

Pelvic floor dysfunction: prevention and non- surgical management

[J] Weight loss interventions

NICE guideline NG210

*Evidence review underpinning recommendations 1.6.5 to 1.6.8
and a research recommendation in the NICE guideline*

December 2021

Final

*These evidence reviews were developed
by the National Guideline Alliance which is
a part of the Royal College of
Obstetricians and Gynaecologists*

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Weight loss interventions

Review question

What is the effectiveness of weight loss interventions for improving symptoms of pelvic floor dysfunction?

Introduction

Obesity is related to an increased prevalence of many chronic conditions. There is an assumption that being overweight or obese may increase pelvic floor dysfunction; a greater body weight results in an increased abdominal pressure, which may affect the pelvic floor detrimentally. However, evidence that weight loss can significantly improve symptoms associated with pelvic floor dysfunction has not been synthesised: this review aims to fill this gap in the evidence.

Summary of the protocol

See Table 1 for a summary of the Population, Intervention, Comparison and Outcome (PICO) characteristics of this review.

Table 1: Summary of the protocol (PICO table)

| | |
|---------------------|--|
| Population | Women and young women (aged 12 years and older) with symptoms associated with pelvic floor dysfunction |
| Intervention | Any intervention with the aim of weight loss will be considered, for example, but not exclusively: <ul style="list-style-type: none"> • Calorie restricted weight loss diets (to include liquid based diets) • Weight loss physical activity programmes • Combined diet and exercise plans for weight loss • Social support groups for weight loss • Behavioural weight loss programmes Combined interventions (those with a mixture of those listed above) will only be included if the primary aim is weight loss |
| Comparison | <ul style="list-style-type: none"> • Any of the above • No treatment/usual care |
| Outcomes | <p>Critical</p> <ul style="list-style-type: none"> • Subjective measure of change in the following symptoms: <ul style="list-style-type: none"> ○ urinary incontinence ○ emptying disorders of the bladder ○ faecal incontinence ○ emptying disorders of the bowel ○ pelvic organ prolapse ○ sexual dysfunction ○ chronic pelvic pain syndromes • Health related quality of life <p>Important</p> <ul style="list-style-type: none"> • Adherence to intervention • Anxiety and depression (only validated scales will be included) • Adverse events leading to withdrawal/discontinuation • Weight loss (as reported in the study: for example change in BMI, percentage weight loss or total weight loss) |

BMI: body mass index.

For details see the review protocol in appendix A.

Methods and process

This evidence review was developed using the methods and process described in [Developing NICE guidelines: the manual](#). Methods specific to this review question are described in the review protocol in appendix A and the methods document (supplementary document 1).

Declarations of interest were recorded according to [NICE's conflicts of interest policy](#).

Clinical evidence

Included studies

Eight publications (Breyer 2018, Myers 2012, Hagovska 2020 Subak 2005 Subak 2009, Gozukara 2014, West 2011, Wing 2011) from 4 randomised controlled trials (RCTs: Hagovska 2020 Subak 2005 Subak 2009; Gozukara 2014) were included for this review. Five of the publications were from the Subak 2009 RCT (Breyer 2018, Myers 2012, Subak 2009, West 2011, Wing 2011).

The included studies are summarised in Table 2.

Studies recruited women with urinary incontinence (Subak 2005, Subak 2009, Gozukara 2014) or overactive bladder (Hagovska 2020) and three studies used a combined low-calorie diet and exercise intervention to promote weight loss (Subak 2005, Subak 2009, Gozukara 2014). The intervention lasted for 6 months for Subak 2009 and Gozukara 2014 and 3 months for Hagovska 2020 and Subak 2005. Subak 2009 also included a 12 months of maintenance therapy for those in the intervention arm.

No evidence was identified for any other symptoms associated with pelvic floor dysfunction.

See the literature search strategy in appendix B and study selection flow chart in appendix C.

Excluded studies

Studies not included in this review are listed, and reasons for their exclusion are provided in appendix K.

Summary of studies included in the evidence review

Summaries of the studies that were included in this review are presented in Table 2.

Table 2: Summary of included studies.

| Study | Population | Intervention | Comparison | Outcomes |
|---|---|---|---|--|
| Breyer 2018 (secondary data analysis of Subak 2009) | See Subak 2009 | See Subak 2009 | See Subak 2009 | <ul style="list-style-type: none"> • Daytime frequency • Nocturia, • Urinary urgency • IPSS Score ≥8 |
| Gozukara 2014 RCT Turkey | N=378 women with urinary incontinence Age: Intervention | <u>6-month behavioural weight loss program and structured education program</u> | <u>Structured education program</u> 4 x 1hr sessions prior to randomisation with | <ul style="list-style-type: none"> • Change in UI symptoms • Change in PFD symptoms • Weight loss |

| Study | Population | Intervention | Comparison | Outcomes |
|--|--|---|--|--|
| | 44.1 (8.6) years; Control 43.8 (9.7) years; BMI: Intervention 32.7 (4.1)kg/m ² ; Control 32.3 (3.6) kg/m ² | Calorie and fat restricted diet (1200-1800 kcal/day). 6 x 1hr group sessions led by a nutrition, exercise and behaviour change internist. Encouraged to increase physical activity. | information on benefits of weight loss, physical activity and healthy eating on urinary complaints. | |
| Hagovska 2020 | N=93 women with overactive bladder Age: Intervention 26.7 (4.8); Control 26.9 (4.9) BMI: Intervention 25 (3); Control 25.1 (4.2) | <u>3-month program for reducing abdominal fat</u> Exercise was performed two times a week for 60-80 minutes and included aerobic training, stretching and strength training for reduction of abdominal fat, deep abdominal activation and superficial abdominal muscles. Eating habits were not changed. | <u>No treatment</u> Women did not undergo exercise, and did not change their everyday life activities or eating habits. | <ul style="list-style-type: none"> • Change in OAB symptoms • Weight loss |
| Myers 2012 (secondary data analysis of Subak 2009) | See Subak 2009 | See Subak 2009 | See Subak 2009 | <ul style="list-style-type: none"> • POP symptoms: cured • POP symptoms: improved bother or cured • POP symptoms: new report of symptom |
| Subak 2005 RCT USA | N=48 women with urinary incontinence Age (median, range): Intervention 50.5 (46-54) years; Control 57.5 (50-62) years BMI (median, range): Intervention 34 (32-40) kg/m ² ; Control 36 (32-38) kg/m ² | <u>3- month Weight loss program</u> 800 calorie or less liquid diet. Weekly group meetings. Encouraged to increase physical activity until exercising 60mins daily and taught standard cognitive and behavioural skills to modify eating and exercise habits. | <u>Wait-list control group</u> No treatment for 3 months then received weight loss program | <ul style="list-style-type: none"> • Change in UI symptoms • Health related quality of life • Weight loss |

| Study | Population | Intervention | Comparison | Outcomes |
|--------------------------|--|--|---|---|
| Subak 2009 RCT USA | N=338 women with urinary incontinence Age: Intervention 53 (11) years; Control 53 (10) years; BMI: Intervention 36 (6)kg/m ² ; Control 36 (5) kg/m ² | <u>6- month Weight loss program</u> Calorie and fat restricted diet (1200-1500 kcal/day). 1hr weekly group meeting. Vouchers for meal replacement shakes. Encouraged to increase physical activity aiming to achieve 200 minutes each week. <u>Followed by 12-month maintenance approach</u> Randomised to either a motivation-focused weight maintenance program or a skill-based maintenance program | <u>Educational sessions</u> 4 x 1hr sessions with general information about weight loss, physical activity and healthful eating habits | <ul style="list-style-type: none"> • Body weight (6 months) • Incontinence episodes/week (6 months; any, stress and urge) |
| West 2011 RCT USA | See Subak 2009 | 18 month follow-up after the initial 6 month intervention from Subak 2009 and then a 12 month maintenance programme for skill-based maintenance or motivational maintenance for the intervention arm | See Subak 2009 | <ul style="list-style-type: none"> • Weight change at 18 months (see Wing 2010 also) |
| Wing 2010 RCT USA | See Subak 2009 | 18 month follow-up after the initial 6 month intervention from Subak 2009 and then a 12 month maintenance programme for skill-based maintenance or motivational maintenance for the intervention arm | See Subak 2009 | <ul style="list-style-type: none"> • Body weight change at 18 months • Total, stress and urge UI episodes/week at 18 months • 24hr involuntary urine loss at 18 months |

CRADI: colorectal anal distress inventory; IIQ: incontinence impact questionnaire; IPSS: International Prostate Symptom Score; OAB-Q: Overactive Bladder Questionnaire; PFDI: Pelvic Floor Distress Inventory; POP: pelvic organ prolapse; POPDI: pelvic organ prolapse distress inventory; SF-36: short form of the quality of life questionnaire; UDI: urinary distress inventory; UI: urinary incontinence.

See the full evidence tables in appendix D and forest plots in appendix E.

Quality assessment of studies included in the evidence review

See the evidence profiles in appendix F.

Economic evidence

Included studies

A single economic search was undertaken for all topics included in the scope of this guideline but no economic studies were identified which were applicable to this review question. See the literature search strategy in appendix B and economic study selection flow chart in appendix G.

Excluded studies

Studies not included in this review are listed, and reasons for their exclusion are provided in appendix K.

Economic model

No economic modelling was undertaken for this review because the committee agreed that other topics were higher priorities for economic evaluation as any interventions recommended were unlikely to have a significant resource impact.

Brief summary of evidence

Weight loss interventions

- Very low to moderate quality evidence showed that weight loss interventions were effective in reducing weight when compared to control.
- Very low to moderate quality evidence showed that weight loss was associated with a clinically important reduction in urinary symptoms but not in POP symptoms.

Rationale and impact

The committee's discussion of the evidence

Interpreting the evidence

The outcomes that matter most

The committee agreed that improvement in symptoms of pelvic floor dysfunction and health related quality of life were the most critical outcomes for this review question. These outcomes are likely to have the most impact on the woman's life, and the interventions included specifically target the management of these symptoms. Anxiety and depression were considered important outcomes as many women report the psychological impact that pelvic floor dysfunction has on their lives. Other important outcomes were adherence to the intervention, adverse events and weight change as these outcomes were considered the most relevant to determining if, and potentially why the intervention was or was not successful.

The quality of the evidence

The quality of evidence for this review was assessed using GRADE and ranged from very low to moderate, with most of the evidence being of low quality. The main concerns with the quality of the evidence for the included outcomes was due to the risk of bias (where all

studies had methodological issues related to assignment, adherence, missing data and selective reporting) and imprecision.

No evidence was identified for the following outcomes: emptying disorders of the bladder, faecal incontinence, emptying disorders of the bowel, sexual dysfunction, chronic pelvic pain syndromes, adherence, anxiety and depression.

Benefits and harms

The evidence presented showed that weight loss interventions were effective in significantly reducing weight of women as compared to control. In one study body weight remained significantly lower in the intervention arm at 18 months when compared to the control arm.

The evidence also showed weight loss significantly reduced urinary incontinence episodes (including both stress and urge incontinence) as compared to control. The committee discussed based on experience that weight exacerbates urinary incontinence symptoms because of the intra-abdominal pressure it puts on the pelvic floor muscles and organs, so weight loss will be beneficial. All the evidence presented was based on women with a BMI greater than 30kg/m²; as such the recommendations are based on this cut-off. However, the committee discussed that very few women come into practice with a BMI that is lower than 25kg/m², and are considered overweight (which they noted was also the risk factor threshold identified in evidence review B). Therefore, these recommendations may also be relevant for these women, but since there was no direct evidence in this group they did not make a recommendation for women with a lower BMI than 30kg/m² even if this was inconsistent with the risk factor threshold identified in evidence review B. There was no evidence that weight loss was detrimental to their pelvic floor symptoms, and therefore women should be supported to lose weight.

All of the included interventions were based on “*supported weight loss programmes*”; therefore, the committee discussed how women should not simply be told to “lose weight” but should be referred to an appropriate programme. The committee acknowledged that healthcare professionals should follow the NICE obesity management guideline to help women lose weight and decided to cross reference a range of possible guidance related to this.

The evidence did not show any effect of weight loss on symptoms of pelvic organ prolapse (POP); the committee however, noted that this differs according to the stage of POP. The committee noted that this lack of effect was potentially related to different stages of POP. For women in late stage, weight loss is likely to have no effect, but weight loss could help considerably for women with early stage POP. They decided that a reduction of intra-abdominal pressure through weight loss may be beneficial in the early stages of pelvic organ prolapse because less weight would press on the pelvic organs and this could improve symptoms. Nonetheless, there was no evidence to support this, so the committee recommended it generally for POP (because of other beneficial health effects) rather than specifically by the stage of POP.

The committee noted that no evidence was identified that specifically studied women explicitly stated as having ‘pelvic floor dysfunction,’ or evidence for any other symptoms of pelvic floor dysfunction other than urinary incontinence, overactive bladder and pelvic organ prolapse. They therefore decided to make a research recommendation to encourage future research which would inform future guidance. If weight loss were an effective intervention, some women may be able to avoid surgery and other invasive interventions.

The committee agreed that although weight loss may help with some symptoms of pelvic floor disorder there is no reason to delay other effective management options while waiting for weight to be lost. This is because weight loss alone is unlikely to be effective for all women and it may be difficult to achieve quickly.

Cost effectiveness and resource use

The recommendation that certain women should be advised that weight loss can help with symptoms has negligible costs as that advice can occur in contacts and consultations that would be happening as part of the management of women's pelvic floor dysfunction, although a little more time may be necessary to discuss options. The guideline cross refers to other UK clinical and public health guidance for guidance and support for weight loss. Whilst no formal economic analysis was undertaken the committee considered their recommendations were likely to be cost-effective given their low cost of implementation and the potential of weight loss to prevent or mitigate symptoms of pelvic floor dysfunction.

Other factors the committee took into account

The committee agreed that weight loss recommendations should be in line with the NICE guidelines on obesity management: [Obesity: identification, assessment and management](#) [CG189], and [Weight management before, during and after pregnancy](#) [PH27]. The committee also referred to relevant NICE guidance for weight management interventions: [Weight management: lifestyle services for overweight or obese adults](#) [PH53] and [Weight management: lifestyle services for overweight or obese children and young people](#) [PH47].

Recommendations supported by this evidence review

This evidence review supports recommendations 1.6.5 to 1.6.8 and a research recommendation on weight loss to reduce symptoms of pelvic floor dysfunction in the NICE guideline.

References

Breyer 2018

Breyer, B. N., Creasman, J. M., Richter, H. E., Myers, D., Burgio, K. L., Wing, R. R., West, D. S., Kusek, J. W., Subak, L. L., Pride,, A Behavioral Weight Loss Program and Nonurinary Incontinence Lower Urinary Tract Symptoms in Overweight and Obese Women with Urinary Incontinence: A Secondary Data Analysis of PRIDE, *Journal of urology*, 199, 215-222, 2018

Gozukara 2014

Gozukara, Y. M., Akalan, G., Tok, E. C., Aytan, H., Ertunc, D., The improvement in pelvic floor symptoms with weight loss in obese women does not correlate with the changes in pelvic anatomy, *International Urogynecology Journal*, 25, 1219-25, 2014

Hagovska 2020

Hagovska, M., Svihra, J., Bukova, A., Drackova, D., Horbacz, A., Nagyova, I., Effect of an Exercise Programme for Reducing Abdominal Fat on Overactive Bladder Symptoms in Young Overweight Women, *Obstetrical and Gynecological Survey*, 75, 471-472, 2020

Myers 2012

Myers, D. L., Sung, V. W., Richter, H. E., Creasman, J., Subak, L. L., Prolapse symptoms in overweight and obese women before and after weight loss, *Female Pelvic Medicine & Reconstructive Surgery*, 18, 55-9, 2012

Subak 2005

Subak, L.L., Whitcomb, E., Shen, H., Saxton, J., Vittinghoff, E., Brown, J.S., Weight loss: A novel and effective treatment for urinary incontinence, *Journal of Urology*, 174, 190-195, 2005

Subak 2009

Subak, L. L., Wing, R., West, D. S., Franklin, F., Vittinghoff, E., Creasman, J. M., Richter, H. E., Myers, D., Burgio, K. L., Gorin, A. A., Macer, J., Kusek, J. W., Grady, D., Pride Investigators, Weight loss to treat urinary incontinence in overweight and obese women, *New England journal of medicine*, 360, 481-90, 2009

West 2011

West, D. S., Gorin, A. A., Subak, L. L., Foster, G., Bragg, C., Hecht, J., Schembri, M., Wing, R. R., Program to Reduce Incontinence by, Diet, Exercise Research, Group, A motivation-focused weight loss maintenance program is an effective alternative to a skill-based approach, *International journal of obesity*, 35, 259-69, 2011

Wing 2010

Wing, R. R., West, D. S., Grady, D., Creasman, J. M., Richter, H. E., Myers, D., Burgio, K. L., Franklin, F., Gorin, A. A., Vittinghoff, E., Macer, J., Kusek, J. W., Subak, L. L., Program to Reduce Incontinence by, Diet, Exercise, Group, Effect of weight loss on urinary incontinence in overweight and obese women: results at 12 and 18 months, *Journal of urology*, 184, 1005-10, 2010

Appendices

Appendix A – Review protocol

Review protocol for review question: What is the effectiveness of weight loss interventions for improving symptoms of pelvic floor dysfunction?

