | 0 | 0 | 0 | 0 |
| Socioeconomic status | 3 (2.3) | 3 (2.3) | 0 | 0 |
| Marital status | 1 (0.8) | 0 | 0 | 1 (0.8) |
| Older age | 2 (1.5) | 1 (0.8) | 0 | 82 (62.6)* |
| Physical health | 10 (7.6) | 0 | 0 | 51 (38.9) |
| Diabetes type 1 | 0 | 0 | 0 | 15 (11.5) |
| Diabetes type 2 | 28 (21.4) | 0 | 0 | 29 (22.1) |
| Diabetes (type 1 and 2 or type not reported) | 0 | 0 | 0 | 3 (2.3) |
| Mental health | 6 (4.6) | 0 | 0 | 76 (58.0) |
| Substance abuse | 0 | 0 | 0 | 58 (44.2) |
| Eating disorder | 0 | 0 | 0 | 35 (26.7) |
| Language | 0 | 0 | 0 | 16 (12.2) |

*Includes eight RCTs recruiting participants up to 75 years, one RCT recruited participants up to 76 years and three RCTs recruited participants aged up to 80 years.

RCTs, randomised controlled trials; REBALANCE, REview of Behaviour And Lifestyle interventions for severe obesity: AN evidenCE synthesis.
adults from older age groups and based on current or planned pregnancy.

Twenty-one (16.0%) trials were judged to have inclusion criteria that might have implicitly excluded certain disadvantaged groups, such as people who do not have healthcare insurance, people who do not belong to a particular religious community group, people without regular internet or telephone access, and English language comprehension. In a further 40 (30.5%) trials, it was unclear if trial recruitment could have implicitly excluded disadvantaged groups. Few trials reported their inclusion criteria so as to include particular underserved groups or were judged to have made efforts to maximise trial recruitment from these groups. When trial recruitment was targeted, this was usually to recruit women only (19 trials) or people with type 2 diabetes (28 trials).

Reporting of baseline characteristics
Details of the number of trials reporting baseline characteristics of their participants by each of the protected characteristic groups are presented in table 2. The majority of trials reported age (99.2%) and sex (97.7%) in their description of baseline participant characteristics. Just over half of the trials (58.0%) reported race or ethnicity. Education history was less well reported (40.5%). SES was reported by 22.9%, and occupation status by 21.4%. Of the trials that were not specifically for people with diabetes, six (4.6%) included diabetes in their reporting of baseline characteristics. Two (1.5%) trials reported whether people lived alone or not (coded as social capital). Few trials reported details of the other protected characteristics, and none reported details of gender reassignment, sexual orientation or pregnancy.

Outcome reporting
Details of the number of trials reporting outcomes by each of the protected characteristics are shown in table 3. Very few trials reported outcomes by protected characteristic groups. Where outcomes were reported by protected characteristics, the most common group was sex (47.3%), followed by race or ethnicity (16.8%).

DISCUSSION
Main findings
Our findings demonstrate that most trialists testing weight management strategies to help adults with severe obesity fail to consider populations who are most at risk of poorer health outcomes. Almost all trials were from high-income countries, where lower SES and income are associated with a greater prevalence of obesity, particularly severe obesity. Few trials reported adapting recruitment to appeal to underserved groups or reported culturally adapting their trial materials for ethnic groups or people with limited English language literacy or numeracy. This is concerning as limiting the accessibility or appeal of trials could limit the representativeness of the trial population, and thus limit the generalisability of trial findings. Only 6.1% of trials gave any justification for their exclusion

Table 2 The number (and per cent) of trials (n=131) reporting protected characteristics at baseline in the REBALANCE systematic review of RCTs