Table 3: Review protocol

| ID | Field | Content |
|----|------------------------------|---|
| 0. | PROSPERO registration number | Not registered |
| 1. | Review title | Weight loss interventions |
| 2. | Review question | What is the effectiveness of weight loss interventions for improving symptoms of pelvic floor dysfunction? |
| 3. | Objective | The objective of this review is to determine whether weight loss interventions can effectively improve symptoms (including urinary incontinence, pelvic organ prolapse, emptying disorders of the bladder, faecal incontinence, emptying disorders of the bowel, sexual dysfunction and chronic pelvic pain syndromes) associated with pelvic floor dysfunction. |
| 4. | Searches | <p>The following databases will be searched:</p> <ul style="list-style-type: none"> • Cochrane Database of Systematic Reviews (CDSR) • Cochrane Central Register of Controlled Trials (CENTRAL) • MEDLINE & Medline in Process • Embase • CINAHL or Emcare • PsycINFO <p>Searches will be restricted by:</p> <ul style="list-style-type: none"> • Date limit: 1980 onwards (see section 10 for justification) • English language • Human studies <p>Other searches:</p> <ul style="list-style-type: none"> • Inclusion lists of potentially relevant systematic reviews |

| ID | Field | Content |
|----|-----------------------------------|--|
| | | <p>The full search strategies for MEDLINE database will be published in the final review.</p> <p>For each search, the principal database search strategy is quality assured by a second information scientist using an adaptation of the PRESS 2015 Guideline Evidence-Based Checklist.</p> |
| 5. | Condition or domain being studied | <p>The following symptoms will be addressed as long as they are associated with pelvic floor dysfunction: urinary incontinence, emptying disorders of the bladder, faecal incontinence, emptying disorders of the bowel, pelvic organ prolapse, sexual dysfunction and chronic pelvic pain syndromes.</p> |
| 6. | Population | <p>Inclusion</p> <ul style="list-style-type: none"> • Women and young women (aged 12 years and older) with symptoms associated with pelvic floor dysfunction <p>Exclusion</p> <ul style="list-style-type: none"> • Studies which include women with urinary incontinence, emptying disorders of the bladder, faecal incontinence, emptying disorders of the bowel, pelvic organ prolapse, sexual dysfunction and chronic pelvic pain syndromes which are not due to pelvic floor dysfunction will be excluded. For example, women who have urinary incontinence due to a neurological condition or pelvic cancer will be excluded. During the screening stage, the reported inclusion/exclusion criteria of studies will be examined carefully. We do not anticipate studies on urinary incontinence, emptying disorders of the bladder or pelvic organ prolapse will explicitly state “<i>associated with pelvic floor dysfunction</i>” therefore this will be a pragmatic decision based on the description of the condition provided by the study authors. Some of these symptoms (for example urinary incontinence) are most often due to a failure in the pelvic floor and therefore unless the exclusion criteria states a different cause, these studies are likely to be included. However, for studies on faecal incontinence, emptying disorders of the bowel, sexual dysfunction and pelvic pain the causes are more numerous. As such for these symptoms unless the study specifically states “<i>associated with pelvic floor dysfunction</i>” they will be excluded. If any ambiguity exists, at least two reviewers will make the final decision if to include or exclude the study. • Men <p>Babies and children under 12 years old</p> |
| 7. | Intervention/Exposure/Test | <p>Any intervention with the aim of weight loss will be considered, for example, but not exclusively:</p> <ul style="list-style-type: none"> • Calorie restricted weight loss diets (to include liquid based diets) • Weight loss physical activity programmes • Combined diet and exercise plans for weight loss • Social support groups for weight loss • Behavioural weight loss programmes |

| ID | Field | Content |
|----|---|--|
| | | <p>Combined interventions (those with a mixture of those listed above) will only be included if the primary aim is weight loss</p> <p>All weight loss interventions included in this review, will fall under those described as “<i>lifestyle</i>”, “<i>behavioural</i>”, “<i>dietary</i>” or “<i>physical activity</i>” interventions for weight loss, as described in NICE guideline CG189: https://www.nice.org.uk/guidance/cg189/chapter/1-Recommendations</p> |
| 8. | Comparator/Reference standard/Confounding factors | <ul style="list-style-type: none"> • Any of the above • No treatment/usual care |
| 9. | Types of study to be included | <ul style="list-style-type: none"> • Systematic reviews of RCTs • RCTs <p>If there is no RCT evidence then other studies designs will be considered, namely</p> <ul style="list-style-type: none"> • Non-randomised controlled studies • Comparative prospective cohort studies <p>The decision to include non RCT study designs will be determined for each of the listed symptoms associated with pelvic floor dysfunction. For example, if we identify an RCT on urinary incontinence but not for pelvic organ prolapse, then we will continue our search for observational studies on pelvic organ prolapse but we will not search for further study designs for urinary incontinence.</p> <p>The decision to include non RCT study designs was made to ensure all relevant symptoms associated with pelvic floor dysfunctions are given equal consideration. Additionally, interventions may influence the various symptoms differently, and it is important this is considered. Within each symptom category (for example faecal incontinence), the committee has agreed a subset of symptoms that are specifically associated with pelvic floor dysfunction, as such each symptom only includes those sub-symptoms which occur as a result of pelvic floor dysfunction (rather than anybody with faecal incontinence). The committee agreed these subsets of symptoms by examining the population search strategy. Therefore, if lower level of evidence is identified it will only be relevant to symptoms that specifically result from pelvic floor dysfunction, rather than the entire population for which there could potentially have been a higher level of evidence.</p> <p>Potentially important confounders which should be considered include BMI, age, ethnicity, dietary factors and physical activity. Appropriate adjustment for these confounders within the included studies will be considered during the GRADE process.</p> |

| ID | Field | Content |
|-----|--------------------------------------|--|
| | | <p>Note: For further details, see the algorithm in appendix H, Developing NICE guidelines: the manual.</p> |
| 10. | Other exclusion criteria | <ul style="list-style-type: none"> • Weight loss surgery • Pharmaceutical weight loss drugs (for example orlistat) • We will only include studies where women are classified as overweight at study start (defined as BMI $\geq 25\text{kg/m}^2$), or where sub-group analysis for these women has been reported • Studies with a mixed population (that is women with symptoms such as urinary incontinence which are associated with pelvic floor dysfunction and women with symptoms that are not associated with pelvic floor dysfunction) will be excluded, unless subgroup analysis for those women with symptoms associated with pelvic floor dysfunction has been reported • Conference abstracts will be excluded because these do not typically provide sufficient information to fully assess risk of bias • Only articles published after 1980 will be included. This was agreed by the committee as this is the date that the condition “pelvic floor dysfunction” was recognised to include agreed terminology on symptoms. https://www.ncbi.nlm.nih.gov/pmc/articles/PMC2815805/ |
| 11. | Context | <p>Studies which explicitly demonstrate a change in outcomes for symptoms associated with pelvic floor dysfunction will be prioritised for decision making in regards to recommendations, and these recommendations will apply to those receiving care in any healthcare settings (for example community, primary, secondary care). However, the context of recommendations is likely broader than just the health care setting itself. Women who are not currently accessing services may benefit from the recommendations in order to make lifestyle changes which could improve symptoms they are experiencing.</p> <p>Specific recommendations for groups listed in the Equality Considerations section of the scope may be also be made as appropriate.</p> |
| 12. | Primary outcomes (critical outcomes) | <ul style="list-style-type: none"> • Subjective measure of change in the following symptoms: <ul style="list-style-type: none"> ○ urinary incontinence, ○ emptying disorders of the bladder, ○ faecal incontinence, ○ emptying disorders of the bowel, ○ pelvic organ prolapse, ○ sexual dysfunction ○ chronic pelvic pain syndromes |

| ID | Field | Content |
|-----|---|--|
| | | <ul style="list-style-type: none"> • Health related QOL <p>For primary outcomes listed, only validated tools will be included (for example: ICIQ-UI, ICIQ-VS, BFLUTS, KHQ, UDI, ISI, ePAQ, POPSS, PISQ, POPQ, FISQ, FIQL, GIQLI, PAC-QM, PAC –SYM, PDI, BPI)</p> |
| 13. | Secondary outcomes (important outcomes) | <ul style="list-style-type: none"> • Adherence to intervention • Anxiety and depression (only validated scales will be included) • Adverse events leading to withdrawal/discontinuation • Weight loss (as reported in the study for example: change in BMI, percentage weight loss or total weight loss) |
| 14. | Data extraction (selection and coding) | <p>All references identified by the searches and from other sources will be uploaded into STAR and de-duplicated.</p> <p>Titles and abstracts of the retrieved citations will be screened to identify studies that potentially meet the inclusion criteria outlined in the review protocol.</p> <p>Dual sifting will be performed on at least 10% of records; 90% agreement is required. Disagreements will be resolved via discussion between the two reviewers, and consultation with senior staff if necessary.</p> <p>Full versions of the selected studies will be obtained for assessment. Studies that fail to meet the inclusion criteria once the full version has been checked will be excluded at this stage. Each study excluded after checking the full version will be listed, along with the reason for its exclusion.</p> <p>A standardised form will be used to extract data from studies. One reviewer will extract relevant data into a standardised form, and this will be quality assessed by a senior reviewer. Information to be extracted from studies includes: study type, study dates, location of study, funding, inclusion and exclusion criteria, participant characteristics, and details of the intervention and comparator.</p> |
| 15. | Risk of bias (quality) assessment | <p>Quality assessment of individual studies will be performed using the following checklists</p> <ul style="list-style-type: none"> • ROBIS tool for systematic reviews • Cochrane RoB tool v.2 for RCTs and quasi-RCTs • Cochrane ROBINS-I tool for non-randomised (clinical) controlled trials and cohort studies <p>The quality assessment will be performed by one reviewer and this will be quality assessed by a senior reviewer.</p> |

| ID | Field | Content |
|-----|-----------------------------|--|
| 16. | Strategy for data synthesis | <p>Depending on the availability of the evidence, the findings will be summarised narratively or quantitatively.</p> <p><u>Data Synthesis</u></p> <p>Where possible, pair wise meta-analyses will be conducted using Cochrane Review Manager software. A fixed effect meta-analysis will be conducted and data will be presented as risk ratios for dichotomous outcomes. Peto odds ratio will be used for outcomes with zero events Mean differences or standardised mean differences will be calculated for continuous outcomes.</p> <p><u>Heterogeneity</u></p> <p>Heterogeneity in the effect estimates of the individual studies will be assessed using the I^2 statistic. I^2 values of greater than 50% and 80% will be considered as significant and very significant heterogeneity, respectively. In the presence of heterogeneity sub-group analysis will be conducted</p> <ol style="list-style-type: none"> 1. According to risk of bias of individual studies 2. According to socioeconomic status of population included 3. By ethnicity of included populations <p>Exact subgroup analysis may vary depending on differences identified within included studies. If heterogeneity cannot be explained through subgroup analysis, then a random effects model will be used for meta-analysis. If heterogeneity remains above 80% reviewers will consider if meta-analysis is appropriate given the characteristics of included</p> <p><u>Minimal important differences (MIDs)</u></p> <p>For outcomes where validated tools are included (for example ICIQ), then the published MIDs will be used.</p> <p>Where no published MID is available, default MIDs will be used:</p> <ul style="list-style-type: none"> • For risk ratios: 0.8 and 1.25. • For continuous outcomes: <ul style="list-style-type: none"> ○ For one study: the MID is calculated as +/-0.5 times the baseline SD of the control arm. ○ For two studies: the MID is calculated as +/-0.5 times the mean of the SDs of the control arms at baseline. If baseline SD is not available, then SD at follow up will be used. ○ For three or more studies (meta-analysed): the MID is calculated by ranking the studies in order of SD in the control arms. The MID is calculated as +/- 0.5 times median SD. ○ For studies that have been pooled using SMD (meta-analysed): +0.5 and -0.5 in the SMD scale are used as MID boundaries. |

| ID | Field | Content | | | | | | | | | | | | | | |
|-------------------------------------|----------------------------------|--|-------------------------------------|--------------|--------------------------|------------|--------------------------|------------|--------------------------|-------------|--------------------------|---------------|--------------------------|------------------|--------------------------|------------------------|
| | | <p><u>Validity</u></p> <p>The confidence in the findings across all available evidence will be evaluated for each outcome using an adaptation of the ‘Grading of Recommendations Assessment, Development and Evaluation (GRADE) toolbox’ developed by the international GRADE working group: http://www.gradeworkinggroup.org/</p> | | | | | | | | | | | | | | |
| 17. | Analysis of sub-groups | <p><u>Stratification</u></p> <p>All data will initially be pooled for overall analysis; however, if data is available, separate analysis will also be conducted on:</p> <ul style="list-style-type: none"> • Women who are pregnant • Women before and after gynaecological surgery • Women aged 65 or older • Women with physical disabilities • Women with cognitive impairment • According to those who do not identify themselves as women, but who have female pelvic organs • Women who successfully lost weight during the intervention and those that did not <p><i>Recommendations will apply to all those with pelvic floor dysfunction unless there is evidence of a difference in these stratified groups</i></p> | | | | | | | | | | | | | | |
| 18. | Type and method of review | <table border="1"> <tr> <td><input checked="" type="checkbox"/></td> <td>Intervention</td> </tr> <tr> <td><input type="checkbox"/></td> <td>Diagnostic</td> </tr> <tr> <td><input type="checkbox"/></td> <td>Prognostic</td> </tr> <tr> <td><input type="checkbox"/></td> <td>Qualitative</td> </tr> <tr> <td><input type="checkbox"/></td> <td>Epidemiologic</td> </tr> <tr> <td><input type="checkbox"/></td> <td>Service Delivery</td> </tr> <tr> <td><input type="checkbox"/></td> <td>Other (please specify)</td> </tr> </table> | <input checked="" type="checkbox"/> | Intervention | <input type="checkbox"/> | Diagnostic | <input type="checkbox"/> | Prognostic | <input type="checkbox"/> | Qualitative | <input type="checkbox"/> | Epidemiologic | <input type="checkbox"/> | Service Delivery | <input type="checkbox"/> | Other (please specify) |
| <input checked="" type="checkbox"/> | Intervention | | | | | | | | | | | | | | | |
| <input type="checkbox"/> | Diagnostic | | | | | | | | | | | | | | | |
| <input type="checkbox"/> | Prognostic | | | | | | | | | | | | | | | |
| <input type="checkbox"/> | Qualitative | | | | | | | | | | | | | | | |
| <input type="checkbox"/> | Epidemiologic | | | | | | | | | | | | | | | |
| <input type="checkbox"/> | Service Delivery | | | | | | | | | | | | | | | |
| <input type="checkbox"/> | Other (please specify) | | | | | | | | | | | | | | | |
| 19. | Language | English | | | | | | | | | | | | | | |
| 20. | Country | England | | | | | | | | | | | | | | |
| 21. | Anticipated or actual start date | TBC | | | | | | | | | | | | | | |
| 22. | Anticipated completion date | August 2021 | | | | | | | | | | | | | | |

| ID | Field | Content | | |
|-----|--|--|----------------|------------------|
| 23. | Stage of review at time of this submission | Review stage | Started | Completed |
| | | Preliminary searches | x | x |
| | | Piloting of the study selection process | x | x |
| | | Formal screening of search results against eligibility criteria | x | x |
| | | Data extraction | x | x |
| | | Risk of bias (quality) assessment | x | x |
| | | Data analysis | x | x |
| 24. | Named contact | <p>5a. Named contact National Guideline Alliance</p> <p>5b Named contact e-mail PreventionofPOP@nice.org.uk</p> <p>5e Organisational affiliation of the review National Institute for Health and Care Excellence (NICE) and the National Guideline Alliance</p> | | |
| 25. | Review team members | <ul style="list-style-type: none"> • NGA technical team | | |
| 26. | Funding sources/sponsor | <p>This systematic review is being completed by the National Guideline Alliance, which is funded by NICE and hosted by the Royal College of Obstetricians and Gynaecologists. NICE funds the National Guideline Alliance to develop guidelines for those working in the NHS, public health, and social care in England.</p> | | |
| 27. | Conflicts of interest | <p>All guideline committee members and anyone who has direct input into NICE guidelines (including the evidence review team and expert witnesses) must declare any potential conflicts of interest in line with NICE's code of practice for declaring and dealing with conflicts of interest. Any relevant interests, or changes to interests, will also be declared publicly at the start of each guideline committee meeting. Before each meeting, any potential conflicts of interest will be considered by the guideline committee Chair and a senior member of the development team. Any decisions to exclude a person from all or part of a meeting will be documented. Any changes to a member's declaration of interests will be recorded in the minutes of the meeting. Declarations of interests will be published with the final guideline.</p> | | |

| ID | Field | Content |
|------|--|--|
| 28. | Collaborators | Development of this systematic review will be overseen by an advisory committee who will use the review to inform the development of evidence-based recommendations in line with section 3 of Developing NICE guidelines: the manual . Members of the guideline committee are available on the NICE website: https://www.nice.org.uk/guidance/indevelopment/gid-ng10123/ |
| 29. | Other registration details | |
| 30. | Reference/URL for published protocol | [Give the citation and link for the published protocol, if there is one.] |
| 31. | Dissemination plans | NICE may use a range of different methods to raise awareness of the guideline. These include standard approaches such as: notifying registered stakeholders of publication publicising the guideline through NICE's newsletter and alerts issuing a press release or briefing as appropriate, posting news articles on the NICE website, using social media channels, and publicising the guideline within NICE. |
| 32. | Keywords | Weight loss Pelvic floor dysfunction |
| 33. | Details of existing review of same topic by same authors | Not applicable |
| 34. | Current review status | <input checked="" type="checkbox"/> Ongoing |
| | | <input type="checkbox"/> Completed but not published |
| | | <input type="checkbox"/> Completed and published |
| | | <input type="checkbox"/> Completed, published and being updated |
| | | <input type="checkbox"/> Discontinued |
| 35.. | Additional information | |
| 36. | Details of final publication | www.nice.org.uk |

BFLUTS: Bristol Female Lower Urinary Tract Symptoms Questionnaire; BPI: Brief pain inventory; BMI: body mass index; CDSR: Cochrane Database of Systematic Reviews; CENTRAL: Cochrane Central Register of Controlled Trials; DARE: Database of Abstracts of Reviews of Effects; ePAQ: Electronic personal health questionnaire; FIQL: Faecal incontinence quality of life scale; FISI: Faecal incontinence severity index; GIQLI: Gastrointestinal quality of life index; GRADE: Grading of Recommendations Assessment, Development and Evaluation; HTA: Health Technology Assessment; ICIQ-UI: International Consultation on Incontinence Questionnaire- Urinary incontinence; ICIQ-VA: International Consultation on Incontinence questionnaire – vaginal symptoms; ISI: Incontinence symptom index; KHQ: Kings health questionnaire; MID: minimally important difference; NGA: National Guideline Alliance; NHS: National health service; NICE: National Institute for Health and Care Excellence; PAC-QL: patient assessment of constipation - quality of life; PAC-SYM: Patient assessment of constipation symptoms; PDI: Pain disability index; PISQ: Pelvic organ prolapse/urinary incontinence sexual questionnaire; POPQ: Pelvic organ prolapse quantification system; POP-SS: Pelvic organ prolapse symptom score; QOL: quality of life; RCT: randomised controlled trial; RoB: risk of bias; SD: standard deviation; UDI: Urinary distress index

Appendix B – Literature search strategies

Literature search strategies for review question: What is the effectiveness of weight loss interventions for improving symptoms of pelvic floor dysfunction?

Clinical Search

Database(s): Medline & Embase (Multifile) – OVID interface

Embase Classic+Embase 1947 to 2021 February 01; Ovid MEDLINE(R) and Epub Ahead of Print, In-Process & Other Non-Indexed Citations and Daily 1946 to February 01, 2021
Date of last search: 2 February 2021

Multifile database codes: emczd = Embase Classic+Embase; ppez= MEDLINE(R) and Epub Ahead of Print, In-Process & Other Non-Indexed Citations and Daily

| # | Searches |
|----|---|
| 1 | exp Urinary Incontinence/ use ppez |
| 2 | Urinary Bladder, Overactive/ use ppez |
| 3 | Nocturia/ use ppez |
| 4 | exp Enuresis/ use ppez |
| 5 | exp urine incontinence/ use emczd |
| 6 | overactive bladder/ use emczd |
| 7 | bladder instability/ use emczd |
| 8 | nocturia/ use emczd |
| 9 | exp enuresis/ use emczd |
| 10 | ((stress\$ or mix\$ or urg\$ or urin\$) adj5 incontinen\$).tw. |
| 11 | (bladder\$ adj5 (overactiv\$ or over activ\$ or over-activ\$ or instabilit\$ or hyper-reflex\$ or hyperreflex\$ or hyper reflex\$ or incontinen\$)).tw. |
| 12 | (detrusor\$ adj5 (overactiv\$ or over activ\$ or over-activ\$ or instabilit\$ or hyper-reflex\$ or hyperreflex\$ or hyper reflex\$)).tw. |
| 13 | ((urgency adj2 frequency) or (frequency adj2 urgency)).tw. |
| 14 | ((urin\$ or bladder\$) adj2 (urg\$ or frequen\$)).tw. |
| 15 | (nocturia\$ or enuresis\$).tw. |
| 16 | (SUI or OAB).tw. |
| 17 | or/1-16 |
| 18 | exp Pelvic Organ Prolapse/ use ppez |
| 19 | exp pelvic organ prolapse/ use emczd |
| 20 | Rectocele/ use ppez |
| 21 | rectocele/ use emczd |
| 22 | (pelvic\$ adj3 organ\$ adj3 prolaps\$).tw. |
| 23 | (urinary adj3 bladder adj3 prolaps\$).tw. |
| 24 | ((vagin\$ or urogenital\$ or genit\$ or uter\$ or viscer\$ or anterior\$ or posterior\$ or apical or pelvi\$ or vault\$ or urethr\$ or bladder\$ or cervi\$ or rectal or rectum) adj3 prolaps\$).tw. |
| 25 | (splanchnoptos\$ or visceroptos\$).tw. |
| 26 | (hernia\$ adj3 (pelvi\$ or vagin\$ or urogenital\$ or uter\$ or bladder\$ or urethr\$ or viscer\$)).tw. |
| 27 | (urethroc?ele\$ or enteroc?ele\$ or sigmoidoc?ele\$ or proctoc?ele\$ or rectoc?ele\$ or cystoc?ele\$ or rectoenteroc?ele\$ or cystourethroc?ele\$).tw. |
| 28 | or/18-27 |
| 29 | Fecal Incontinence/ use ppez |
| 30 | feces incontinence/ use emczd |
| 31 | ((faecal or fecal or faeces or feces or fecally or faecally or anal or anally or stool or stools or bowel or double or defecat\$ or defaecat\$) adj5 (incontinence or incontinent or urge\$ or leak or leaking or leakage or soiling or seeping or seepage or impacted or impaction)).tw. |
| 32 | or/29-31 |
| 33 | Urinary Retention/ use ppez |
| 34 | Dysuria/ use ppez |
| 35 | urine retention/ use emczd |
| 36 | dysuria/ use emczd |
| 37 | (urin\$ adj3 (retention\$ or retain\$)).tw. |
| 38 | dysuria\$.tw. |
| 39 | (voiding adj (disorder\$ or dysfunction\$ or problem\$)).tw. |
| 40 | (empty\$ adj disorder\$ adj3 (bowel\$ or bladder\$ or vesical\$ or stool\$)).tw. |
| 41 | ((urogeni\$ or anorec\$ or ano-rec\$ or ano rec\$) adj3 dysfunction\$).tw. |
| 42 | defecation disorder/ use emczd |
| 43 | *Constipation/ use ppez |
| 44 | *constipation/ use emczd |
| 45 | Fecal Impaction/ use ppez |

| # | Searches |
|-----|---|
| 46 | Feces Impaction/ use emczd |
| 47 | constipat\$.ti. |
| 48 | ((difficult\$ or delay\$ or irregular\$ or infrequen\$ or pain\$) adj3 (defecat\$ or defaecat\$ or stool\$ or faecal or fecal or faeces or feces or fecally or faecally or bowel movement\$)).tw. |
| 49 | coprostasis.tw. |
| 50 | (obstruct\$ adj3 (defecat\$ or defaecat\$)).tw. |
| 51 | ((defecat\$ or defaecat\$ or evacuat\$) adj3 (disorder\$ or dysfunction\$)).tw. |
| 52 | outlet\$ dysfunction\$ constipa\$.tw. |
| 53 | (dys?ynerg\$ adj (defecat\$ or defaecat\$)).tw. |
| 54 | (pelvi\$ adj3 dyskines\$).tw. |
| 55 | pelvi\$ outlet\$ obstruct\$.tw. |
| 56 | anismus\$.tw. |
| 57 | puborectal\$ contract\$.tw. |
| 58 | ((rectal or rectum) adj3 urge\$).tw. |
| 59 | or/33-58 |
| 60 | female sexual dysfunction/ use emczd |
| 61 | (female adj sex\$ adj (dysfunct\$ or satisf\$ or problem\$ or symptom\$ or arous\$ or activit\$ or disorder\$)).tw. |
| 62 | Dyspareunia/ use ppez |
| 63 | dyspareunia/ use emczd |
| 64 | (sex\$ adj3 pain\$).tw. |
| 65 | (dyspareun\$ or anodyspareun\$).tw. |
| 66 | (obstruct\$ adj3 intercourse).tw. |
| 67 | (vagin\$ adj3 laxity\$).tw. |
| 68 | (vagin\$ adj wind).tw. |
| 69 | orgasm disorder/ use emczd |
| 70 | (female adj orgasm\$ adj (disorder\$ or deficienc\$ or dysfunction\$ or problem\$)).tw. |
| 71 | anorgasm\$.tw. |
| 72 | Vaginismus/ use ppez |
| 73 | vaginism/ use emczd |
| 74 | vaginismus\$.tw. |
| 75 | (vagin\$ adj penetrat\$ adj disorder\$).tw. |
| 76 | Vulvodynia/ use ppez |
| 77 | vulvodynia/ use emczd |
| 78 | vulvodynia\$.tw. |
| 79 | (vagin\$ adj dry\$).tw. |
| 80 | hypoactive sexual desire disorder/ use emczd |
| 81 | hypoactiv\$ sex\$ desire\$.tw. |
| 82 | sexual arousal disorder/ use emczd |
| 83 | (sex\$ adj arous\$ adj disorder\$).tw. |
| 84 | (genitourin\$ adj syndrom\$ adj5 menopaus\$).tw. |
| 85 | or/60-84 |
| 86 | Pelvic Pain/ use ppez |
| 87 | pelvic pain/ use emczd |
| 88 | ((pelvi\$ or lumbopelvi\$ or lumbo-pelvi\$ or genito-pelvi\$ or genitopelvi\$) adj3 pain\$).tw. |
| 89 | (pubi\$ adj3 (pain\$ or dysfunction\$)).tw. |
| 90 | (pudend\$ adj3 neuralg\$).tw. |
| 91 | proctalgia\$.tw. |
| 92 | (tension\$ adj myalgia\$).tw. |
| 93 | or/86-92 |
| 94 | Pelvic Floor/ use ppez |
| 95 | Pelvic Floor Disorders/ use ppez |
| 96 | pelvis floor/ use emczd |
| 97 | pelvic floor disorder/ use emczd |
| 98 | (pelvi\$ adj (floor\$ or diaphragm\$) adj3 (dysfunction\$ or disorder\$ or fail\$ or impair\$ or incompeten\$ or insufficien\$ or dyssynerg\$ or symptom\$ or laxity or change\$ or care\$ or health\$ or wellbeing\$ or well-being\$ or prevent\$ or rehabilitat\$)).tw. |
| 99 | (pelvi\$ adj (dysfunction\$ or disorder\$ or fail\$ or impair\$ or incompeten\$ or insufficien\$ or dyssynerg\$ or symptom\$ or laxity or care\$ or health\$ or wellbeing\$ or well-being\$ or prevent\$ or rehabilitat\$)).tw. |
| 100 | or/94-99 |
| 101 | Weight Loss/ use ppez |
| 102 | Weight Reduction Programs/ use ppez |
| 103 | weight reduction/ use emczd |
| 104 | *body weight loss/ use emczd |
| 105 | body weight control/ use emczd |
| 106 | body weight change/ use emczd |
| 107 | weight loss program/ use emczd |
| 108 | (weight adj2 (los\$ or reduc\$) adj3 (modif\$ or therap\$ or intervention\$ or strateg\$ or program\$ or management or scheme\$ or group\$ or pathway)).tw. |
| 109 | (weight adj management).tw. |
| 110 | ((calori\$ or hypocalori\$) adj2 (restrict\$ or diet\$)).tw. |
| 111 | or/101-110 |

| # | Searches |
|-----|---|
| 112 | (17 or 28 or 32 or 59 or 85 or 93 or 100) and 111 |
| 113 | limit 112 to english language |
| 114 | limit 113 to yr="1980 -Current" [General Exclusions filter applied] |

Database(s): Cochrane Library – Wiley interface

Cochrane Database of Systematic Reviews, Issue 2 of 12, February 2021; Cochrane Central Register of Controlled Trials, Issue 2 of 12, February 2021

Date of last search: 2 February 2021

| # | Search |
|-----|---|
| #1 | MeSH descriptor: [Urinary Incontinence] explode all trees |
| #2 | MeSH descriptor: [Urinary Bladder, Overactive] this term only |
| #3 | MeSH descriptor: [Nocturia] this term only |
| #4 | MeSH descriptor: [Enuresis] explode all trees |
| #5 | ((stress* or mix* or urg* or urin*) NEAR/5 incontinen*)):ti,ab,kw |
| #6 | ((bladder* NEAR/5 (overactiv* or over activ* or over-activ* or instabilit* or hyper-reflex* or hyperreflex* or hyper reflex* or incontinen*)):ti,ab,kw |
| #7 | ((detrusor* NEAR/5 (overactiv* or over activ* or over-activ* or instabilit* or hyper-reflex* or hyperreflex* or hyper reflex*)):ti,ab,kw |
| #8 | ((urgency NEAR/2 frequency) or (frequency NEAR/2 urgency)):ti,ab,kw |
| #9 | ((urin* or bladder*) NEAR/2 (urg* or frequen*)):ti,ab,kw |
| #10 | ((nocturia* or enuresis*)):ti,ab,kw |
| #11 | ((SUI or OAB)):ti,ab,kw |
| #12 | #1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7 OR #8 OR #9 OR #10 OR #11 |
| #13 | MeSH descriptor: [Pelvic Organ Prolapse] explode all trees |
| #14 | MeSH descriptor: [Rectocele] this term only |
| #15 | ((pelvic* NEAR/3 organ* NEAR/3 prolaps*)):ti,ab,kw |
| #16 | ((urinary NEAR/3 bladder NEAR/3 prolaps*)):ti,ab,kw |
| #17 | ((vagin* or urogenital* or genit* or uter* or viscer* or anterior* or posterior* or apical or pelvi* or vault* or urethr* or bladder* or cervi* or rectal or rectum) NEAR/3 prolaps*)):ti,ab,kw |
| #18 | ((splanchnoptos* or visceroptos*)):ti,ab,kw |
| #19 | ((hernia* NEAR/3 (pelvi* or vagin* or urogenital* or uter* or bladder* or urethr* or viscer*)):ti,ab,kw |
| #20 | ((urethroc?ele* or enteroc?ele* or sigmoidoc?ele* or proctoc?ele* or rectoc?ele* or cystoc?ele* or rectoenteroc?ele* or cystourethroc?ele*)):ti,ab,kw |
| #21 | #13 OR #14 OR #15 OR #16 OR #17 OR #18 OR #19 OR #20 |
| #22 | MeSH descriptor: [Fecal Incontinence] this term only |
| #23 | ((faecal or fecal or faeces or feces or fecally or faecally or anal or anally or stool or stools or bowel or double or defecat* or defaecat*) NEAR/5 (incontinence or incontinent or urge* or leak or leaking or leakage or soiling or seeping or seepage or impacted or impaction)):ti,ab,kw |
| #24 | #22 OR #23 |
| #25 | MeSH descriptor: [Urinary Retention] this term only |
| #26 | MeSH descriptor: [Dysuria] this term only |
| #27 | ((urin* NEAR/3 (retention* or retain*)):ti,ab,kw |
| #28 | (dysuria*):ti,ab,kw |
| #29 | ((voiding NEXT (disorder* or dysfunction* or problem*)):ti,ab,kw |
| #30 | ((empty* NEXT disorder* NEAR/3 (bowel* or bladder* or vesical* or stool*)):ti,ab,kw |
| #31 | ((urogeni* or anorec* or ano-rec* or ano rec*) NEAR/3 dysfunction*)):ti,ab,kw |
| #32 | MeSH descriptor: [Constipation] this term only |
| #33 | MeSH descriptor: [Fecal Impaction] this term only |
| #34 | (constipat*):ti |
| #35 | ((difficult* or delay* or irregular* or infrequen* or pain*) NEAR/3 (defecat* or defaecat* or stool* or faecal or fecal or faeces or feces or fecally or faecally or bowel movement*)):ti,ab,kw |
| #36 | (coprostasis):ti,ab,kw |
| #37 | ((obstruct* NEAR/3 (defecat* or defaecat*)):ti,ab,kw |
| #38 | ((defecat* or defaecat* or evacuat*) NEAR/3 (disorder* or dysfunction*)):ti,ab,kw |
| #39 | (outlet* dysfunction* constipa*):ti,ab,kw |
| #40 | ((dys?ynerg* NEXT (defecat* or defaecat*)):ti,ab,kw |
| #41 | ((pelvi* NEAR/3 dyskines*)):ti,ab,kw |
| #42 | (pelvi* outlet* obstruct*):ti,ab,kw |
| #43 | (anismus*):ti,ab,kw |
| #44 | (puborectal* contract*):ti,ab,kw |
| #45 | ((rectal or rectum) NEAR/3 urge*)):ti,ab,kw |
| #46 | #25 OR #26 OR #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 #40 OR #41 OR #42 OR #43 OR #44 OR #45 |
| #47 | ((female NEXT sex* NEXT (dysfunct* or satisf* or problem* or symptom* or arous* or activit* or disorder*)):ti,ab,kw |
| #48 | MeSH descriptor: [Dyspareunia] this term only |
| #49 | ((sex* NEAR/3 pain*)):ti,ab,kw |
| #50 | (dyspareun* or anodyspareun*):ti,ab,kw |
| #51 | ((obstruct* NEAR/3 intercourse)):ti,ab,kw |
| #52 | ((vagin* NEAR/3 laxity*)):ti,ab,kw |

| # | Search |
|-----|---|
| #53 | ((vagin* NEXT wind)):ti,ab,kw |
| #54 | ((female NEXT orgasm* NEXT (disorder* or deficienc* or dysfunction* or problem*)):ti,ab,kw |
| #55 | (anorgasm*):ti,ab,kw |
| #56 | MeSH descriptor: [Vaginismus] this term only |
| #57 | (vaginismus*):ti,ab,kw |
| #58 | ((vagin* NEXT penetrat* NEXT disorder*)):ti,ab,kw |
| #59 | MeSH descriptor: [Vulvodynia] this term only |
| #60 | (vulvodynia*):ti,ab,kw |
| #61 | ((vagin* NEXT dry*)):ti,ab,kw |
| #62 | (hypoactiv* NEXT sex* NEXT desire*):ti,ab,kw |
| #63 | ((sex* NEXT arous* NEXT disorder*)):ti,ab,kw |
| #64 | ((genitourin* NEXT syndrom* NEAR/5 menopaus*)):ti,ab,kw |
| #65 | #47 OR #48 OR #49 OR #50 OR #51 OR #52 OR #53 OR #54 OR #55 OR #56 OR #57 OR #58 OR #59 OR #60 OR #61 OR #62 OR #63 OR #64 |
| #66 | MeSH descriptor: [Pelvic Pain] this term only |
| #67 | ((pelvi* or lumbopelvi* or lumbo-pelvi* or genito-pelvi* or genitopelvi*) NEAR/3 pain*)):ti,ab,kw |
| #68 | ((pubi* NEAR/3 (pain* or dysfunction*)):ti,ab,kw |
| #69 | ((pudend* NEAR/3 neuralg*)):ti,ab,kw |
| #70 | (proctalgia*):ti,ab,kw |
| #71 | ((tension* NEXT myalgia*)):ti,ab,kw |
| #72 | #66 OR #67 OR #68 OR #69 OR #70 OR #71 |
| #73 | MeSH descriptor: [Pelvic Floor] this term only |
| #74 | MeSH descriptor: [Pelvic Floor Disorders] this term only |
| #75 | ((pelvi* adj (floor* or diaphragm*) NEAR/3 (dysfunction* or disorder* or fail* or impair* or incompeten* or insufficien* or dyssynerg* or symptom* or laxity or change* or care* or health* or wellbeing* or well-being* or prevent* or rehabilitat*)):ti,ab,kw |
| #76 | ((pelvi* NEXT (dysfunction* or disorder* or fail* or impair* or incompeten* or insufficien* or dyssynerg* or symptom* or laxity or care* or health* or wellbeing* or well-being* or prevent* or rehabilitat*)):ti,ab,kw |
| #77 | #73 OR #74 OR #75 OR #76 |
| #78 | MeSH descriptor: [Weight Loss] this term only |
| #79 | MeSH descriptor: [Weight Reduction Programs] this term only |
| #80 | ((weight NEAR/2 (los* or reduc*) NEAR/3 (modif* or therap* or intervention* or strateg* or program* or management or scheme* or group* or pathway)):ti,ab,kw |
| #81 | ((weight NEXT management)):ti,ab,kw |
| #82 | ((calori* or hypocalori*) NEAR/2 (restrict* or diet*)):ti,ab,kw |
| #83 | #78 OR #79 OR #80 OR #81 OR #82 |
| #84 | (#12 OR #21 OR #24 OR #46 OR #65 OR #72 OR #77) AND #83 |

Database(s): Database of Abstracts of Reviews of Effects (DARE); HTA Database – CRD interface