<table>
<thead>
<tr>
<th>Protected characteristic</th>
<th>Protected characteristic is reported at baseline for each intervention arm</th>
<th>Protected characteristic is reported at baseline for the whole trial</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>126 (96.2%)</td>
<td>4 (3.0%)</td>
<td>130 (99.2%)</td>
</tr>
<tr>
<td>Physical health</td>
<td>10 (7.6%)</td>
<td>0</td>
<td>10 (7.6%)</td>
</tr>
<tr>
<td>Mental health</td>
<td>10 (7.6%)</td>
<td>0</td>
<td>10 (7.6%)</td>
</tr>
<tr>
<td>Diabetes</td>
<td>6 (4.6%)</td>
<td>0</td>
<td>6 (4.6%)</td>
</tr>
<tr>
<td>Sex</td>
<td>126 (96.2%)</td>
<td>2 (1.5%)</td>
<td>128 (97.7%)</td>
</tr>
<tr>
<td>Gender reassignment</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Sexual orientation</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Marriage/civil partnership status</td>
<td>38 (29.0%)</td>
<td>0</td>
<td>38 (29.0%)</td>
</tr>
<tr>
<td>Pregnancy</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Place of residence/housing</td>
<td>6 (4.6%)</td>
<td>0</td>
<td>6 (4.6%)</td>
</tr>
<tr>
<td>Occupation status</td>
<td>27 (20.6%)</td>
<td>1 (0.8%)</td>
<td>28 (21.4%)</td>
</tr>
<tr>
<td>Education/literacy</td>
<td>51 (38.9%)</td>
<td>2 (1.5%)</td>
<td>53 (40.5%)</td>
</tr>
<tr>
<td>Socioeconomic status</td>
<td>29 (22.1%)</td>
<td>1 (0.8%)</td>
<td>30 (22.9%)</td>
</tr>
<tr>
<td>Social capital</td>
<td>2 (1.5%)</td>
<td>0</td>
<td>2 (1.5%)</td>
</tr>
<tr>
<td>Religion/belief</td>
<td>1 (0.8%)</td>
<td>0</td>
<td>1 (0.8%)</td>
</tr>
<tr>
<td>Race/ethnicity</td>
<td>74 (56.5%)</td>
<td>2 (1.5%)</td>
<td>76 (58.0%)</td>
</tr>
<tr>
<td>PROGRESS-Plus</td>
<td>2 (1.5%)</td>
<td>0</td>
<td>2 (1.5%)</td>
</tr>
</tbody>
</table>

<http://bmjopen.bmj.com/> BMJ Open: first published as 10.1136/bmjopen-2021-054459 on 31 May 2022. Downloaded from http://bmjopen.bmj.com/ on December 2, 2022 by guest. Protected by copyright.
criteria, yet more than half excluded participation for age or mental health reasons. Where justification for exclusion was reported, the rationale included excluding people who were deemed likely to have poor intervention adherence or were more likely to be lost to follow-up, such as people with substance abuse, cognitive impairment or mental health problems. Excluding these groups could lead to an unrealistic estimation of the real-world effectiveness of interventions. Just over half of the trials reported participants’ race or ethnicity, and only around one-fifth reported SES. Where outcomes were reported for underserved groups, the most common analysis was by sex (47.3%), followed by race or ethnicity (16.8%); however, where analyses were presented as subgroups, it was often unclear whether these analyses were planned or unplanned. Similarly, some smaller trials might have been underpowered to detect differences in treatment effects between subgroups, but this was also unclear from trial reporting. This finding was also demonstrated by Liu and colleagues, who highlighted a lack of transparent reporting of intentions to analyse race and ethnicity subgroups in Cochrane intervention reviews. Only 3.1% of the trials we reviewed reported outcomes according to SES. Few trials reported outcomes by the other protected characteristics.

Although we were limited by the available information in the published reports, our findings are concerning. In almost all trials, it is difficult to assess the generalisability of findings to the wider population of adults with severe obesity. There is clear evidence that underserved groups with lower incomes, less education, lower SES, intellectual and physical disabilities, or poorer mental health are more at risk of obesity, particularly severe obesity, in high-income countries, especially the USA which provided the majority of trials examined. We do not have relevant data to be able to comment on the reasons for poor reporting. Nevertheless, the lack of reporting for characteristics reflecting underserved groups suggests that trial investigators did not consider or faced barriers that prevented their inclusion in the design, recruitment, and analysis or reporting of their interventions.

Our finding that few trials adapted their recruitment methods or interventions to appeal to underserved groups suggests lack of engagement with underserved people with obesity in the design of services. This is important, given that a systematic review of qualitative research by Sutcliffe and colleagues showed how service users have perspectives that should inform weight management services to improve their reach. From systematic reviews, researchers have clearly demonstrated the need

### Table 3 The number (and per cent) of trials (n=131) reporting outcome data for protected characteristics in the REBALANCE systematic review of RCTs

<table>
<thead>
<tr>
<th>Trial recruitment was targeted at people from the protected characteristic group</th>
<th>One or more outcome(s) reported for the protected characteristic in planned subgroup analysis</th>
<th>One or more outcome(s) reported for the protected characteristic—unclear if subgroup analysis was preplanned</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Older age</td>
<td>2 (1.5%)*</td>
<td>2 (1.5%)</td>
<td>3 (2.3%)</td>
</tr>
<tr>
<td>Physical health</td>
<td>10 (7.6%)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Mental health</td>
<td>6 (4.6%)</td>
<td>0</td>
<td>2 (1.5%)</td>
</tr>
<tr>
<td>Diabetes</td>
<td>28 (21.3%)</td>
<td>0</td>
<td>1 (0.8%)</td>
</tr>
<tr>
<td>Sex</td>
<td>23 (17.5%)†</td>
<td>17 (13.0%)</td>
<td>22 (16.8%)</td>
</tr>
<tr>
<td>Gender reassignment</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Sexual orientation</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Marriage/civil partnership status</td>
<td>1 (0.8%)</td>
<td>1 (0.8%)</td>
<td>3 (2.3%)</td>
</tr>
<tr>
<td>Pregnancy</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Place of residence/housing</td>
<td>0</td>
<td>2 (1.5%)</td>
<td>0</td>
</tr>
<tr>
<td>Occupation status</td>
<td>2 (1.5%)</td>
<td>2 (1.5%)</td>
<td>4 (3.1%)</td>
</tr>
<tr>
<td>Education/literacy</td>
<td>0</td>
<td>1 (0.8%)</td>
<td>6 (4.6%)</td>
</tr>
<tr>
<td>Socioeconomic status</td>
<td>5 (3.8%)</td>
<td>4 (3.1%)</td>
<td>3 (2.3%)</td>
</tr>
<tr>
<td>Social capital</td>
<td>1 (0.8%)</td>
<td>3 (2.3%)</td>
<td>2 (1.5%)</td>
</tr>
<tr>
<td>Religion/belief</td>
<td>1 (0.8%)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Race/ethnicity</td>
<td>8 (6.1%)</td>
<td>5 (3.8%)</td>
<td>9 (6.9%)</td>
</tr>
<tr>
<td>PROGRESS-Plus</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

*Both trials recruited participants aged ≥65 years.
†Nineteen women-only trials, four men-only trials.
PROGRESS, Place of residence, Race/ethnicity/culture/language, Occupation, Gender/sex, Religion, Education, Socioeconomic status, Social capital; RCTs, randomised controlled trials; REBALANCE, REview of Behaviour And Lifestyle interventions for severe obesity; AN evidence synthesis.
to involve communities in all stages of research in order to enhance the engagement and generalisability of that research, acknowledging that this requires extended time frames and greater costs.15 16 For example, Ni She and colleagues109 undertook a rapid realist review of the mechanisms and resources needed to engage underserved, seldom-heard groups in health and social care research, with items grouped by an expert panel under the headings of environmental and social planning, service provision, guidelines, fiscal measures, communication and marketing, and regulation and legislation. In the USA, Arnegard and colleagues103 have also called on the NIH’s stakeholder groups to redouble their efforts to encourage sex/gender-aware reporting of biomedical investigations. We endorse this call following the findings from our previous systematic review of weight management interventions for men with obesity.8 Our review highlighted the paucity of evidence for men, who are less likely to take part in weight management interventions, and the lack of engagement of men in all aspects of intervention design, and optimal trial recruitment processes of weight management.8

While the reasons for the under-representation of underserved groups in RCTs are likely to be complex and multifaceted, with many known and unknown barriers to participation, there is evidence that, for some groups, willingness to participate is not a predominant factor.104 Mindful of the need to improve the engagement of underserved groups in research in the UK, the NIHR set up the INCLUDE Project,19 which has led to the INCLUDE ethnicity framework105; providing four key inequalities for adults with obesity114 found that primary care-delivered tailored weight loss programmes and group weight loss interventions had the most evidence of potential effectiveness in reducing obesity, at least in the short term, among low-income women, but there were few individual-level intervention studies and a lack of long-term evidence of effectiveness.

**Strengths and limitations**

We used categories informed by the PROGRESS-Plus framework32 and the UK Equality Act 201035 as the key characteristics for identifying those underserved participants who should be considered for study design, public–patient involvement, recruitment, analysis and reporting, not just for trials of weight management, but trials generally. O’Neill and colleagues32 have shown how the prior PROGRESS framework can be used as an equity lens for systematic reviews and methodological studies; however, NIHR’s INCLUDE Project has recently published a more extensive list of categories of underserved groups to consider with regard to representation in trials.115

Our literature search attempted to identify all long-term randomised trials published since 1990 for adults with BMI ≥35 kg/m² irrespective of the type of lifestyle intervention, including comparisons with orlistat and bariatric surgery. Although we included publications in any language from any country, we cannot exclude the possibility that we failed to find some trials, particularly those from low-income countries, which might not be listed in the databases we searched.