Date of last search: 2 February 2021

| Line | Search |
|------|--|
| 1 | MeSH DESCRIPTOR Urinary Incontinence EXPLODE ALL TREES IN DARE,HTA |
| 2 | MeSH DESCRIPTOR Urinary Bladder, Overactive IN DARE,HTA |
| 3 | MeSH DESCRIPTOR Nocturia IN DARE,HTA |
| 4 | MeSH DESCRIPTOR Enuresis EXPLODE ALL TREES IN DARE,HTA |
| 5 | ((stress* or mix* or urg* or urin*) NEAR5 incontinen*) IN DARE, HTA |
| 6 | ((bladder* NEAR5 (overactiv* or over activ* or over-activ* or instabilit* or hyper-reflex* or hyperreflex* or hyper reflex* or incontinen*)) IN DARE, HTA |
| 7 | ((detrusor* NEAR5 (overactiv* or over activ* or over-activ* or instabilit* or hyper-reflex* or hyperreflex* or hyper reflex*)) IN DARE, HTA |
| 8 | ((urgency NEAR2 frequency) or (frequency NEAR2 urgency)) IN DARE, HTA |
| 9 | ((urin* or bladder*) NEAR2 (urg* or frequen*)) IN DARE, HTA |
| 10 | ((nocturia* or enuresis*)) IN DARE, HTA |
| 11 | ((SUI or OAB)) IN DARE, HTA |
| 12 | MeSH DESCRIPTOR Pelvic Organ Prolapse EXPLODE ALL TREES IN DARE,HTA |
| 13 | MeSH DESCRIPTOR Rectocele IN DARE,HTA |
| 14 | ((pelvic* NEAR3 organ* NEAR3 prolaps*)) IN DARE, HTA |
| 15 | ((urinary NEAR3 bladder NEAR3 prolaps*)) IN DARE, HTA |
| 16 | ((vagin* or urogenital* or genit* or uter* or viscer* or anterior* or posterior* or apical or pelvi* or vault* or urethr* or bladder* or cervi* or rectal or rectum) NEAR3 prolaps*)) IN DARE, HTA |
| 17 | ((splanchnoptos* or visceroptos*)) IN DARE, HTA |
| 18 | ((hernia* NEAR3 (pelvi* or vagin* or urogenital* or uter* or bladder* or urethr* or viscer*)) IN DARE, HTA |
| 19 | ((urethroc?ele* or enteroc?ele* or sigmoidoc?ele* or proctoc?ele* or rectoc?ele* or cystoc?ele* or rectoenteroc?ele* or cystourethroc?ele*)) IN DARE, HTA |
| 20 | MeSH DESCRIPTOR Fecal Incontinence IN DARE,HTA |
| 21 | ((faecal or fecal or faeces or feces or fecally or faecally or anal or anally or stool or stools or bowel or double or defecat* or defaecat*) NEAR5 (incontinence or incontinent or urge* or leak or leaking or leakage or soiling or seeping or seepage or impacted or impaction)) IN DARE, HTA |
| 22 | MeSH DESCRIPTOR Urinary Retention IN DARE,HTA |

| Line | Search |
|------|--|
| 23 | ((urin* NEAR3 (retention* or retain*)) IN DARE, HTA |
| 24 | (dysuria*) IN DARE, HTA |
| 25 | ((voiding NEXT (disorder* or dysfunction* or problem*)) IN DARE, HTA |
| 26 | ((empty* NEXT disorder* NEAR3 (bowel* or bladder* or vesical* or stool*)) IN DARE, HTA |
| 27 | ((urogeni* or anorec* or ano-rec* or ano rec*) NEAR3 dysfunction*) IN DARE, HTA |
| 28 | MeSH DESCRIPTOR Constipation IN DARE,HTA |
| 29 | MeSH DESCRIPTOR Fecal Impaction IN DARE,HTA |
| 30 | (constipat*):TI IN DARE, HTA |
| 31 | ((difficult* or delay* or irregular* or infrequen* or pain*) NEAR3 (defecat* or defaecat* or stool* or faecal or fecal or faeces or feces or fecally or faecally or bowel movement*)) IN DARE, HTA |
| 32 | (coprosthiasis) IN DARE, HTA |
| 33 | ((obstruct* NEAR3 (defecat* or defaecat*)) IN DARE, HTA |
| 34 | ((defecat* or defaecat* or evacuat*) NEAR3 (disorder* or dysfunction*)) IN DARE, HTA |
| 35 | (outlet* NEXT dysfunction* NEXT constipa*) IN DARE, HTA |
| 36 | ((dys?ynerg* NEXT (defecat* or defaecat*)) IN DARE, HTA |
| 37 | ((pelvi* NEAR3 dyskines*)) IN DARE, HTA |
| 38 | (pelvi* NEXT outlet* NEXT obstruct*) IN DARE, HTA |
| 39 | (anismus*) IN DARE, HTA |
| 40 | (puborectal* contract*) IN DARE, HTA |
| 41 | ((rectal or rectum) NEAR3 urge*) IN DARE, HTA |
| 42 | ((female NEXT sex* NEXT (dysfunct* or satisf* or problem* or symptom* or arous* or activit* or disorder*)) IN DARE, HTA |
| 43 | MeSH DESCRIPTOR Dyspareunia IN DARE,HTA |
| 44 | ((sex* NEAR3 pain*)) IN DARE, HTA |
| 45 | ((dyspareun* or anodyspareun*) IN DARE, HTA |
| 46 | ((obstruct* NEAR3 intercourse)) IN DARE, HTA |
| 47 | ((vagin* NEAR3 laxity*)) IN DARE, HTA |
| 48 | ((vagin* NEXT wind)) IN DARE, HTA |
| 49 | ((female NEXT orgasm* adj (disorder* or deficienc* or dysfunction* or problem*)) IN DARE, HTA |
| 50 | (anorgasm*) IN DARE, HTA |
| 51 | MeSH DESCRIPTOR Vaginismus IN DARE,HTA |
| 52 | (vaginismus*) IN DARE, HTA |
| 53 | ((vagin* NEXT penetrat* NEXT disorder*)) IN DARE, HTA |
| 54 | MeSH DESCRIPTOR vulvodynia IN DARE,HTA |
| 55 | (vulvodynia*) IN DARE, HTA |
| 56 | ((vagin* NEXT dry*)) IN DARE, HTA |
| 57 | (hypoactiv* NEXT sex* NEXT desire*) IN DARE, HTA |
| 58 | ((sex* NEXT arous* NEXT disorder*)) IN DARE, HTA |
| 59 | ((genitourin* NEXT syndrom* NEAR5 menopaus*)) IN DARE, HTA |
| 60 | MeSH DESCRIPTOR Pelvic Pain IN DARE,HTA |
| 61 | ((pelvi* or lumbopelvi* or lumbo-pelvi* or genito-pelvi* or genitopelvi*) NEAR3 pain*) IN DARE, HTA |
| 62 | ((pubi* NEAR3 (pain* or dysfunction*)) IN DARE, HTA |
| 63 | ((pudend* NEAR3 neuralg*)) IN DARE, HTA |
| 64 | (proctalgia*) IN DARE, HTA |
| 65 | ((tension* NEXT myalgia*)) IN DARE, HTA |
| 66 | MeSH DESCRIPTOR Pelvic Floor IN DARE,HTA |
| 67 | MeSH DESCRIPTOR Pelvic Floor Disorders IN DARE,HTA |
| 68 | ((pelvi* adj (floor* or diaphragm*) NEAR3 (dysfunction* or disorder* or fail* or impair* or incompeten* or insufficien* or dyssynerg* or symptom* or laxity or change* or care* or health* or wellbeing* or well-being* or prevent* or rehabilitat*)) IN DARE, HTA |
| 69 | ((pelvi* NEXT (dysfunction* or disorder* or fail* or impair* or incompeten* or insufficien* or dyssynerg* or symptom* or laxity or care* or health* or wellbeing* or well-being* or prevent* or rehabilitat*)) IN DARE, HTA |
| 70 | #1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #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 OR #30 OR #31 OR #32 OR #33 OR #34 OR #35 OR #36 OR #37 OR #38 OR #39 OR #40 OR #41 OR #42 OR #43 OR #44 OR #45 OR #46 OR #47 OR #48 OR #49 OR #50 OR #51 OR #52 OR #53 OR #54 OR #55 OR #56 OR #57 OR #58 OR #59 OR #60 OR #61 OR #62 OR #63 OR #64 OR #65 OR #66 OR #67 OR #68 OR #69 |
| 71 | MeSH DESCRIPTOR Weight Loss IN DARE,HTA |
| 72 | MeSH DESCRIPTOR Weight Reduction Programs IN DARE,HTA |
| 73 | ((weight NEAR2 (los* or reduc*) NEAR3 (modif* or therap* or intervention* or strateg* or program* or management or scheme* or group* or pathway*)) IN DARE, HTA |
| 74 | ((weight NEXT management)) IN DARE, HTA |
| 75 | ((calori* or hypocalori*) NEAR2 (restrict* or diet*)) IN DARE, HTA |
| 76 | #71 OR #72 OR #73 OR #74 OR #75 |
| 77 | #70 AND #76 |

Database(s): EMCare – OVID interface

Date of last search: 2 February 2021

| # | Searches |
|---|-------------------------|
| 1 | exp urine incontinence/ |

| # | Searches |
|----|---|
| 2 | overactive bladder/ |
| 3 | bladder instability/ |
| 4 | nocturia/ |
| 5 | enuresis/ |
| 6 | exp enuresis/ |
| 7 | ((stress\$ or mix\$ or urg\$ or urin\$) adj5 incontinen\$).tw. |
| 8 | (bladder\$ adj5 (overactiv\$ or over activ\$ or over-activ\$ or instabili\$ or hyper-reflex\$ or hyperreflex\$ or hyper reflex\$ or incontinen\$)).tw. |
| 9 | (detrusor\$ adj5 (overactiv\$ or over activ\$ or over-activ\$ or instabili\$ or hyper-reflex\$ or hyperreflex\$ or hyper reflex\$)).tw. |
| 10 | ((urgency adj2 frequency) or (frequency adj2 urgency)).tw. |
| 11 | ((urin\$ or bladder\$) adj2 (urg\$ or frequen\$)).tw. |
| 12 | (nocturia\$ or enuresis\$).tw. |
| 13 | (SUI or OAB).tw. |
| 14 | or/1-13 |
| 15 | exp pelvic organ prolapse/ |
| 16 | rectocele/ |
| 17 | (pelvic\$ adj3 organ\$ adj3 prolaps\$).tw. |
| 18 | (urinary adj3 bladder adj3 prolaps\$).tw. |
| 19 | ((vagin\$ or urogenital\$ or genit\$ or uter\$ or viscer\$ or anterior\$ or posterior\$ or apical or pelvi\$ or vault\$ or urethr\$ or bladder\$ or cervi\$ or rectal or rectum) adj3 prolaps\$).tw. |
| 20 | (splanchnoptos\$ or visceroptos\$).tw. |
| 21 | (hernia\$ adj3 (pelvi\$ or vagin\$ or urogenital\$ or uter\$ or bladder\$ or urethr\$ or viscer\$)).tw. |
| 22 | (urethroc?ele\$ or enteroc?ele\$ or sigmoidoc?ele\$ or proctoc?ele\$ or rectoc?ele\$ or cystoc?ele\$ or rectoenteroc?ele\$ or cystourethroc?ele\$).tw. |
| 23 | or/15-22 |
| 24 | feces incontinence/ |
| 25 | ((faecal or fecal or faeces or feces or fecally or faecally or anal or anally or stool or stools or bowel or double or defecat\$ or defaecat\$) adj5 (incontinence or incontinent or urge\$ or leak or leaking or leakage or soiling or seeping or seepage or impacted or impaction)).tw. |
| 26 | 24 or 25 |
| 27 | urine retention/ |
| 28 | dysuria/ |
| 29 | (urin\$ adj3 (retention\$ or retain\$)).tw. |
| 30 | dysuria\$.tw. |
| 31 | (voiding adj (disorder\$ or dysfunction\$ or problem\$)).tw. |
| 32 | (empty\$ adj disorder\$ adj3 (bowel\$ or bladder\$ or vesical\$ or stool\$)).tw. |
| 33 | ((urogeni\$ or anorec\$ or ano-rec\$ or ano rec\$) adj3 dysfunction\$).tw. |
| 34 | defecation disorder/ |
| 35 | *constipation/ |
| 36 | feces impaction/ |
| 37 | constipat\$.ti. |
| 38 | ((difficult\$ or delay\$ or irregular\$ or infrequen\$ or pain\$) adj3 (defecat\$ or defaecat\$ or stool\$ or faecal or fecal or faeces or feces or fecally or faecally or bowel movement\$)).tw. |
| 39 | coprosthesis.tw. |
| 40 | (obstruct\$ adj3 (defecat\$ or defaecat\$)).tw. |
| 41 | ((defecat\$ or defaecat\$ or evacuat\$) adj3 (disorder\$ or dysfunction\$)).tw. |
| 42 | outlet\$ dysfunction\$ constipa\$.tw. |
| 43 | (dys?ynerg\$ adj (defecat\$ or defaecat\$)).tw. |
| 44 | (pelvi\$ adj3 dyskines\$).tw. |
| 45 | pelvi\$ outlet\$ obstruct\$.tw. |
| 46 | anismus\$.tw. |
| 47 | puborectal\$ contract\$.tw. |
| 48 | ((rectal or rectum) adj3 urge\$).tw. |
| 49 | or/27-48 |
| 50 | female sexual dysfunction/ |
| 51 | (female adj sex\$ adj (dysfunct\$ or satisf\$ or problem\$ or symptom\$ or arous\$ or activit\$ or disorder\$)).tw. |
| 52 | dyspareunia/ |
| 53 | (sex\$ adj3 pain\$).tw. |
| 54 | (dyspareun\$ or anodyspareun\$).tw. |
| 55 | (obstruct\$ adj3 intercourse).tw. |
| 56 | (vagin\$ adj3 laxity\$).tw. |
| 57 | (vagin\$ adj wind).tw. |
| 58 | orgasm disorder/ |
| 59 | (female adj orgasm\$ adj (disorder\$ or deficienc\$ or dysfunction\$ or problem\$)).tw. |
| 60 | anorgasm\$.tw. |
| 61 | vaginism/ |
| 62 | vaginismus\$.tw. |
| 63 | (vagin\$ adj penetrat\$ adj disorder\$).tw. |
| 64 | vulvodinia/ |
| 65 | vulvodinia\$.tw. |

| # | Searches |
|----|---|
| 66 | (vagin\$ adj dry\$.)\$.tw. |
| 67 | hypoactive sexual desire disorder/ |
| 68 | hypoactiv\$ sex\$ desire\$.tw. |
| 69 | sexual arousal disorder/ |
| 70 | (sex\$ adj arous\$ adj disorder\$.)\$.tw. |
| 71 | (genitourin\$ adj syndrom\$ adj5 menopaus\$.)\$.tw. |
| 72 | or/50-71 |
| 73 | pelvic pain/ |
| 74 | ((pelvi\$ or lumbopelvi\$ or lumbo-pelvi\$ or genito-pelvi\$ or genitopelvi\$) adj3 pain\$.)\$.tw. |
| 75 | (pubi\$ adj3 (pain\$ or dysfunction\$)).tw. |
| 76 | (pudend\$ adj3 neuralg\$.)\$.tw. |
| 77 | proctalgia\$.tw. |
| 78 | (tension\$ adj myalgia\$.)\$.tw. |
| 79 | or/73-78 |
| 80 | pelvis floor/ |
| 81 | pelvic floor disorder/ |
| 82 | (pelvi\$ adj (floor\$ or diaphragm\$) adj3 (dysfunction\$ or disorder\$ or fail\$ or impair\$ or incompeten\$ or insufficien\$ or dyssynerg\$ or symptom\$ or laxity or change\$ or care\$ or health\$ or wellbeing\$ or well-being\$ or prevent\$ or rehabilitat\$)).tw. |
| 83 | (pelvi\$ adj (dysfunction\$ or disorder\$ or fail\$ or impair\$ or incompeten\$ or insufficien\$ or dyssynerg\$ or symptom\$ or laxity or care\$ or health\$ or wellbeing\$ or well-being\$ or prevent\$ or rehabilitat\$)).tw. |
| 84 | or/80-83 |
| 85 | weight reduction/ |
| 86 | *body weight loss/ |
| 87 | body weight control/ |
| 88 | body weight change/ |
| 89 | weight loss program/ |
| 90 | (weight adj2 (los\$ or reduc\$) adj3 (modif\$ or therap\$ or intervention\$ or strateg\$ or program\$ or management or scheme\$ or group\$ or pathway)).tw. |
| 91 | (weight adj management).tw. |
| 92 | ((calori\$ or hypocalori\$) adj2 (restrict\$ or diet\$)).tw. |
| 93 | or/85-92 |
| 94 | (14 or 23 or 26 or 49 or 72 or 79 or 84) and 93 |
| 95 | limit 94 to (english language and yr="1980 -Current") [General Exclusions filter applied] |

Database(s): PsycINFO 1806 to January Week 4 2021 – OVID interface

Date of last search: 2 February 2021

| # | Searches |
|----|---|
| 1 | exp Urinary Incontinence/ |
| 2 | ((stress\$ or mix\$ or urg\$ or urin\$) adj5 incontinen\$.)\$.tw. |
| 3 | (bladder\$ adj5 (overactiv\$ or over activ\$ or over-activ\$ or instabilit\$ or hyper-reflex\$ or hyperreflex\$ or hyper reflex\$ or incontinen\$)).tw. |
| 4 | (detrusor\$ adj5 (overactiv\$ or over activ\$ or over-activ\$ or instabilit\$ or hyper-reflex\$ or hyperreflex\$ or hyper reflex\$)).tw. |
| 5 | ((urgency adj2 frequency) or (frequency adj2 urgency)).tw. |
| 6 | ((urin\$ or bladder\$) adj2 (urg\$ or frequen\$)).tw. |
| 7 | (nocturia\$ or enuresis\$.)\$.tw. |
| 8 | (SUI or OAB).tw. |
| 9 | (pelvic\$ adj3 organ\$ adj3 prolaps\$.)\$.tw. |
| 10 | (urinary adj3 bladder adj3 prolaps\$.)\$.tw. |
| 11 | ((vagin\$ or urogenital\$ or genit\$ or uter\$ or viscer\$ or anterior\$ or posterior\$ or apical or pelvi\$ or vault\$ or urethr\$ or bladder\$ or cervi\$ or rectal or rectum) adj3 prolaps\$.)\$.tw. |
| 12 | (splachnoptos\$ or visceroptos\$.)\$.tw. |
| 13 | (hernia\$ adj3 (pelvi\$ or vagin\$ or urogenital\$ or uter\$ or bladder\$ or urethr\$ or viscer\$)).tw. |
| 14 | (urethroc?ele\$ or enteroc?ele\$ or sigmoidoc?ele\$ or proctoc?ele\$ or rectoc?ele\$ or cystoc?ele\$ or rectoenteroc?ele\$ or cystourethroc?ele\$).tw. |
| 15 | exp Fecal Incontinence/ |
| 16 | ((faecal or fecal or faeces or feces or fecally or faecally or anal or anally or stool or stools or bowel or double or defecat\$ or defaecat\$) adj5 (incontinence or incontinent or urge\$ or leak or leaking or leakage or soiling or seeping or seepage or impacted or impaction)).tw. |
| 17 | (urin\$ adj3 (retention\$ or retain\$)).tw. |
| 18 | dysuria\$.tw. |
| 19 | (voiding adj (disorder\$ or dysfunction\$ or problem\$)).tw. |
| 20 | (empty\$ adj disorder\$ adj3 (bowel\$ or bladder\$ or vesical\$ or stool\$)).tw. |
| 21 | ((urogeni\$ or anorec\$ or ano-rec\$ or ano rec\$) adj3 dysfunction\$.)\$.tw. |
| 22 | exp Constipation/ |
| 23 | constipat\$.ti. |
| 24 | ((difficult\$ or delay\$ or irregular\$ or infrequen\$ or pain\$) adj3 (defecat\$ or defaecat\$ or stool\$ or faecal or fecal or faeces or feces or fecally or faecally or bowel movement\$)).tw. |
| 25 | coprostasis.tw. |