While we originally contacted authors for all available additional materials relating to our main research question for the REBALANCE Project,10 we did not recontact authors to obtain additional information relating to the current analysis. We were also limited by poor reporting by trial authors. Some trials were statistically underpowered to detect subgroup differences, and this might explain under-reporting of underserved characteristics; however, this was unclear from the trial reports. It is also possible that some trialists were unable to obtain relevant baseline data for some underserved groups if this was deemed sensitive by an ethics committee, for example, sexual orientation. Nevertheless, we consider that most characteristics are pivotal to interpreting these trials into real-world guidance and services, so we would expect their presentation in trial publications, especially at baseline. We assessed long-term RCT evidence because it is most likely to inform guidance on weight management.28–31 Other study designs, interventions with shorter follow-up and unpublished studies may have been more inclusive in their designs and reporting.

**Recommendations for research**

Trialists should improve reporting of their justification of inclusion and exclusion criteria to meet current Consolidated Standards of Reporting Trials (CONSORT) statement guidelines,116 and report sufficient data to allow comparisons between their populations and the populations for whom the interventions apply. Including core criteria for baseline reporting within the CONSORT checklist116 could help to improve the completeness of reporting of these factors. NIHR’s INCLUDE Project’s ethnicity framework provides important factors to consider with regard to ethnic groups (https://www.trial-forge.org/trial-forge-centre/include/), which can help provide transferable considerations for other underserved...
groups. However, a wider equity lens may be needed in the face of groups with multiple disadvantages. Although guidance for research will aid considerations of equity, we do not yet have ways of assessing when proportional representation in larger trials and subgroup reporting for underserved groups is insufficient. This research should be explicitly conducted with and for these underserved groups, ensuring user involvement in all stages of the research process.

CONCLUSIONS AND RECOMMENDATIONS FOR PRACTICE

Long-term RCTs of weight management in people with BMI $\geq$35 kg/m² have inadequate representation of and engagement with underserved groups, who are particularly relevant for health and social care services. Thus, guidance for weight management research on how to improve the representation of underserved groups in clinical trials may improve the appropriateness of that research and help inform greater engagement of underserved communities with health and social care services.

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Competing interests None declared.

Patient and public involvement Patients and/or the public were not involved in the design, or conduct, or reporting, or dissemination plans of this research.

Patient consent for publication Not required.

Ethics approval Ethical approval was not required.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available upon reasonable request. All data relevant to the study are included in the article, uploaded as supplemental information, or are available from the NIHR journals library (REBALANCE) Review of Behaviour And Lifestyle interventions for severe obesity: AN evidence synthesis (nihr.ac.uk)

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REFERENCES

1 Booth HP, Charlton J, Guilford MC. Socioeconomic inequality in morbid obesity with body mass index more than 40 kg/m² in the United States and England. SSM Popul Health 2017;3:172–8.


101 De las Nueces D, Harwood K, DiGirolamo A. A systematic review of community-based participatory research to enhance clinical trials in racial and ethnic minority groups. Health Serv Res 2012;47:1383–86.


105 Trial Forge [Internet]. Aberdeen, Scotland: Trial Forge; 2020 [cited February 2012]. The INCLUDE ethnicity framework. Available: https://www.trialforge.org/trial-forge-centre/include/


109 Trivedi RB, Humphreys K. Participant exclusion criteria in treatment research on neurological disorders: are unrepresentative study samples problematic? J Neurol Neurosurg Psychiatry 2014;85:489–96.


113 Olstad DL, Ancilotto R, Teychenne M, et al. Can targeted policies reduce obesity and improve obesity-related behaviours in black and white adults?


Appendix 1  Literature search strategies

Literature searching was conducted in June 2016 and updated in April/May 2017. The following databases were searched without language restriction: MEDLINE, MEDLINE Epub Ahead of Print & MEDLINE In-Process, PsychINFO, Cumulative Index to Nursing and Allied Health Literature (CINAHL), Science Citation Index (SCI), Cochrane Central Register of Controlled Trials (CENTRAL), and ClinicalTrials.gov. Ovid MEDLINE and EMBASE autoalerts have been in place since 2002 and results are screened regularly for long-term RCTs on obesity management in adults (≥ 1 year of follow-up). Copies of relevant reports are retained and the reviewers hand-searched these reports eligible studies. Therefore, search strategies for MEDLINE and EMBASE excluded the results of the autoalert search (using the Boolean operator NOT). A supplementary search of MEDLINE was undertaken to identify systematic reviews of severe or morbid obesity, and reference lists were scrutinised for additional studies.

RCTs for Review 1 (weight loss or weight maintenance programmes for adults with obesity and Review 5 (weight loss or weight maintenance programmes for adults with obesity compared with bariatric surgery)
MEDLINE and EMBASE
Ovid multifile search: http://shibboleth.ovid.com/
Database: Embase <1988 to 2017 Week 17>, Ovid MEDLINE(R) Epub Ahead of Print, In-Process & Other Non-Indexed Citations, Ovid MEDLINE(R) Daily and Ovid MEDLINE(R) <1946 to Present> 25th April 2017
Date of Search: 25th April 2017
1 Obesity, Morbid/dh, dt, su, th use ppez
2 Morbid Obesity/dt, su, th use emed
3 Obesity, Morbid/ use ppez
4 Morbid Obesity/ use emed
5 (obes$ adj3 (morbid$ or severe$)).tw.
6 (morbid obesity or severe obesity).kw.
7 (bmi adj ("27$" or "28$" or "29$")) .tw.
8 (bmi adj ("30$" or "31$" or "32$" or "33$" or "34$" or "35$" or "36$" or "37$" or "38$" or "39$")).tw.
9 (bmi adj ("40$" or "41$" or "42$" or "43$" or "44$" or "45$" or "46$" or "47$" or "48$" or "49$")) .tw.
10 (bmi adj ("50$" or "51$" or "52$" or "53$" or "54$" or "55$")).tw.
11 (body mass index adj ("35$" or "36$" or "37$" or "38$" or "39$")).tw.
12 (body mass index adj ("40$" or "41$" or "42$" or "43$" or "44$" or "45$" or "46$" or "47$" or "48$" or "49$")) .tw.
13 (body mass index adj ("50$" or "51$" or "52$" or "53$" or "54$" or "55$")).tw.
14 or/3-12
15 exp bariatric surgery/
16 bariatric.tw,kw.
17 (gastric adj3 (bypass or bypass or band$ or balloon$)).tw.
18 (gastroplast$ or gastrect$).tw.
19 (jejunoileal adj3 (bypass or by pass)).tw.
20 lipectom$.tw
21 diet therapy/ use ppez or caloric restriction/ use ppez or diet, carbohydrate-restricted/ use ppez or diet, fat-restricted/ use ppez or diet, reducing/ use ppez
22 diet/ use emed or low calory diet/ use emed or low carbohydrate diet/ use emed
23 (diet adj3 (restrict$ or reduc$ or modif$ or calorie$)).tw.
24 (calori$ adj3 (reduc$ or restrict$ or limit$)).tw.
25 exp anti-obesity agents/ use ppez
26 tetrahydrolipstatin/ use emed
27 sibutramine/ use emed
28 rimonabant/ use emed
29 (appetite adj3 (reduc$ or depress$)).tw.
30 (orlistat or xenical).tw,rm.
31 (sibutramin$ or arcalion or reductil or medaria or meridia).tw,rm.
32 (rimonabant or acomplia or zimulti).tw,rm.
33 exp exercise/
34 exp sports/
35 (exercise or aerobic or physical activ$).tw.
36 weight loss/ use ppez
37 weight reduction/ use emed
38 (weight adj3 (reduc$ or loss or lose)).tw.
39 Weight Reduction Programs/ use ppez
40 exp Body Weight Management/ use emed
41 Behavior Therapy/
42 Health Education/
43 Counseling/ use ppez
44 Nutritional counseling/ use emed
45 Life Style/ use ppez
46 Lifestyle modification/ use emed
47 (modif$ adj3 (behavior or diet or eat? or eating)).tw.
48 ((diet or weight) adj3 (educat$ or counsel$)).tw.
49 or/15-46
50 1 or 2 or (14 and 49)
51 randomized controlled trial.pt.
52 controlled clinical trial.pt
53 randomi?ed.ab.
54 placebo.ab.
55 drug therapy.fs.
56 randomly.ab.
57 trial.ab.
58 groups.ab
59 nonhuman/ not human/
60 exp clinical trial/ use emed
61 randomization/ use emed
62 51 or 52 or 53 or 54 or 55 or 56 or 57 or 58 or 60 or 61
63 exp animals/ use ppez not humans/ use ppez
64 nonhuman/ use emed or animal/ use emed
65 64 not exp human/ use emed
66 62 not (63 or 65)
67 50 and 66
68 67 not (abstract or letter or note or editorial or comment).pt.
69 limit 68 to yr="1990 -Current"
70 obesity/
71 (obesity adj2 (morbid or diabet$)).tw.
72 obesity, morbid/ use ppez
73 morbid obesity/ use emed
74 obes$.tw.
75 weight loss/ use ppez
76 weight reduction/ use emed
77 (weight adj1 (los$ or reduc$ or maint$ or control)).tw.
78 (diet adj5 weight).tw.
79 overweight.tw.
80 or/70-79
81 80 and 66
82 69 not 81