| # | Searches |
|----|---|
| 26 | (obstruct\$ adj3 (defecat\$ or defaecat\$)).tw. |
| 27 | ((defecat\$ or defaecat\$ or evacuat\$) adj3 (disorder\$ or dysfunction\$)).tw. |
| 28 | outlet\$ dysfunction\$ constipa\$.tw. |
| 29 | (dys?ynerg\$ adj (defecat\$ or defaecat\$)).tw. |
| 30 | (pelvi\$ adj3 dyskines\$).tw. |
| 31 | pelvi\$ outlet\$ obstruct\$.tw. |
| 32 | anismus\$.tw. |
| 33 | puborectal\$ contract\$.tw. |
| 34 | ((rectal or rectum) adj3 urge\$).tw. |
| 35 | exp Female Sexual Dysfunction/ |
| 36 | (female adj sex\$ adj (dysfunct\$ or satisf\$ or problem\$ or symptom\$ or arous\$ or activit\$ or disorder\$)).tw. |
| 37 | exp Dyspareunia/ |
| 38 | (sex\$ adj3 pain\$).tw. |
| 39 | (dyspareun\$ or anodyspareun\$).tw. |
| 40 | (obstruct\$ adj3 intercourse).tw. |
| 41 | (vagin\$ adj3 laxity\$).tw. |
| 42 | (vagin\$ adj wind).tw. |
| 43 | (female adj orgasm\$ adj (disorder\$ or deficienc\$ or dysfunction\$ or problem\$)).tw. |
| 44 | anorgasm\$.tw. |
| 45 | exp Vaginismus/ |
| 46 | vaginismus\$.tw. |
| 47 | (vagin\$ adj penetrat\$ adj disorder\$).tw. |
| 48 | vulvodynia\$.tw. |
| 49 | (vagin\$ adj dry\$).tw. |
| 50 | exp Inhibited Sexual Desire/ |
| 51 | hypoactiv\$ sex\$ desire\$.tw. |
| 52 | (sex\$ adj arous\$ adj disorder\$).tw. |
| 53 | (genitourin\$ adj syndrom\$ adj5 menopaus\$).tw. |
| 54 | ((pelvi\$ or lumbopelvi\$ or lumbo-pelvi\$ or genito-pelvi\$ or genitopelvi\$) adj3 pain\$).tw. |
| 55 | (pubi\$ adj3 (pain\$ or dysfunction\$)).tw. |
| 56 | (pudend\$ adj3 neuralg\$).tw. |
| 57 | proctalgia\$.tw. |
| 58 | (tension\$ adj myalgia\$).tw. |
| 59 | (pelvi\$ adj (floor\$ or diaphragm\$) adj3 (dysfunction\$ or disorder\$ or fail\$ or impair\$ or incompeten\$ or insufficien\$ or dyssynerg\$ or symptom\$ or laxity or change\$ or care\$ or health\$ or wellbeing\$ or well-being\$ or prevent\$ or rehabilitat\$)).tw. |
| 60 | (pelvi\$ adj (dysfunction\$ or disorder\$ or fail\$ or impair\$ or incompeten\$ or insufficien\$ or dyssynerg\$ or symptom\$ or laxity or care\$ or health\$ or wellbeing\$ or well-being\$ or prevent\$ or rehabilitat\$)).tw. |
| 61 | or/1-60 |
| 62 | exp Weight Loss/ |
| 63 | weight control/ |
| 64 | (weight adj2 (los\$ or reduc\$) adj3 (modif\$ or therap\$ or intervention\$ or strateg\$ or program\$ or management or scheme\$ or group\$ or pathway)).tw. |
| 65 | (weight adj management).tw. |
| 66 | ((calori\$ or hypocalori\$) adj2 (restrict\$ or diet\$)).tw. |
| 67 | or/62-66 |
| 68 | 61 and 67 |
| 69 | limit 68 to (english language and yr="1980 -Current") [General Exclusions filter applied] |

Economic Search

One global search was conducted for economic evidence across the guideline.

Database(s): NHS Economic Evaluation Database (NHS EED); HTA Database – CRD interface

Date of last search: 3 February 2021

| # | Searches |
|----|--|
| 1 | MeSH DESCRIPTOR Pelvic Floor IN NHSEED,HTA |
| 2 | MeSH DESCRIPTOR Pelvic Floor Disorders IN NHSEED,HTA |
| 3 | MeSH DESCRIPTOR Urinary Bladder, Overactive IN NHSEED,HTA |
| 4 | (((pelvi* NEXT (floor* or diaphragm*) NEAR3 (dysfunction* or disorder* or fail* or impair* or incompeten* or insufficien* or dyssynerg* or symptom* or laxity or change* or care* or health* or wellbeing* or well-being* or prevent* or rehabilitat* or weak* or hypertonic* or overactiv* or over activ* or over-activ*)))) IN NHSEED, HTA |
| 5 | MeSH DESCRIPTOR Urinary Incontinence EXPLODE ALL TREES IN NHSEED,HTA |
| 6 | MeSH DESCRIPTOR Urinary Bladder, Overactive IN NHSEED,HTA |
| 7 | (((stress* or mix* or urg* or urin*) NEAR5 incontinen*)) IN NHSEED, HTA |
| 8 | (((bladder* NEAR5 (overactiv* or over activ* or over-activ* or instabilit* or hyper-reflex* or hyperreflex* or hyper reflex* or incontinen*)) IN NHSEED, HTA |
| 9 | (((detrusor* NEAR5 (overactiv* or over activ* or over-activ* or instabilit* or hyper-reflex* or hyperreflex* or hyper reflex*)) IN NHSEED, HTA |
| 10 | (((urgency NEAR2 frequency) or (frequency NEAR2 urgency))) IN NHSEED, HTA |
| 11 | (((urin* or bladder*) NEAR2 (urg* or frequen*)) IN NHSEED, HTA |
| 12 | (((SUI or OAB))) IN NHSEED, HTA |
| 13 | MeSH DESCRIPTOR Pelvic Organ Prolapse EXPLODE ALL TREES IN NHSEED,HTA |
| 14 | MeSH DESCRIPTOR Rectocele IN NHSEED,HTA |
| 15 | (((pelvic* NEAR3 organ* NEAR3 prolaps*)) IN NHSEED, HTA |
| 16 | (((urinary NEAR3 bladder NEAR3 prolaps*)) IN NHSEED, HTA |
| 17 | (((vagin* or urogenital* or genit* or uter* or viscer* or anterior* or posterior* or apical or pelvi* or vault* or urethr* or bladder* or cervi* or rectal or rectum) NEAR3 prolaps*)) IN NHSEED, HTA |
| 18 | (((splanchnoptos* or visceroptos*)) IN NHSEED, HTA |
| 19 | (((hernia* NEAR3 (pelvi* or vagin* or urogenital* or uter* or bladder* or urethr* or viscer*)) IN NHSEED, HTA |
| 20 | (((urethroc?ele* or enteroc?ele* or sigmoidoc?ele* or proctoc?ele* or rectoc?ele* or cystoc?ele* or rectoenteroc?ele* or cystourethroc?ele*)) IN NHSEED, HTA |
| 21 | MeSH DESCRIPTOR Fecal Incontinence IN NHSEED,HTA |
| 22 | (((faecal or fecal or faeces or feces or fecally or faecally or anal or anally or stool or stools or bowel or double or defecat* or defaecat*) NEAR5 (incontinence or incontinent or urge* or leak or leaking or leakage or soiling or seeping or seepage or impacted or impaction))) IN NHSEED, HTA |
| 23 | MeSH DESCRIPTOR Urinary Retention IN NHSEED,HTA |
| 24 | (((urin* NEAR3 (retention* or retain*)) IN NHSEED, HTA |
| 25 | (((voiding NEXT (disorder* or dysfunction* or problem*)) IN NHSEED, HTA |
| 26 | (((empty* NEXT disorder* NEAR3 (bowel* or bladder* or vesical* or stool*)) IN NHSEED, HTA |
| 27 | (((urogeni* or anorec* or ano-rec* or ano rec*) NEAR3 dysfunction*)) IN NHSEED, HTA |
| 28 | MeSH DESCRIPTOR Fecal Impaction IN NHSEED,HTA |
| 29 | (((difficult* or delay* or irregular* or infrequen* or pain*) NEAR3 (defecat* or defaecat* or stool* or faecal or fecal or faeces or feces or fecally or faecally or bowel movement*)) IN NHSEED, HTA |
| 30 | (((obstruct* NEAR3 (defecat* or defaecat*)) IN NHSEED, HTA |
| 31 | (((defecat* or defaecat* or evacuat*) NEAR3 (disorder* or dysfunction*)) IN NHSEED, HTA |
| 32 | (((outlet* NEXT dysfunction* NEXT constipa*)) IN NHSEED, HTA |
| 33 | (((dys?ynerg* NEXT (defecat* or defaecat*)) IN NHSEED, HTA |
| 34 | (((pelvi* NEAR3 dyskines*)) IN NHSEED, HTA |
| 35 | (((pelvi* NEXT outlet* NEXT obstruct*)) IN NHSEED, HTA |
| 36 | (((anismus*)) IN NHSEED, HTA |
| 37 | (((puborectal* NEXT contract*)) IN NHSEED, HTA |
| 38 | (((rectal or rectum) NEAR3 urge*)) IN NHSEED, HTA |
| 39 | (((female NEXT sex* NEXT (dysfunct* or satisf* or problem* or symptom* or arous* or activit* or disorder*)) IN NHSEED, HTA |
| 40 | (((obstruct* NEAR3 intercourse))) IN NHSEED, HTA |
| 41 | (((vagin* NEAR3 laxity*)) IN NHSEED, HTA |
| 42 | (((vagin* NEXT wind))) IN NHSEED, HTA |
| 43 | MeSH DESCRIPTOR Vaginismus IN NHSEED,HTA |
| 44 | (((vaginismus*)) IN NHSEED, HTA |
| 45 | (((vagin* NEXT penetrat* NEXT disorder*)) IN NHSEED, HTA |
| 46 | (#1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #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 OR #30 OR #31 OR #32 OR #33 OR #34 OR #35 OR #36 OR #37 OR #38 OR #39 OR #40 OR #41 OR #42 OR #43 OR #44 OR #45) IN NHSEED, HTA |

Database(s): Medline & Embase (Multifile) – OVID interface

Embase Classic+Embase 1947 to 2021 February 01; Ovid MEDLINE(R) and Epub Ahead of Print, In-Process & Other Non-Indexed Citations and Daily 1946 to February 01, 2021
Date of last search: 3 February 2021

Multifile database codes: emczd = Embase Classic+Embase; ppez= MEDLINE(R) and Epub Ahead of Print, In-Process & Other Non-Indexed Citations and Daily

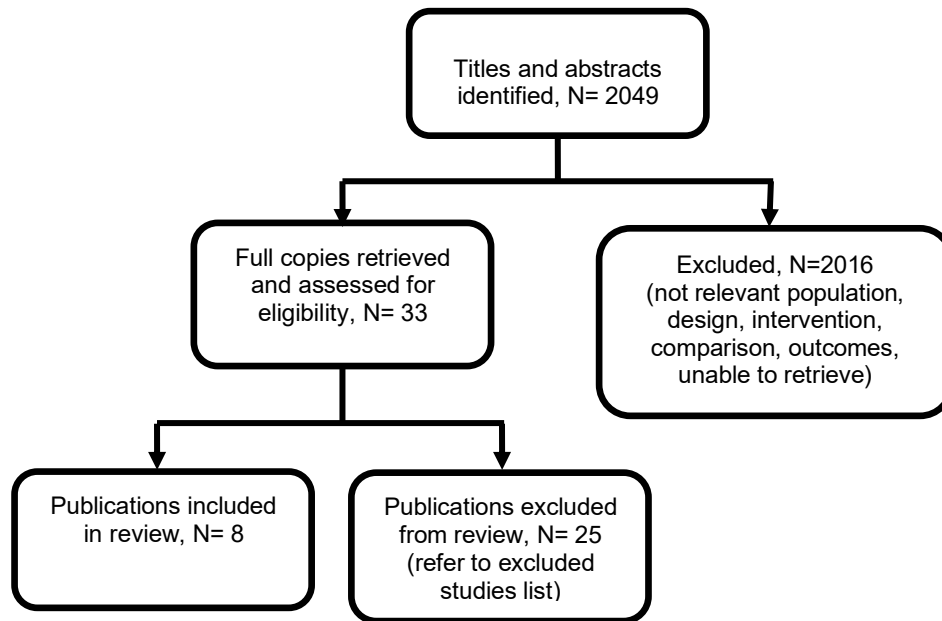
| # | Searches |
|----|--|
| 1 | Pelvic Floor/ use ppez |
| 2 | Pelvic Floor Disorders/ use ppez |
| 3 | pelvis floor/ use emczd |
| 4 | pelvic floor disorder/ use emczd |
| 5 | (pelvi\$ adj (floor\$ or diaphragm\$) adj3 (dysfunction\$ or disorder\$ or fail\$ or impair\$ or incompeten\$ or insufficien\$ or dyssynerg\$ or symptom\$ or laxity or change\$ or care\$ or health\$ or wellbeing\$ or well-being\$ or prevent\$ or rehabilitat\$ or weak\$ or hypertonic\$ or overactiv\$ or over activ\$ or over-activ\$)).tw. |
| 6 | (pelvi\$ adj (dysfunction\$ or disorder\$ or fail\$ or impair\$ or incompeten\$ or insufficien\$ or dyssynerg\$ or symptom\$ or laxity or care\$ or health\$ or wellbeing\$ or well-being\$ or prevent\$ or rehabilitat\$ or weak\$ or hypertonic\$ or overactiv\$ or over activ\$ or over-activ\$)).tw. |
| 7 | or/1-6 |
| 8 | exp *Urinary Incontinence/ use ppez |
| 9 | *Urinary Bladder, Overactive/ use ppez |
| 10 | exp *urine incontinence/ use emczd |
| 11 | *overactive bladder/ use emczd |
| 12 | *bladder instability/ use emczd |
| 13 | ((stress\$ or mix\$ or urg\$ or urin\$) adj5 incontinen\$).ti. |
| 14 | (bladder\$ adj5 (overactiv\$ or over activ\$ or over-activ\$ or instabilit\$ or hyper-reflex\$ or hyperreflex\$ or hyper reflex\$ or incontinen\$)).ti. |
| 15 | (detrusor\$ adj5 (overactiv\$ or over activ\$ or over-activ\$ or instabilit\$ or hyper-reflex\$ or hyperreflex\$ or hyper reflex\$)).ti. |
| 16 | ((urgency adj2 frequency) or (frequency adj2 urgency)).ti. |
| 17 | ((urin\$ or bladder\$) adj2 (urg\$ or frequen\$)).ti. |
| 18 | (SUI or OAB).ti. |
| 19 | or/8-18 |
| 20 | exp *Pelvic Organ Prolapse/ use ppez |
| 21 | exp *pelvic organ prolapse/ use emczd |
| 22 | *Rectocele/ use ppez |
| 23 | *rectocele/ use emczd |
| 24 | (pelvic\$ adj3 organ\$ adj3 prolaps\$).ti. |
| 25 | (urinary adj3 bladder adj3 prolaps\$).ti. |
| 26 | ((vagin\$ or urogenital\$ or genit\$ or uter\$ or viscer\$ or anterior\$ or posterior\$ or apical or pelvi\$ or vault\$ or urethr\$ or bladder\$ or cervi\$ or rectal or rectum) adj3 prolaps\$).ti. |
| 27 | (splachnoptos\$ or visceroptos\$).ti. |
| 28 | (hernia\$ adj3 (pelvi\$ or vagin\$ or urogenital\$ or uter\$ or bladder\$ or urethr\$ or viscer\$)).ti. |
| 29 | (urethro?ele\$ or enteroc?ele\$ or sigmoidoc?ele\$ or proctoc?ele\$ or rectoc?ele\$ or cystoc?ele\$ or rectoenteroc?ele\$ or cystourethro?ele\$).ti. |
| 30 | or/20-29 |
| 31 | *Fecal Incontinence/ use ppez |
| 32 | *feces incontinence/ use emczd |
| 33 | ((faecal or fecal or faeces or feces or fecally or faecally or anal or anally or stool or stools or bowel or double or defecat\$ or defaecat\$) adj5 (incontinence or incontinent or urge\$ or leak or leaking or leakage or soiling or seeping or seepage or impacted or impaction)).ti. |
| 34 | or/31-33 |
| 35 | Urinary Retention/ use ppez |
| 36 | urine retention/ use emczd |
| 37 | (urin\$ adj3 (retention\$ or retain\$)).tw. |
| 38 | (voiding adj (disorder\$ or dysfunction\$ or problem\$)).tw. |
| 39 | (empty\$ adj disorder\$ adj3 (bowel\$ or bladder\$ or vesical\$ or stool\$)).tw. |
| 40 | ((urogeni\$ or anorec\$ or ano-rec\$ or ano rec\$) adj3 dysfunction\$).tw. |
| 41 | defecation disorder/ use emczd |
| 42 | Fecal Impaction/ use ppez |
| 43 | Feces Impaction/ use emczd |
| 44 | ((difficult\$ or delay\$ or irregular\$ or infrequen\$ or pain\$) adj3 (defecat\$ or defaecat\$ or stool\$ or faeces or feces or bowel movement\$)).tw. |
| 45 | (obstruct\$ adj3 (defecat\$ or defaecat\$)).tw. |
| 46 | ((defecat\$ or defaecat\$ or evacuat\$) adj3 (disorder\$ or dysfunction\$)).tw. |
| 47 | outlet\$ dysfunction\$ constipa\$.tw. |
| 48 | (dys?ynerg\$ adj (defecat\$ or defaecat\$)).tw. |
| 49 | (pelvi\$ adj3 dyskines\$).tw. |
| 50 | pelvi\$ outlet\$ obstruct\$.tw. |
| 51 | anismus\$.tw. |
| 52 | puborectal\$ contract\$.tw. |
| 53 | ((rectal or rectum) adj3 urge\$).tw. |

| # | Searches |
|----|---|
| 54 | or/35-53 |
| 55 | female sexual dysfunction/ use emczd |
| 56 | (female adj sex\$ adj (dysfunct\$ or satisf\$ or problem\$ or symptom\$ or arous\$ or activit\$ or disorder\$)).tw. |
| 57 | (obstruct\$ adj3 intercourse).tw. |
| 58 | (vagin\$ adj3 laxity\$).tw. |
| 59 | (vagin\$ adj wind).tw. |
| 60 | Vaginismus/ use ppez |
| 61 | vaginism/ use emczd |
| 62 | vaginismus\$.tw. |
| 63 | (vagin\$ adj penetrat\$ adj disorder\$).tw. |
| 64 | or/55-63 |
| 65 | 7 or 19 or 30 or 34 or 54 or 64 |
| 66 | Economics/ use ppez |
| 67 | Value of life/ use ppez |
| 68 | exp "Costs and Cost Analysis"/ use ppez |
| 69 | exp Economics, Hospital/ use ppez |
| 70 | exp Economics, Medical/ use ppez |
| 71 | Economics, Nursing/ use ppez |
| 72 | Economics, Pharmaceutical/ use ppez |
| 73 | exp "Fees and Charges"/ use ppez |
| 74 | exp Budgets/ use ppez |
| 75 | health economics/ use emczd |
| 76 | exp economic evaluation/ use emczd |
| 77 | exp health care cost/ use emczd |
| 78 | exp fee/ use emczd |
| 79 | budget/ use emczd |
| 80 | funding/ use emczd |
| 81 | budget*.ti,ab. |
| 82 | cost*.ti. |
| 83 | (economic* or pharmaco?economic*).ti. |
| 84 | (price* or pricing*).ti,ab. |
| 85 | (cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab. |
| 86 | (financ* or fee or fees).ti,ab. |
| 87 | (value adj2 (money or monetary)).ti,ab. |
| 88 | or/66-87 |
| 89 | 65 and 88 |
| 90 | limit 89 to english language |

Appendix C – Clinical evidence study selection

Study selection for: What is the effectiveness of weight loss interventions for improving symptoms of pelvic floor dysfunction?

Figure 1: Study selection flow chart



Appendix D – Evidence tables

Evidence tables for review question: What is the effectiveness of weight loss interventions for improving symptoms of pelvic floor dysfunction?

Table 4: Evidence tables

| Study details | Participants | Interventions | Methods | Outcomes | Comments |
|--|---|---|--|---|---|
| <p>Full citation</p> <p>Breyer, B. N., Creasman, J. M., Richter, H. E., Myers, D., Burgio, K. L., Wing, R. R., West, D. S., Kusek, J. W., Subak, L. L., Pride, A Behavioral Weight Loss Program and Nonurinary Incontinence Lower Urinary Tract Symptoms in Overweight and Obese Women with Urinary Incontinence: A Secondary Data Analysis of PRIDE, Journal of urology, 199, 215-222, 2018</p> <p>Ref Id</p> <p>1118105</p> <p>Country/ies where the study was carried out</p> <p>USA</p> <p>Study type</p> <p>Randomised controlled trial</p> | <p>Sample size</p> <p>See Subak 2009</p> <p>Characteristics</p> <p>See Subak 2009</p> <p>Inclusion criteria</p> <p>See Subak 2009</p> <p>Exclusion criteria</p> <p>See Subak 2009</p> | <p>Interventions</p> <p>See Subak 2009</p> | <p>Details</p> <p>Non-UI lower urinary tract storage symptoms (LUTS), including daytime urinary frequency, nocturia, and urinary urgency, were measured by a participant-completed 7-day voiding diary and self-report questionnaires completed at baseline and 6 months.</p> <p>The International Prostate Symptom Score (IPSS), which has been validated for use in women, was administered to obtain the composite outcome of non-UI LUTS.</p> | <p>Results</p> <p>Outcomes at baseline and after 6months</p> <p><u>Daytime Frequency:</u> Pre: Intervention 109/214 (48%); Control 63/90 (56%) Post: Intervention 93/214 (43%); Control 47/90 (52%)</p> <p><u>Nocturia</u> Pre: Intervention 112/214 (50%); Control 61/90 (54%) Post: Intervention 83/214 (38%); Control 47/90 (48%)</p> <p><u>Urinary Urgency</u> Pre: Intervention 138/214 (61%); Control 71/90 (63%) Post: Intervention 89/214 (40%); Control 46/90 (47%)</p> <p><u>International Prostate Symptom Score (IPSS) Score \geq8</u></p> | <p>Limitations</p> <p>See Subak 2009</p> <p>Other information</p> <p>See Subak 2009</p> |

| Study details | Participants | Interventions | Methods | Outcomes | Comments |
|--|---|--|--|--|--|
| <p>Aim of the study See Subak 2009</p> <p>Study dates See Subak 2009</p> <p>Source of funding See Subak 2009</p> | | | | <p>Pre: Intervention 137/214 (61%); Control 72/90 (64%) Post: Intervention 76/214 (35%); Control 40/90 (41%)</p> | |
| <p>Full citation Gozukara, Y. M., Akalan, G., Tok, E. C., Aytan, H., Ertunc, D., The improvement in pelvic floor symptoms with weight loss in obese women does not correlate with the changes in pelvic anatomy, International Urogynecology Journal, 25, 1219-25, 2014</p> <p>Ref Id 541434</p> <p>Country/ies where the study was carried out Turkey</p> <p>Study type Randomised Controlled Trial</p> <p>Aim of the study</p> | <p>Sample size n=378</p> <p>Characteristics Age: Intervention 44.1 (8.6) years; Control 43.8 (9.7) years; BMI: Intervention 32.7 (4.1) kg/m²; Control 32.3 (3.6) kg/m² Parity: Intervention 3.1 (2.0); Control 2.9 (1.9) Vaginal birth: Intervention 139 (85.3%); Control 132 (83.5%) Postmenopausal: Intervention 51 (31.3%); Control 48 (30.4%) Stress UI: Intervention 66 (40.5%); Control 62 (39.2%)</p> | <p>Interventions 6-month behavioural weight loss program (intervention) or a structured education program (control)</p> <p>Intervention group: a calorie and fat restricted diet of 1,200–1,800 kcal daily, depending on initial weight, with less than 30 % of calories from fat, and was designed to produce an average loss of 7–9 % of initial body weight within 6 months.</p> <p>Participants met monthly for 6 months in groups of 15–20 for 1-h sessions that were led by an internist in nutrition, exercise, and behaviour change. Participants were provided with sample meal plans, encouraged to gradually increase physical activity, and self-</p> | <p>Details Pelvic anatomy was assessed in all participants before interventions and at the end of the study period with the POP-Q system. UI was assessed with a 3-day voiding diary. The Pelvic Floor Distress Inventory-20 (PFDI-20) assessed the impact of pelvic floor disorders on quality of life (QOL)</p> | <p>Results Study completed by n=163 women in the intervention group and n=158 women in the control group <u>Body weight (kg):</u> Pre: Intervention 85.1 (9.7); Control 85.9 (9.6) Post: Intervention 78.3 (10.4); Control 86.4 (11.7)</p> <p>PFDI Scores (mean (SE)): <u>Urinary Distress Inventory (UDI)-6:</u> Pre: Intervention 14.5 (1.4); Control 14.6 (2.4) Post: Intervention 13.0 (2.4); Control 14.7 (2.9)</p> <p><u>Colorectal-Anal Distress Inventory (CRADI)-8</u> Pre: Intervention 10.8 (2.1); Control 10.6 (1.5) Post: Intervention 10.1 (2.1); Control 10.0 (1.6)</p> | <p>Limitations Cochrane risk of bias (Version 2.0)</p> <p>Domain 1: Randomisation: Low risk 1.1: Yes, patients were randomly allocated to treatments 1.2: Yes, randomisation used tamper proof envelopes 1.3: No, no significant differences between groups at baseline</p> <p>Domain 2: Deviations from intended interventions: Some risk 2.1: Yes, participants not blinded 2.2: Yes, carers and people delivering the interventions not blinded 2.3: No information whether there were any deviations from the intended intervention</p> |

| Study details | Participants | Interventions | Methods | Outcomes | Comments |
|--|---|---|---------|---|---|
| <p>To investigate the effects of weight loss on urinary incontinence episodes and pelvic floor anatomy</p> <p>Study dates June 2008 and December 2008</p> <p>Source of funding None reported</p> | <p>Urge UI: Intervention 34 (20.9%); Control 40 (25.3%)</p> <p>Stress + Urge UI: Intervention 63 (38.7%); Control 56 (35.4%)</p> <p>POP-Q stage 0-I: Intervention 116 (71.2%); Control 108 (68.45%)</p> <p>POP-Q stage II-IV: Intervention 47 (28.8%); Control 50 (31.6%)</p> <p>Inclusion criteria Having five or more episodes of any UI in a 3-day voiding diary and a BMI over 25 kg/m².</p> <p>Exclusion criteria Women who had used medical therapy for incontinence or made any attempt at weight loss within the previous month and women with urinary tract infection, pregnancy, or parturition in the previous 6 months and previous genitourinary surgery.</p> <p>Patients with UI due to neurological or functional origins, or with significant systemic and</p> | <p>monitoring of diet and exercise.</p> | | <p><u>Pelvic Organ Prolapse Distress Inventory (POPDI)-6</u> Pre: Intervention 19.9 (2.9); Control 18.5 (1.6) Post: Intervention 14.5 (3.1); Control 17.4 (1.8)</p> <p><u>PFDI total</u> Pre: Intervention 45.3 (4.4); Control 43.6 (2.8) Post: Intervention 37.3 (3.9); Control 42.1 (3.2)</p> <p>UI episodes (mean (SE)): <u>Stress incontinence</u> Pre: Intervention 7.96 (0.17); Control 7.08 (0.15) Post: Intervention 3.11 (0.13); Control 7.03 (0.13)</p> <p><u>Urge Incontinence</u> Pre: Intervention 2.85 (0.11); Control 2.74 (0.09) Post: Intervention 1.08 (0.08); Control 2.48 (0.08)</p> | <p>Domain 3: Missing outcome data: Low risk 3.1: Probably no, 14% of the intervention group and 16% in the control group were lost to follow-up 3.2: Probably no, no evidence that the results were not biased by missing outcome data 3.3: Probably no, missingness of the outcome was not dependent on its true value</p> <p>Domain 4: Measurement of the outcome: Low risk 4.1: Probably no, outcomes clearly defined, but some information on how they were assessed and by whom 4.2: Probably no, outcomes unlikely to differ between treatment arms 4.3: No, outcome assessors were blinded</p> <p>Domain 5: Selection of the reported result: Some concerns 5.1: No, no pre-panned analysis or protocol available 5.2: No, descriptive data presented 5.3: No, data presented as expected</p> <p>Domain 6: Overall judgment of bias: Some concerns</p> |

| Study details | Participants | Interventions | Methods | Outcomes | Comments |
|---|---|---|--|---|---|
| | <p>genitourinary medical conditions, and women who required assistance during their daily activities.</p> <p>Patients who were using any medication that potentially affects urinary continence (e.g., cholinergic and anticholinergic agents, certain antihypertensives, diuretics, opioids, and certain psychotropic drugs)</p> | | | | |
| <p>Full citation</p> <p>Hagovska, M., Svihra, J., Bukova, A., Drackova, D., Horbacz, A., Nagyova, I., Effect of an Exercise Programme for Reducing Abdominal Fat on Overactive Bladder Symptoms in Young Overweight Women, Obstetrical and Gynecological Survey, 75, 471-472, 2020</p> <p>Ref Id</p> <p>1287224</p> <p>Country/ies where the study was carried out</p> <p>Slovakia</p> <p>Study type</p> <p>RCT</p> | <p>Sample size</p> <p>n=93</p> <p>Characteristics</p> <p>Age (mean, SD): Intervention group 26.7 (4.8); Control 26.9 (4.9) years</p> <p>OAB-q - SS symptoms score (mean, SD): Intervention group 10.9 (10.6); Control group 11.1 (8.6)</p> <p>OAB-q - HR-quality of life (mean, SD): Intervention group 94.6 (6.6); Control group 93.0 (7.6)</p> <p>BMI (mean, SD): Intervention group 25.0 (3.0); control group 25.1 (4.2)</p> | <p>Interventions</p> <p>Intervention (n=46): Programme for reducing abdominal fat, with activation of deep abdominal muscles and strengthening of the surface abdominal muscles. The duration of the intervention was 12 weeks. The women did not change their movement and physical activities other than the intervention. In addition, their eating habits were not changed during this study. Exercise was performed two times a week for 60–80 min under the supervision of a sports trainer and physiotherapist. The training programme had elements of: (1) Aerobic training: a stationary bicycle for 20 min; (2)</p> | <p>Details</p> <p>OAB symptoms were measured using the OAB-q.: This questionnaire focuses on the symptoms of an overactive bladder in the last 4 weeks. It contains six questions, the symptom score (0 without symptoms, 100 = the most symptoms) and 13 questions that assess quality of life (100 = best quality of life, 0 = the worst quality of life)</p> | <p>Results</p> <p>OAB-Q: SS-symptom score (mean, SD; final score; 10 weeks)</p> <p>Intervention (n=34): 1 (1.3)</p> <p>Control (n=36): 11.9 (1.4)</p> <p>OAB-Q: HR-quality of life score (mean, SD; final score; 10 weeks)</p> <p>Intervention (n=34): 100 (0.9)</p> <p>Control (n=36): 93 (0.9)</p> <p>Weight (mean, SD; kg; final score; 10 weeks)</p> <p>Intervention (n=34): 61.8 (1.6)</p> <p>Control (n=36): 66.9 (1.7)</p> <p>BMI (mean, SD; kg/m2; final score; 10 weeks)</p> <p>Intervention (n=34): 23.2 (0.6)</p> <p>Control (n=36): 25 (0.6)</p> | <p>Limitations</p> <p>Limitations</p> <p>Cochrane risk of bias (Version 2.0)</p> <p>Domain 1: Randomisation: Low risk</p> <p>1.1: Probably yes, patients were randomly allocated to treatments using Microsoft Excel</p> <p>1.2: No information</p> <p>1.3: Probably yes, no significant differences between groups</p> <p>Domain 2: Deviations from intended interventions: Some risk</p> <p>2.1: Yes, participants not blinded</p> <p>2.2: Yes, carers and people delivering the interventions not blinded</p> <p>2.3: Yes, states that twelve women did not complete the exercise</p> |

| Study details | Participants | Interventions | Methods | Outcomes | Comments |
|---|---|--|---------|----------|--|
| <p>Aim of the study To reduce symptoms of OAB through a 3-month exercise programme in young overweight women with OAB</p> <p>Study dates Between March and September 2018</p> <p>Source of funding This work was supported by the Slovak Research and Development Agency</p> | <p>Body weight (mean, SD), kg: Intervention group 66 (8.4); Control group 67 (11.5)</p> <p>Inclusion criteria Women aged 18–35, BMI 25–29.9, waist circumference > 88 cm and OAB defined as urgency usually accompanied by frequency and nocturia, with or without urge urinary incontinence (UUI), with the absence of urinary tract infections and other pathologies</p> <p>Exclusion criteria Stress urinary incontinence (SUI), surgical treatment of gynaecological and urological diseases, urinary tract infection, oncological and neurological urinary tract disease, incomplete questionnaires and refusal to participate in the study</p> | <p>Dynamic warm-up: stretching muscles by slow and controlled movements, 10 min; (3) Strength training: for reduction of abdominal fat, deep abdominal muscle activation (M. transversus abdominis, M. obliquus abdominis internus), 20 min, strengthening of superficial abdominal muscles (M. obliquus abdominis externus, M. rectus abdominis). (4) Static stretching: passive stretching of lower limbs and abdominal muscles, 10 min.</p> <p>Control (n=47): The control group did not undergo the exercise programme. The women did not change their everyday life activities or their eating habits.</p> | | | <p>programme in the experimental group, and 11 did not appear for examination in the control group. Further, the level of adherence was 75%</p> <p>2.4: Probably yes, drop out and non-adherence is likely to effect the outcome</p> <p>2.5: No, drop out is similar between the groups</p> <p>Domain 3: Missing outcome data: Low risk</p> <p>3.1: Probably no, 26% of the intervention group and 23% in the control group were lost to follow-up or excluded from final analysis</p> <p>3.2: Probably no, no evidence that the results were not biased by missing outcome data</p> <p>3.3: Probably no, missingness of the outcome was not dependent on its true value</p> <p>Domain 4: Measurement of the outcome: High risk</p> <p>4.1: Probably no, outcomes clearly defined and some information on how they were assessed</p> <p>4.2: Probably no, outcomes unlikely to differ between treatment arms</p> <p>4.3: Yes, due to self-report</p> |

| Study details | Participants | Interventions | Methods | Outcomes | Comments |
|--|---|--|--|---|---|
| | | | | | <p>4.4: Probably yes, as the control group did not have any intervention</p> <p>4.5: Probably yes, as the control group did not have any intervention</p> <p>Domain 5: Selection of the reported result: Some concerns</p> <p>5.1: No, no pre-planned analysis or protocol available</p> <p>5.2: No, descriptive data presented</p> <p>5.3: No, data presented as expected</p> <p>Domain 6: Overall judgment of bias: High risk</p> <p>Other information</p> |
| <p>Full citation</p> <p>Myers, D. L., Sung, V. W., Richter, H. E., Creasman, J., Subak, L. L., Prolapse symptoms in overweight and obese women before and after weight loss, Female Pelvic Medicine & Reconstructive Surgery, 18, 55-9, 2012</p> <p>Ref Id</p> <p>541569</p> <p>Country/ies where the study was carried out</p> | <p>Sample size See Subak 2009</p> <p>Characteristics See Subak 2009</p> <p>Inclusion criteria See Subak 2009</p> <p>Exclusion criteria See Subak 2009</p> | <p>Interventions See Subak 2009</p> | <p>Details See Subak 2009</p> | <p>Results</p> <p>Change in baseline for 'Any POP Symptoms' at 6 months</p> <p><u>Cured</u> Post: Intervention 73/NR (68%); Control 35/NR (71%) (Estimated by NGA team to be 73/107 vs 35/49 for the purpose of data analysis)</p> <p><u>Improved bother or cured</u> Post: Intervention 82/NR (77%); Control 38/NR (78%) (Estimated by NGA</p> | <p>Limitations See Subak 2009</p> <p>Other information The number of participants for each outcome was calculated by the NGA team as these were not reported in the paper</p> |

| Study details | Participants | Interventions | Methods | Outcomes | Comments |
|--|--|---|---|---|---|
| <p>USA</p> <p>Study type Randomised controlled trial</p> <p>Aim of the study See Subak 2009</p> <p>Study dates See Subak 2009</p> <p>Source of funding See Subak 2009</p> | | | | <p>team to be 82/107 vs 38/49 for the purpose of data analysis)</p> <p><u>New report of the symptoms</u> Post: Intervention 35/NR (16%); Control 20/NR (21%) (Estimated by NGA team to be 35/214 vs 20/94 for the purpose of data analysis)</p> | |
| <p>Full citation</p> <p>Subak, L. L., Wing, R., West, D. S., Franklin, F., Vittinghoff, E., Creasman, J. M., Richter, H. E., Myers, D., Burgio, K. L., Gorin, A. A., Macer, J., Kusek, J. W., Grady, D., Pride Investigators, Weight loss to treat urinary incontinence in overweight and obese women, New England journal of medicine, 360, 481-90, 2009</p> <p>Ref Id</p> <p>1118467</p> | <p>Sample size n=338</p> <p>Characteristics Age: Intervention 53 (11) years; Control 53 (10) years; BMI: Intervention 36 (6) kg/m²; Control 36 (5) kg/m² Race - White: Intervention 171 (75.7%); Control 91 (81.2%) Race - Black: Intervention 47 (20.8%); Control 17 (15.2%)</p> | <p>Interventions All participants were given a self-help behavioural-treatment booklet for improving bladder control. Incontinence was not discussed further in either the control group or the weight-loss group. Control group: participated in four education sessions at months 1, 2, 3, and 4. During these 1-hour group sessions, which included 10 to 15 women, general information was presented about weight loss, physical activity, and healthful eating habits,</p> | <p>Details Participants were randomly assigned at a 2:1 ratio to an intensive 6-month behavioural weight-loss program (intervention group) or to a structured four-session education program (the control group). Demographic characteristics, medical and behavioural history, and history of incontinence was collected. The participants were trained to complete a 7-day diary of voiding, and interviewers reviewed the</p> | <p>Results Study completed by n=226 women in the intervention group and n=112 women in the control group. Unless stated, data provided on n=214 intervention and n=90 control <u>Body weight (kg) (n=221 intervention and n=97 control):</u> Pre: Intervention 98 (17); Control 95 (16) Post: Intervention 90 (17); Control 94 (17)</p> <p><u>Any incontinence (no./wk)</u> Pre: Intervention 24 (18); Control 24 (16)</p> | <p>Limitations Cochrane risk of bias (Version 2.0)</p> <p>Domain 1: Randomisation: Low risk 1.1: Yes, patients were randomly allocated to treatments 1.2: Yes, randomisation used tamper proof envelopes 1.3: No, no significant differences between groups at baseline</p> <p>Domain 2: Deviations from intended interventions: Some risk</p> |