PsycINFO
Ovid: http://shibboleth.ovid.com/
Database: PsycINFO <1987 to April Week 3 2017>

Date of Search: 25th April 2017
1 (obes$ adj3 (morbid$ or severe$)).tw.
2 (bmi adj ("27$" or "28$" or "29$"))tw.
3 (bmi adj ("30$" or "31$" or "32$" or "33$" or "34$" or "35$" or "36$" or "37$" or "38$" or "39$"))tw.
4 (bmi adj ("40$" or "41$" or "42$" or "43$" or "44$" or "45$" or "46$" or "47$" or "48$" or "49$"))tw.
5 (bmi adj ("50$" or "51$" or "52$" or "53$" or "54$" or "55$"))tw.
6 (body mass index adj ("27$" or "28$" or "29$"))tw.
7 (body mass index adj ("30$" or "31$" or "32$" or "33$" or "34$" or "35$" or "36$" or "37$" or "38$" or "39$"))tw.
8 (body mass index adj ("40$" or "41$" or "42$" or "43$" or "44$" or "45$" or "46$" or "47$" or "48$" or "49$"))tw.
9 (body mass index adj ("50$" or "51$" or "52$" or "53$" or "54$" or "55$"))tw. (7)
10 or/1-9
11 clinical trials/
12 randomi?ed.tw,md.
13 randomly.tw,md.
14 trial?.tw,md
15 or/11-14
16 10 and 15

CINAHL
http://search.ebscohost.com
!990- 25th April 2017
**Date of Search: 25th April 2017**

S1 (MH "Obesity, Morbid") OR TX ( obes* N3 (morbid or sever*))
S2 TX ( bmi N3 (27 or 28 or 29 or 30 or 31 or 32 or 33 or 34 or 35 or 36 or 37 or 38 or 39). ) OR TX ( bmi N3 (40 or 41 or 42 or 43 or 44 or 45 or 46 or 47 or 48 or 49). ) OR TX ( bmi N3 (50 or 51 or 52 or 53 or 54 or 55). )
S3 TX ( body mass index N3 (35 or 36 or 37 or 38 or 39). ) OR TX ( body mass index N3 (40 or 41 or 42 or 43 or 44 or 45 or 46 or 47 or 48 or 49). ) OR TX ( body mass index N3 (50 or 51 or 52 or 53 or 54 or 55). )
S4 S1 OR S2 OR S3
S5 (MH "Bariatric Surgery+")
S6 TX bariatric OR TX gastroplasty
S7 TX bariatric OR TX gastroplast* AND TX gastrect*
S8 TX ( gastric N3 (by pass or by pass or band* or balloon*) ) OR TX ( jejunooideal N3 (bypass or by pass ) OR TX lipectom*
S9 (MH "Diet Therapy") OR (MH "Diet, Reducing") OR (MH "Restricted Diet") OR (MH "Diet, Fat-Restricted") OR (MH "Diet, Low Carbohydrate")
S10 TX ( diet N3 (restrict* or reduc* or modif* or calorie*)) ) OR TX ( calori* N3 (reduc* or restrict* or limit*). )
S11 (MH "Antiobesity Agents+")
S12 TX ( orlistat or xenical ) OR TX ( sibutramin$ or arcalion or reductil or medaria or meridia )
OR TX ( rimonabant or acomplia or zimulti )
S13 (MH “Exercise+”)
S14 TX exercise or aerobic or physical activ*).
S15 (MH “Physical Fitness”)
S16 (MH "Weight Loss")
S17 TX weight N3 (reduc* or loss or lose)).
S18 (MH "Weight Reduction Programs”)
S19 (MH "Counseling")
S20 (MH "Life Style") OR (MH "Life Style Changes")
S21 TX ( modif* N3 (behavior or behaviour or diet or eat* or eating ) OR TX ( (diet or weight) N3 (educat* or counsel*))).

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placed on this supplemental material which has been supplied by the author(s) BMJ Open
doi: 10.1136/bmjopen-2021-054459
S22 S5 OR S6 OR S7 OR S8 OR S9 OR S10 OR S11 OR S12 OR S13 OR S14 OR S15 OR S16
OR S17 OR S18 OR S19 OR S21
S23 S4 AND S22 Limiters - Published Date: 19900101-20161231
S24 S4 AND S22 Limiters - Exclude MEDLINE records
Science Citation Index
www.webofknowledge.com
1988 - 25th April 2017
Date of Search: 25th April 2017
# 1 (TS=(morbid NEAR/3 obes$)) AND DOCUMENT TYPES: (Article)
# 2 (TS=(severe$ NEAR/3 obes$)) AND DOCUMENT TYPES: (Article)
# 3 (TS=(bmi NEAR/3 (27 or 28 or 29 or 30 or 31 or 32 or 33 or 34 or 35 or 36 or 37 or 38 or
39))) AND DOCUMENT TYPES: (Article)
# 4 ((TS=(bmi NEAR/3 (40 or 41 or 42 or 43 or 44 or 45 or 46 or 47 or 48 or 49)))) AND
DOCUMENT TYPES: (Article)
# 5 ((TS=(bmi NEAR/3 (50 or 51 or 52 or 53 or 54 or 55)))) AND DOCUMENT TYPES: (Article)
# 6 #5 OR #4 OR #3 OR #2 OR #1
# 7 (TS=(randomized or randomised or randomly)) AND DOCUMENT TYPES: (Article)
# 8 (TS=clinical trial*) AND DOCUMENT TYPES: (Article)
# 9 #8 OR #7
# 10 #9 AND #6
# 11 (TS=(bariatric or gastroplast* or gastrect*)) AND DOCUMENT TYPES: (Article)
# 12 (TS=((gastric or jejunoileal) NEAR/3 bypass)) AND DOCUMENT TYPES: (Article)
# 13 ((TS=((gastric or jejunoileal) NEAR/3 "by pass"))) AND DOCUMENT TYPES: (Article)
# 14 (TS=(gastric NEAR/3 (band* or balloon*))) AND DOCUMENT TYPES: (Article)
# 15 (TS=(diet NEAR/3 (restrict* or reduc* or modif* or calorie*))). AND DOCUMENT TYPES: (Article)
# 16 (TS=(calori* NEAR/3 (reduc* or restrict* or limit*)].) AND DOCUMENT TYPES: (Article)
# 17 (TS=(appetite NEAR/3 (reduc* or depress*))). AND DOCUMENT TYPES: (Article)
# 18 (TS=(orlistat or xenical)). AND DOCUMENT TYPES: (Article)
# 19 (TS=(sibutramin or arcalion or reductil or medaria or meridia.)) AND DOCUMENT TYPES: (Article)
# 20 (TS=(rimonabant or acomplia or zimulti.)) AND DOCUMENT TYPES: (Article)
# 21 (TS=(exercise or aerobic or physical active*).) AND DOCUMENT TYPES: (Article)
# 22 (TS=(weight NEAR/3 (reduc* or loss or lose))). AND DOCUMENT TYPES: (Article)
# 23 (TS=(modif* NEAR/3 (behavior or behaviour or diet or eat? or eating))). AND DOCUMENT TYPES: (Article)
# 24 (TS=(diet NEAR/3 (educat* or counsel*)].) AND DOCUMENT TYPES: (Article)
# 25 (TS=(weight NEAR/3 (educat* or counsel*)]).) AND DOCUMENT TYPES: (Article)
# 26 #25 OR #24 OR #23 OR #22 OR #21 OR #20 OR #19 OR #18 OR #17 OR #16 OR #15 OR #14 OR #13 OR #12 OR #11
#27 #10 AND #26