| Study details | Participants | Interventions | Methods | Outcomes | Comments |
|---|---|--|--|--|---|
| <p>Country/ies where the study was carried out</p> <p>USA</p> <p>Study type</p> <p>Randomised Controlled Trial</p> <p>Aim of the study</p> <p>The aim of the Program to Reduce Incontinence by Diet and Exercise (PRIDE), was to determine whether a behavioural weight-reduction intervention for overweight and obese women with incontinence would result in greater reductions in the frequency of incontinence episodes at 6 months as compared with a control group.</p> <p>Study dates</p> <p>July 2004 to April 2006</p> <p>Source of funding</p> <p>Supported by grants from the National Institute of Diabetes and Digestive and Kidney Diseases (NIDDK) (U01 DK067860, U01 DK067861, and U01 DK067862) and from the</p> | <p>Race - Other: Intervention 12 (3.6%); Control 4 (3.5%)</p> <p>Parity: Intervention 2 (1); Control 2 (1)</p> <p>Postmenopausal: Intervention 115/209 (55%); Control 62/107 (57.9%)</p> <p>Hysterectomy: Intervention 70/225 (31.1%); Control 29/112 (25.9%)</p> <p>Stress UI: Intervention 8 (3.5%); Control 10 (8.9%)</p> <p>Urge UI: Intervention 33 (14.6%); Control 8 (7.1%)</p> <p>Stress predominant: Intervention 36 (15.9%); Control 21 (18.8%)</p> <p>Urge predominant: Intervention 71 (31.4%); Control 37 (33.0%)</p> <p>Mixed incontinence with no predominant type: Intervention 78 (34.5%); Control 36 (32.1%)</p> <p>Inclusion criteria</p> <p>Women who were at least 30 years of age, had a BMI of 25 to 50, and at baseline reported 10 or</p> | <p>according to a structured protocol.</p> <p>Intervention group: The weight-loss program was designed to produce an average loss of 7 to 9% of initial body weight within the first 6 months of the program. The participants in the weight-loss program met weekly for 6 months in groups of 10 to 15 for 1-hour sessions that were led by experts in nutrition, exercise, and behaviour change and were based on a structured protocol. The participants were given a standard reduced-calorie diet (1200 to 1500 kcal per day), with a goal of providing no more than 30% of the calories from fat. To improve adherence, the participants were provided with sample meal plans and were given vouchers for a meal-replacement product (Slim-Fast) to be used for two meals a day during months 1 to 4 and for one meal a day thereafter. The participants were encouraged to gradually increase physical activity (brisk walking or activities of similar intensity) until they were active for at least 200 minutes each week. Behavioural skills, including self-monitoring,</p> | <p>diaries with the participants to answer questions and reconcile inconsistencies. For the purposes of analysis, each woman was then classified as having stress-only incontinence, stress-predominant incontinence (that is, at least two thirds of the total number of episodes were stress episodes), urge-only incontinence, urge-predominant incontinence (that is, at least two thirds of the total number of episodes were urge episodes), or mixed incontinence (that is at least two types were reported, but no type constituted two thirds of the total number of episodes).</p> | <p>Post: Intervention 13 (15); Control 17 (19)</p> <p>Stress incontinence (no./wk)</p> <p>Pre: Intervention 9 (11); Control 10 (10)</p> <p>Post: Intervention 4 (7); Control 7 (9)</p> <p>Urge incontinence (no./wk)</p> <p>Pre: Intervention 14 (14); Control 13 (15)</p> <p>Post: Intervention 8 (11); Control 10 (15)</p> | <p>2.1: Yes, participants not blinded</p> <p>2.2: Yes, carers and people delivering the interventions not blinded</p> <p>2.3: No information whether there were any deviations from the intended intervention</p> <p>Domain 3: Missing outcome data: Low risk</p> <p>3.1: Probably no, 5% of the intervention group and 24% in the control group were lost to follow-up or did not complete all measures</p> <p>3.2: Probably no, no evidence that the results were not biased by missing outcome data</p> <p>3.3: Probably no, missingness of the outcome was not dependent on its true value</p> <p>Domain 4: Measurement of the outcome: Low risk</p> <p>4.1: Probably no, outcomes clearly defined and some information on how they were assessed and by whom (women's self-reported following training and clinical assessment)</p> <p>4.2: Probably no, outcomes unlikely to differ between treatment arms</p> <p>4.3: No, outcome assessors were blinded</p> |

| Study details | Participants | Interventions | Methods | Outcomes | Comments |
|---------------------------------------|---|---|---------|----------|---|
| Office of Research on Women's Health. | <p>more urinary-incontinence episodes in a 7-day diary of voiding. Willing and able to monitor their food intake and physical activity for 1 week, to be able to walk unassisted for two blocks (approximately 270 m) without stopping, and to agree not to initiate new treatments for incontinence or weight reduction for the duration of the study. Previous medical therapy for incontinence or obesity did not affect eligibility.</p> <p>Exclusion criteria The use of medical therapy for incontinence or weight loss within the previous month, current urinary tract infection or four or more urinary tract infections in the previous year, a history of incontinence of neurologic or functional origin (due to factors not involving the lower urinary tract, such as chronic impairment of physical or cognitive functioning), previous surgery for incontinence or urethral surgery, major medical or genitourinary tract conditions, pregnancy or parturition in the previous 6 months,</p> | stimulus control, and problem-solving, were emphasised. | | | <p>Domain 5: Selection of the reported result: Some concerns 5.1: No, no pre-planned analysis or protocol available 5.2: No, descriptive data presented 5.3: No, data presented as expected</p> <p>Domain 6: Overall judgment of bias: Some concerns</p> <p>Other information Linked studies:</p> <p>Breyer 2018 - additional data on daytime frequency, nocturia, urinary urgency and IPSS Score ≥ 8 Huang 2009 - additional data on sexual function of the women with UI, data not relevant Myers 2012 - additional data on POP symptoms of the women with UI, data not relevant West 2011 - additional data on the 12 month maintenance programme for skill-based maintenance or motivational maintenance Wing 2010 - additional data on improvements in UI based on % weight loss. Data merged for</p> |

| Study details | Participants | Interventions | Methods | Outcomes | Comments |
|--|--|--|--|--|---|
| | type 1 or type 2 diabetes mellitus requiring medical therapy that increased the risk of hypoglycaemia, and uncontrolled hypertension. | | | | both intervention and control groups Wing 2010 - additional data on the 12 month maintenance programme for skill-based maintenance or motivational maintenance |
| <p>Full citation Subak,L.L., Whitcomb,E., Shen,H., Saxton,J., Vittinghoff,E., Brown,J.S., Weight loss: A novel and effective treatment for urinary incontinence, Journal of Urology, 174, 190-195, 2005</p> <p>Ref Id 144400</p> <p>Country/ies where the study was carried out USA</p> <p>Study type Randomised Controlled Trial</p> <p>Aim of the study To evaluated the effect of weight loss on urinary incontinence (UI) in overweight and obese women</p> | <p>Sample size n=48</p> <p>Characteristics Age (median, range): Intervention 50.5 (46-54) years; Control 57.5 (50-62) years</p> <p>BMI (median, range): Intervention 34 (32-40) kg/m²; Control 36 (32-38) kg/m²</p> <p>Parity (median, range): Intervention 1 (0-2); Control 1 (0-2)</p> <p>No. menopausal: Intervention 8 (40%); Control 13 (65%)</p> <p>Urge UI alone: Intervention 3 (12%); Control 2 (9%) Mixed, urge predominate UI: Intervention 9 (38%); Control 11 (48%)</p> <p>Stress UI: Intervention 3 (12%); Control 0 (0)</p> | <p>Interventions A 3-month intensive group based medical and behavioural weight loss program. Participants were placed on a standard low calorie liquid diet (800 kcals per day or less), encouraged to increase physical activity gradually until they were exercising 60 minutes daily, and were taught standard cognitive and behavioural skills to assist in modifying eating and exercise habits. Participants met weekly in group sessions led by a nutritionist, exercise physiologist or behavioural therapist and followed a structured protocol</p> | <p>Details Women were then randomised to either immediate (immediate intervention, 20) or delayed (wait-list control, 19) enrolment in the weight reduction program. Women in the immediate intervention group began the weight reduction program and continued for 3 months. Women in the wait-list control group had no intervention for 3 months and then entered the weight reduction program. Women were followed for 6 months after completing the weight reduction program. A 7-day voiding diary was used to measure the primary outcome of percent change in number of weekly urinary incontinent episodes. Quality of life measures included the Incontinence Impact Questionnaire (IIQ), Urogenital Distress Inventory (UDI) and Short</p> | <p>Results Data are median (IQR)</p> <p><u>Body weight</u> Pre: Intervention 99 (86-109); Control 94 (80-101) Post: Intervention 84 (71-91); Control 96 (82-102)</p> <p><u>Stress incontinent episodes/wk</u> Pre: Intervention 8 (2-13); Control 8 (3-15) Post: Intervention 0 (0-4); Control 8 (1-15)</p> <p><u>Urge incontinent episodes/wk</u> Pre: Intervention 8 (4-20); Control 7 (2-13) Post: Intervention 2 (0-10); Control 5 (3-14)</p> <p><u>IIQ Score</u> Pre: Intervention 109 (53-143); Control 99 (72-164) Post: Intervention 37 (11-86); Control 89 (56-139)</p> <p><u>UDI Score</u></p> | <p>Limitations Cochrane risk of bias (Version 2.0)</p> <p>Domain 1: Randomisation: Low risk 1.1: Yes, patients were randomly allocated to treatments 1.2: Yes, randomisation used sealed opaque consecutively numbered envelopes 1.3: Probably no, participants significantly older in the control group</p> <p>Domain 2: Deviations from intended interventions: Some risk 2.1: Yes, participants not blinded 2.2: Yes, carers and people delivering the interventions not blinded 2.3: No information whether there were any deviations from the intended intervention</p> <p>Domain 3: Missing outcome data: Low risk</p> |

| Study details | Participants | Interventions | Methods | Outcomes | Comments |
|---|---|---------------|---|--|--|
| <p>Study dates January 1999 to March 2000</p> <p>Source of funding Supported by research awards from Mount Zion Health Services, Inc. and the University of California, San Francisco Academic Senate, Committee on Research.</p> | <p>Mixed, stress predominate UI: Intervention 9 (38%); Control 10 (43%)</p> <p>Inclusion criteria Women 18 to 80 years old with BMI between 25 and 45 kg/m², urinary incontinence for at least 3 months and at least 4 incontinent episodes in a 7-day urinary diary. Prior incontinence therapies (including surgery) were not exclusions from study eligibility. Participants currently using incontinence therapy were included in the study but were asked to not change treatment during study.</p> <p>Exclusion criteria Pregnancy, urinary tract infection, significant medical condition, pelvic cancer, neurological condition possibly associated with incontinence, interstitial cystitis or potential inability to complete the study.</p> | | <p>Form 36 (SF-36).12,13 The IIQ assesses incontinence specific quality of life and the UDI quantifies incontinence symptom bother. Both measures are scored on a continuous scale (IIQ range 0 to 400, UDI range 0 to 300) with higher scores indicating greater effect on quality of life. Urodynamic evaluation included cystometrogram (sitting at 45 degrees of recline), urethral pressure profile and pressure flow voiding studies. Measurements conformed to the recommendations of the International Continence Society</p> | <p>Pre: Intervention 158 (142-192); Control 200 (128-233) Post: Intervention 104 (67-122); Control 195 (156-228)</p> <p><u>SF-36 physical component</u> Pre: Intervention 46 (42-51); Control 47 (44-51) Post: Intervention 55 (49-58); Control 47 (41-50)</p> <p><u>SF-36 mental component</u> Pre: Intervention 48 (45-50); Control 48 (45-54) Post: Intervention 48 (46-49); Control 51 (48-54)</p> | <p>3.1: Probably no, 20% of the intervention group and 13% in the control group were lost to follow-up or excluded from final analysis 3.2: Probably no, no evidence that the results were not biased by missing outcome data 3.3: Probably no, missingness of the outcome was not dependent on its true value</p> <p>Domain 4: Measurement of the outcome: Low risk 4.1: Probably no, outcomes clearly defined and some information on how they were assessed and by whom (women's self-reported and blinded clinical assessment) 4.2: Probably no, outcomes unlikely to differ between treatment arms 4.3: No, outcome assessors were blinded</p> <p>Domain 5: Selection of the reported result: Some concerns 5.1: No, no pre-panned analysis or protocol available 5.2: No, descriptive data presented 5.3: No, data presented as expected</p> |

| Study details | Participants | Interventions | Methods | Outcomes | Comments |
|--|---|--|--|--|---|
| | | | | | Domain 6: Overall judgment of bias: Some concerns |
| <p>Full citation West, D. S., Gorin, A. A., Subak, L. L., Foster, G., Bragg, C., Hecht, J., Schembri, M., Wing, R. R., Program to Reduce Incontinence by, Diet, Exercise Research, Group, A motivation-focused weight loss maintenance program is an effective alternative to a skill-based approach, International journal of obesity, 35, 259-69, 2011</p> <p>Ref Id 1118299</p> <p>Country/ies where the study was carried out USA</p> <p>Study type Randomised Controlled Trial</p> <p>Aim of the study See Subak 2009</p> <p>Study dates See Subak 2009</p> | <p>Sample size See Subak 2009</p> <p>Characteristics See Subak 2009</p> <p>Inclusion criteria See Subak 2009</p> <p>Exclusion criteria See Subak 2009</p> | <p>Interventions See Subak 2009 and Wing 2010</p> | <p>Details See Subak 2009</p> | <p>Results See Subak 2009 and Wing 2010</p> | <p>Limitations See Subak 2009</p> <p>Other information See Subak 2009</p> |

| Study details | Participants | Interventions | Methods | Outcomes | Comments |
|--|---|--|---|---|--|
| <p>Source of funding See Subak 2009</p> | | | | | |
| <p>Full citation Wing, R. R., West, D. S., Grady, D., Creasman, J. M., Richter, H. E., Myers, D., Burgio, K. L., Franklin, F., Gorin, A. A., Vittinghoff, E., Macer, J., Kusek, J. W., Subak, L. L., Program to Reduce Incontinence by, Diet, Exercise, Group, Effect of weight loss on urinary incontinence in overweight and obese women: results at 12 and 18 months, Journal of urology, 184, 1005-10, 2010</p> <p>Ref Id 1118193</p> <p>Country/ies where the study was carried out USA</p> <p>Study type Randomised controlled trial</p> <p>Aim of the study See Subak 2009</p> | <p>Sample size See Subak 2009</p> <p>Characteristics See Subak 2009</p> <p>Inclusion criteria See Subak 2009</p> <p>Exclusion criteria See Subak 2009</p> | <p>Interventions Following the initial 6 months follow up from Subak 2009, women were asked if they wished to continue with a 12 month maintenance approach. Women in the intervention group (n=226) were randomised into either a novel motivation-focused weight maintenance programme (n=113) or a skill-based maintenance program (n=113). Women in the control group, remained in the control group.</p> | <p>Details Urinary incontinence was assessed with a 7-day voiding diary which participants were trained to use. 24-hour involuntary urine loss was measured using a standardised pad test. Participants were also asked whether, compared to baseline, incontinence episodes were less frequent (yes, no, uncertain), leakage was smaller (yes, no, uncertain) and whether leakage was improved (5-point scale), and were asked to rate overall satisfaction with changes in incontinence (5-point scale).</p> | <p>Results Data given as percent change and 95% CI - Data for Intervention groups merged (skills based maintenance and motivation based maintenance)</p> <p><u>Body weight (kg) change, (n=201 intervention motivation focused and intervention skill based and n=91 control):</u> Baseline to 18 months: Intervention -5.5 (-6.7, -4.3); Control -1.6 (-3.4, 0.7)</p> <p><u>Total UI episodes/wk, (n=197 intervention motivation focused and intervention skill based and n=90 control):</u> Baseline to 18 months: Intervention -62 (-67, -55); Control -55 (-65, -43)</p> <p><u>Stress UI episodes/wk, (n=197 intervention motivation focused and intervention skill based and n=90 control):</u> Baseline to 18 months: Intervention -69 (-76, -61); Control -62 (-73, -48)</p> | <p>Limitations See Subak 2009</p> <p>Other information</p> |

| Study details | Participants | Interventions | Methods | Outcomes | Comments |
|---|--------------|---------------|---------|---|----------|
| <p>Study dates See Subak 2009</p> <p>Source of funding See Subak 2009</p> | | | | <p><u>Urge UI episodes/wk, (n=197 intervention motivation focused and intervention skill based and n=90 control):</u> Baseline to 18 months: Intervention -56 (-64, -46); Control -49 (-64, -28)</p> <p><u>24hr involuntary urine loss, (n=197 intervention motivation focused and intervention skill based and n=90 control):</u> Baseline to 18 months: Intervention -55 (-62, -46); Control -52 (-65, -35)</p> | |

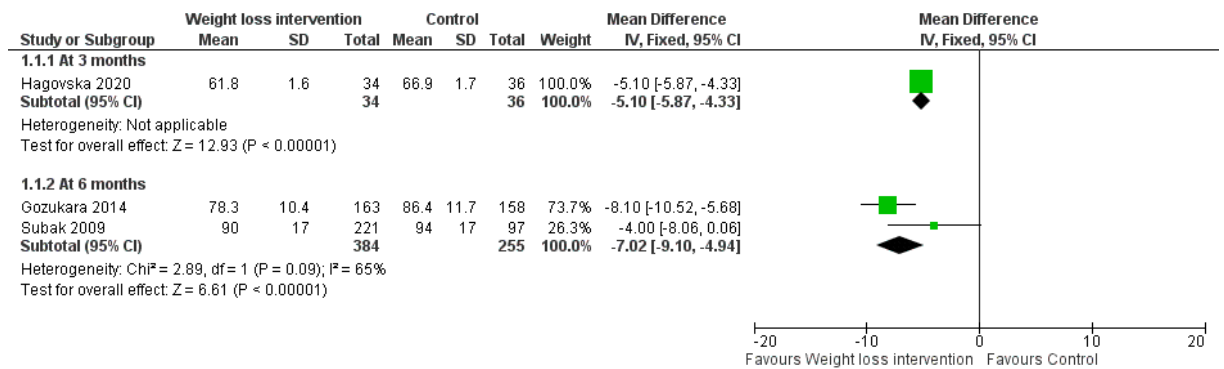
CRADI: colorectal anal distress inventory; IIQ: incontinence impact questionnaire; IPSS: International Prostate Symptom Score; PFDI: Pelvic Floor Distress Inventory; POP: pelvic organ prolapse; POPDI: pelvic organ prolapse distress inventory; SF-36: short form of the quality of life questionnaire; UDI: urinary distress inventory; UI: urinary incontinence.

Appendix E – Forest plots

Forest plots for review question: What is the effectiveness of weight loss interventions for improving symptoms of pelvic floor dysfunction?