Cochrane Central Register of Controlled Trials
www.thecochranelibrary.com

Date of Search: 25th April 2017

#1 MeSH descriptor: [Obesity, Morbid] explode all trees and with qualifier(s): [Diet therapy - DH, Surgery - SU, Therapy - TH]
#2 MeSH descriptor: [Obesity, Morbid] explode all trees
#3 obes* near/3 (morbid* or severe*):ti,ab,kw (Word variations have been searched)
#4 bmi near/1 (27* or 28* or 29* or 30* or 31* or 32* or 33* or 34* or 35* or 36* or 37* or 38* or 39*):ti,ab,kw or bmi near/1 (40* or 41* or 42* or 43* or 44* or 45* or 46* or 47* or 48* or 49*):ti,ab,kw or bmi near/1 (50* or 51* or 52* or 53* or 54* or 55*):ti,ab,kw (Word variations have been searched)
#5 body mass index near/1 (27* or 28* or 29* or 30* or 31* or 32* or 33* or 34* or 35* or 36* or 37* or 38* or 39*):ti,ab,kw or body mass index near/1 (40* or 41* or 42* or 43* or 44* or 45* or 46* or 47* or 48* or 49*):ti,ab,kw or body mass index near/1 (50* or 51* or 52* or 53* or 54* or 55*):ti,ab,kw (Word variations have been searched)
#6 #2 or #3 or #4 or #5
#7 MeSH descriptor: [Bariatric Surgery] explode all trees
#8 bariatric:ti,ab,kw (Word variations have been searched)
#9 gastric near/3 (bypass or by pass or band* or balloon*):ti,ab,kw or gastroplast* or gastrect*:ti,ab,kw or lipectom*:ti,ab,kw or jejunoileal near/3 (bypass or by pass):ti,ab,kw (Word variations have been searched)
#10 MeSH descriptor: [Diet Therapy] explode all trees
#11 MeSH descriptor: [Diet, Reducing] explode all trees
#12 MeSH descriptor: [Diet, Fat-Restricted] explode all trees
#13 MeSH descriptor: [Diet, Carbohydrate-Restricted] explode all trees
#14 MeSH descriptor: [Caloric Restriction] explode all trees
#15 diet near/3 (restrict* or reduc* or modif* or calorie*):ti,ab,kw or calori* near/3 (reduc* or restrict* or limit*):ti,ab,kw (Word variations have been searched)
#16 MeSH descriptor: [Anti-Obesity Agents] explode all trees
#17 appetite near/3 (reduc* or depress*):ti,ab,kw or orlistat or xenical:ti,ab,kw or sibutramin or arcalion or reductil or medaria or meridia:ti,ab,kw or rimonabant or acomplia or zimulti:ti,ab,kw (Word variations have been searched)
#18 MeSH descriptor: [Exercise] explode all trees
#19 MeSH descriptor: [Sports] explode all trees
#20 exercise or aerobic* or physical activ*:ti,ab,kw (Word variations have been searched)
#21 MeSH descriptor: [Weight Loss] explode all trees
#22 MeSH descriptor: [Weight Reduction Programs] explode all trees
#23 weight near/3 (reduc* or loss or lose) .:ti,ab,kw (Word variations have been searched)
#24 weight near/3 (reduc* or loss or lose) .:ti,ab,kw (Word variations have been searched)
#25 MeSH descriptor: [Behavior Therapy] explode all trees
#26 MeSH descriptor: [Health Education] explode all trees
#27 MeSH descriptor: [Counseling] explode all trees
#28 MeSH descriptor: [Life Style] explode all trees
#29 diet near/3 (educat* or counsel*):ti,ab,kw or modif* near/3 (behaviour or behavior or diet or eat or eating):ti,ab,kw or weight near/3 (educat* or counsel*):ti,ab,kw (Word variations have been searched)
#30 #7 or #8 or #9 or #10 or #11 or #12 or #13 or #14 or #15 or #16 or #17 or #18 or #19 or #20 or #21 or #22 or #23 or #24 or #25 or #26 or #27 or #28 or #29
#31 #6 and #30
#32 #1 or #31 Publication Year from 1990 to 2016, in Trials
#33 conference abstract (Word variations have been searched)
#34 #32 not #33
Appendix 2   PRISMA flow diagram

Database searches
MEDLINE/Embase   575
PsycINFO        120
CINAHL         372
SCI            970
CENTRAL       3011
Total           5048
After deduplication 3052

Excluded
N= 2011

Selected for full-text assessment
N= 141

Excluded studies
N=122
BMI <35 or unclear  N= 39
Not RCT         N= 19
Not relevant intervention/unclear  N= 6
Assessment <12 months  N= 15
No relevant outcome   N= 27
Secondary publication  N= 13
Not obtained         N= 3

Included studies
N= 19

Identified from other sources (previous auto-alert searches, previous review studies & qualitative searches)
N= 112

Total included studies
N= 131
Appendix 3 List of included studies


O'Neil PM, Miller-Kovach K, Tuerk PW, Becker LE, Wadden TA, Fujioka K, et al. Randomized Controlled Trial of a Nationally Available Weight Control Program Tailored for Adults with Type 2 Diabetes. *Obesity* 2016;**24**:2269-77.


Pekkarinen T, Kaukua J, Mustajoki P. Long-term weight maintenance after a 17-week weight loss intervention with or without a one-year maintenance program: A randomized controlled trial. *Journal of Obesity* 2015;Article 651460


Tsai AG, Wadden TA, Rogers MA, Day SC, Moore RH, Islam BJ. A Primary Care Intervention for Weight Loss: Results of a Randomized Controlled Pilot Study. *Obesity* 2010;18:1614-8.