This section includes forest plots only for outcomes that are meta-analysed. Outcomes from single studies are not presented here; the quality assessment for such outcomes is provided in the GRADE profiles in appendix F.

Figure 2: Weight loss at 3 and 6 months for women with urinary incontinence



Appendix F – GRADE tables

GRADE tables for review question: What is the effectiveness of weight loss interventions for improving symptoms of pelvic floor dysfunction?

Table 5: Clinical evidence profile for weight loss interventions vs control

| Quality assessment | | | | | | | No of patients | | Effect | | Quality | Importance |
|--|-------------------|----------------------------|--------------------------|-------------------------|---------------------------|----------------------|----------------|-------------------------|-------------------|---|----------|------------|
| No of studies | Design | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | Weight loss | Control (UI population) | Relative (95% CI) | Absolute | | |
| Body weight (kg) (follow-up 3 months; Better indicated by lower values) | | | | | | | | | | | | |
| 1 Hagovska 2020 | randomised trials | very serious ¹⁴ | no serious inconsistency | no serious indirectness | no serious imprecision | none | 53 | 56 | - | MD 5.10 kg lower (4.33 to 5.87kg lower) | LOW | CRITICAL |
| Body weight (kg) (follow-up 3 months; Better indicated by lower values) | | | | | | | | | | | | |
| 1 Subak 2005 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | serious ¹³ | none | 19 | 20 | - | Median 12kg lower. Median(IQR) Intervention 84 (71-91); Control 96 (82-102) | LOW | CRITICAL |
| Body weight (kg) (follow-up 6 months; Better indicated by lower values) | | | | | | | | | | | | |
| 2 ² | randomised trials | serious ¹ | serious ³ | no serious indirectness | serious ⁴ | none | 384 | 255 | - | MD 7.02 lower (9.1 to 4.94 lower) | VERY LOW | CRITICAL |
| Body weight (% change) (follow-up 18 months; Better indicated by lower values) | | | | | | | | | | | | |
| 1 Subak 2009 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | no serious imprecision | none | 201 | 91 | - | MD 4.03 lower (4.26 to 3.8 lower) | MODERATE | CRITICAL |
| Total UI episodes/wk (follow-up 6 months; Better indicated by lower values) | | | | | | | | | | | | |
| Subak 2009 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | very serious ⁵ | none | 214 | 90 | - | MD 4 lower (8.41 lower to 0.41 higher) | VERY LOW | CRITICAL |
| Total UI episodes/wk (% change) (follow-up 18 months; Better indicated by lower values) | | | | | | | | | | | | |

| Quality assessment | | | | | | | No of patients | | Effect | | Quality | Importance |
|---|-------------------|----------------------|--------------------------|-------------------------|---------------------------|----------------------|----------------|-------------------------|-------------------|--|----------|------------|
| No of studies | Design | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | Weight loss | Control (UI population) | Relative (95% CI) | Absolute | | |
| Subak 2009 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | no serious imprecision | none | 197 | 90 | - | MD 7.01 lower (8.26 to 5.76 lower) | MODERATE | CRITICAL |
| 24hr involuntary urine loss (% change) (follow-up 18 months; Better indicated by lower values) | | | | | | | | | | | | |
| Subak 2009 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | serious ⁶ | none | 197 | 90 | - | MD 3.51 lower (5.21 to 1.81 lower) | LOW | CRITICAL |
| PFDI-20 total (follow-up 6 months; Better indicated by lower values, scale from 0-300) | | | | | | | | | | | | |
| Gozukara 2014 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | very serious ⁷ | none | 163 | 158 | - | MD 4.8 lower (5.58 to 4.02 lower) | VERY LOW | CRITICAL |
| Stress incontinence per 3 days (follow-up 6 months; Better indicated by lower values) | | | | | | | | | | | | |
| Gozukara 2014 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | no serious imprecision | none | 163 | 158 | - | MD 3.92 lower (3.95 to 3.89 lower) | MODERATE | CRITICAL |
| Urge incontinence per 3 days (follow-up 6 months; Better indicated by lower values) | | | | | | | | | | | | |
| Gozukara 2014 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | no serious imprecision | none | 163 | 158 | - | MD 1.4 lower (1.42 to 1.38 lower) | MODERATE | CRITICAL |
| Stress incontinence episodes/wk (follow-up 3 months; Better indicated by lower values) | | | | | | | | | | | | |
| Subak 2005 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | serious ¹³ | none | 19 | 20 | - | Median 8 fewer. Median(IQR) Intervention 0 (0-4); Control 8 (1-15) | LOW | CRITICAL |
| Stress incontinence episodes/wk (follow-up 6 months; Better indicated by lower values) | | | | | | | | | | | | |
| Subak 2009 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | serious ⁸ | none | 214 | 90 | - | MD 3 lower (5.08 to 0.92 lower) | LOW | CRITICAL |
| Urge incontinence episodes/wk (follow-up 3 months; Better indicated by lower values) | | | | | | | | | | | | |

| Quality assessment | | | | | | | No of patients | | Effect | | Quality | Importance |
|---|-------------------|----------------------|--------------------------|-------------------------|----------------------------|----------------------|----------------|-------------------------|-------------------|---|----------|------------|
| No of studies | Design | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | Weight loss | Control (UI population) | Relative (95% CI) | Absolute | | |
| 1 Subak 2005 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | serious ¹³ | none | 19 | 20 | - | Median 3 fewer. Median(IQR) : Intervention 2 (0-10); Control 5 (3-14) | LOW | CRITICAL |
| Urge incontinence episodes/wk (follow-up 6 months; Better indicated by lower values) | | | | | | | | | | | | |
| Subak 2009 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | very serious ¹⁰ | none | 214 | 90 | - | MD 2 lower (5.43 lower to 1.43 higher) | VERY LOW | CRITICAL |
| IIQ score (follow-up 3 months; Better indicated by lower values, scale from 0-400) | | | | | | | | | | | | |
| Subak 2005 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | serious ¹³ | none | 19 | 20 | - | Median 52 lower. Median(IQR) Intervention 37 (11-86); Control 89 (56-139) | LOW | CRITICAL |
| UDI score (follow-up 3 months; Better indicated by lower values, scale from 0-300) | | | | | | | | | | | | |
| Subak 2005 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | serious ¹³ | none | 19 | 20 | - | Median 91 lower. Median(IQR) Intervention 104 (67-122); Control 195 (156-228) | LOW | CRITICAL |
| SF-36 Physical component (follow-up 3 months; Better indicated by higher values, scale from 0-100) | | | | | | | | | | | | |
| Subak 2005 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | serious ¹³ | none | 19 | 20 | - | Median 8 higher. Median(IQR) : Intervention 55 (49-58); Control 47 (41-50) | LOW | IMPORTANT |
| SF-36 Mental component (follow-up 3 months; Better indicated by higher values, scale from 0-100) | | | | | | | | | | | | |
| Subak 2005 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | serious ¹³ | none | 19 | 20 | - | Median 3 lower. Median(IQR) : Intervention 48 (46-49); Control 51 (48-54) | LOW | IMPORTANT |

| Quality assessment | | | | | | | No of patients | | Effect | | Quality | Importance |
|---|-------------------|----------------------|--------------------------|-------------------------|----------------------------|----------------------|----------------|-------------------------|------------------------|---|----------|------------|
| No of studies | Design | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | Weight loss | Control (UI population) | Relative (95% CI) | Absolute | | |
| Daytime frequency (follow-up 6 months) | | | | | | | | | | | | |
| Subak 2009 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | serious ¹¹ | none | 93/214 (43.5%) | 47/90 (52.2%) | RR 0.83 (0.65 to 1.07) | 89 fewer per 1000 (from 183 fewer to 37 more) | LOW | CRITICAL |
| Nocturia (follow-up 6 months) | | | | | | | | | | | | |
| Subak 2009 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | serious ¹¹ | none | 83/214 (38.8%) | 47/90 (52.2%) | RR 0.74 (0.57 to 0.96) | 136 fewer per 1000 (from 21 fewer to 225 fewer) | LOW | CRITICAL |
| Urinary urgency (follow-up 6 months) | | | | | | | | | | | | |
| Subak 2009 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | serious ¹¹ | none | 89/214 (41.6%) | 46/90 (51.1%) | RR 0.81 (0.63 to 1.05) | 97 fewer per 1000 (from 189 fewer to 26 more) | LOW | CRITICAL |
| IPSS score >8 (follow-up 6 months) | | | | | | | | | | | | |
| Subak 2009 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | serious ¹¹ | none | 76/214 (35.5%) | 40/90 (44.4%) | RR 0.8 (0.6 to 1.07) | 89 fewer per 1000 (from 178 fewer to 31 more) | LOW | CRITICAL |
| POP symptoms – number of people reporting cure (follow-up 6 months) | | | | | | | | | | | | |
| Subak 2009 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | very serious ¹² | none | 73/107 (68.2%) | 35/49 (71.4%) | RR 0.96 (0.77 to 1.19) | 29 fewer per 1000 (from 164 fewer to 136 more) | VERY LOW | CRITICAL |
| POP symptoms – number of people reporting improved bother or cure (follow-up 6 months) | | | | | | | | | | | | |
| Subak 2009 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | very serious ¹² | none | 82/107 (76.6%) | 38/49 (77.6%) | RR 0.99 (0.82 to 1.19) | 8 fewer per 1000 (from 140 fewer to 147 more) | VERY LOW | CRITICAL |
| POP symptoms – number of people reporting new symptom (follow-up 6 months) | | | | | | | | | | | | |
| Subak 2009 | randomised trials | serious ¹ | no serious inconsistency | no serious indirectness | very serious ¹² | none | 35/214 (16.4%) | 20/94 (21.3%) | RR 0.77 (0.47 to 1.26) | 49 fewer per 1000 (from 113 fewer to 55 more) | VERY LOW | CRITICAL |
| OAB symptoms – OAB-Q symptom score (follow up 3 months) | | | | | | | | | | | | |

| Quality assessment | | | | | | | No of patients | | Effect | | Quality | Importance |
|--|-------------------|----------------------------|--------------------------|-------------------------|------------------------|----------------------|----------------|-------------------------|-------------------|--------------------------------------|----------|------------|
| No of studies | Design | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | Weight loss | Control (UI population) | Relative (95% CI) | Absolute | | |
| 1 Hagovska 2020 | randomised trials | very serious ¹⁴ | no serious inconsistency | no serious indirectness | no serious imprecision | none | 34 | 36 | - | MD 10.9 lower (11.53 to 10.27 lower) | LOW | CRITICAL |
| OAB symptoms – OAB-Q quality of life score (follow up 3 months) | | | | | | | | | | | | |
| 1 Hagovska 2020 | randomised trials | very serious ¹⁴ | no serious inconsistency | no serious indirectness | no serious imprecision | none | 34 | 36 | - | MD 7 higher (6.58 to 7.42 higher) | LOW | CRITICAL |
| BMI (follow up 3 months) | | | | | | | | | | | | |
| 1 Hagovska 2020 | randomised trials | very serious ¹⁴ | no serious inconsistency | no serious indirectness | serious ¹⁵ | none | 34 | 36 | - | MD 1.8 lower (2.11 to 1.49) | VERY LOW | CRITICAL |

CI: confidence interval; BMI: body mass; IIQ; Incontinence Impact Questionnaire; MID: minimal important difference; MD: mean difference; OAB: overactive bladder; PISQ: Pelvic organ prolapse/urinary incontinence sexual questionnaire; POPQ: Pelvic organ prolapse quantification system; POP-SS: Pelvic organ prolapse symptom score; POP: Pelvic organ; QOL: quality of life; SD: standard deviation SF-36: short form 36; UI: urinary incontinence; UDI: Urinary distress index

1 Serious risk of bias in the evidence contributing to the outcomes as per RoB 2

2 Subak 2009 and Gozukara 2014

3 Serious heterogeneity unexplained by subgroup analysis

4 95% CI crosses 1 MID (0.5x control group SD, MID=7.2)

5 95% CI crosses 2 MIDs (0.5 x control group SD, MID = 9.5)

6 95% CI crosses 1 MID (0.5x control group SD, MID=3.9)

7 95% CI crosses 2 MIDs (50 points - Ma 2019)

8 95% CI crosses 1 MID (0.5x control group SD, MID=4.5)

9 95% CI crosses 1 MID (0.5x control group SD, MID=1.6)

10 95% CI crosses 2 MIDs (0.5x control group SD, MID=7.5)

11 95% CI crosses 1 MID (0.8 or 1.25)

12 95% CI crosses 2 MIDs (0.8 and 1.25)

13 Subjective assessment of imprecision

14 Very serious risk of bias in the evidence contributing to the outcomes as per RoB 2

15 95% CI crosses 1 MID (0.5x control group SD, MID=2.1)

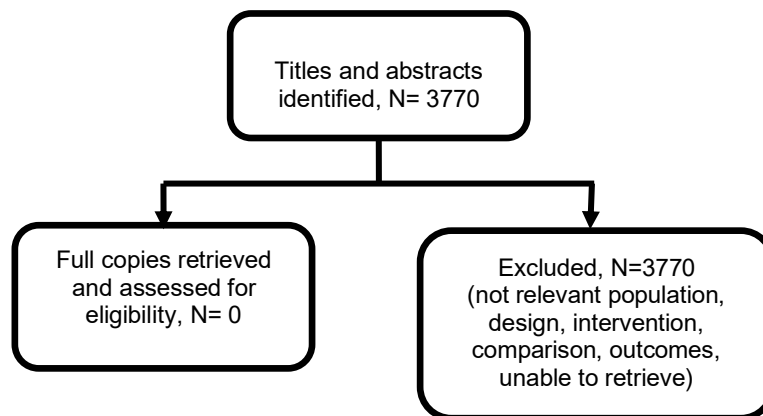
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Appendix G – Economic evidence study selection

Economic evidence study selection for review question: What is the effectiveness of weight loss interventions for improving symptoms of pelvic floor dysfunction?

No economic evidence was identified which was applicable to this review question.

Figure 3: Study selection flow chart



Appendix H – Economic evidence tables

Economic evidence tables for review question: What is the effectiveness of weight loss interventions for improving symptoms of pelvic floor dysfunction?

No evidence was identified which was applicable to this review question.

Appendix I – Economic evidence profiles

Economic evidence profiles for review question: What is the effectiveness of weight loss interventions for improving symptoms of pelvic floor dysfunction?

No economic evidence was identified which was applicable to this review question.

Appendix J – Economic analysis

Economic evidence analysis for review question: What is the effectiveness of weight loss interventions for improving symptoms of pelvic floor dysfunction?

No economic analysis was conducted for this review question.

Appendix K – Excluded studies

Excluded studies for review question: What is the effectiveness of weight loss interventions for improving symptoms of pelvic floor dysfunction?

Clinical studies

Table 6: Excluded studies and reasons for their exclusion

| Study | Reason for exclusion |
|---|---|
| Allahdin,S., Oo,N., An overview of treatment of overactive bladder syndrome in women, <i>Journal of Obstetrics and Gynaecology</i> , 32, 217-221, 2012 | Literature review |
| Aversa, A., Bruzziches, R., Francomano, D., Greco, E. A., Violi, F., Lenzi, A., Donini, L. M., Weight Loss by Multidisciplinary Intervention Improves Endothelial and Sexual Function in Obese Fertile Women, <i>Journal of Sexual Medicine</i> , 10, 1024-1033, 2013 | Population does not have PFD |
| Ballard,A.C., Richter,H.E., Impact of obesity and weight loss on urinary and bowel incontinence symptoms in women, <i>Sexuality, Reproduction and Menopause</i> , 9, S1-S7, 2011 | Literature review |
| Brandt, L. J., Prather, C. M., Quigley, E. M. M., Schiller, L. R., Schoenfeld, P., Talley, N. J., Systematic review on the management of chronic constipation in North America, <i>American Journal of Gastroenterology</i> , 100, S5-S22, 2005 | Systematic review - included studies checked for relevance and none found |
| Cameron,A.P., Haraway,A.M., The treatment of female stress urinary incontinence: An evidenced-based review, <i>Open Access Journal of Urology</i> , 3, 109-120, 2011 | Literature review |
| Cheskin, L. J., Burnett, A. L., A behavioural weight-loss programme was better than an education programme for urinary incontinence in overweight and obese women, <i>Evidence-Based Medicine</i> , 14, 118, 2009 | Abstract |
| de Oliveira, M. C. E., de Oliveira de Lima, V. C., Pegado, R., Silva-Filho, E. M., Fayh, A. P. T., Micussi, M. T., Comparison of pelvic floor muscle training isolated and associated with weight loss: a randomized controlled trial, <i>Archives of gynecology and obstetrics</i> , 300, 1343-1351, 2019 | No usable results |
| Gomelsky,A., Dmochowski,R.R., Treatment of mixed urinary incontinence in women, <i>Current Opinion in Obstetrics and Gynecology</i> , 23, 371-375, 2011 | Literature review |
| Greer,W.J., Richter,H.E., Bartolucci,A.A., Burgio,K.L., Obesity and pelvic floor disorders: A systematic review, <i>Obstetrics and Gynecology</i> , 112, 341-349, 2008 | Systematic review - included studies checked for relevance |
| Huang, A. J., Stewart, A. L., Hernandez, A. L., Shen, H., Subak, L. L., Program to Reduce Incontinence by, Diet, Exercise,, Sexual function among overweight and obese women with urinary incontinence in a randomized controlled trial of an intensive behavioral weight loss intervention, <i>Journal of urology</i> , 181, 2235-42, 2009 | Linked to Subak 2009, no useable data |
| Hill, J. E., Christian, D., Shaw, K., Clegg, A., Weight loss interventions as an option for a lifestyle treatment in urinary incontinence, <i>British Journal of Community Nursing</i> Br J Community Nurs, 25, 616-619, 2020 | Incorrect study design: commentary on a review. |
| Imamura, M., Williams, K., Wells, M., McGrother, C., Lifestyle interventions for the treatment of urinary incontinence in adults, <i>Cochrane Database of Systematic Reviews</i> , 2015 | Systematic review - included studies checked for relevance |
| Kolotkin, R. L., Zunker, C., Ostbye, T., Sexual functioning and obesity: A review, <i>Obesity</i> , 20, 2325-2333, 2012 | Literature review |
| Moore,K.N., Saltmarche,A., Query,B., Urinary incontinence. Non-surgical management by family physicians, <i>Canadian Family Physician</i> , 49, 602-610, 2003 | Literature review |

| Study | Reason for exclusion |
|---|--|
| Norton, C., Whitehead, W. E., Bliss, D. Z., Harari, D., Lang, J., Management of fecal incontinence in adults, <i>Neurourology and urodynamics</i> , 29, 199-206, 2010 | Literature review |
| Nct., Effect of Pelvic Floor Muscle Training on Urinary Incontinence Reports in Obese Women Undergoing a Low Calorie Diet Prior to Bariatric Surgery, https://clinicaltrials.gov/show/NCT04159467 , 2019 | Incorrect intervention/comparison |
| Olivera, C. K., Meriwether, K., El-Nashar, S., Grimes, C. L., Chen, C. C. G., Orejuela, F., Antosh, D., Gleason, J., Kim-Fine, S., Wheeler, T., McFadden, B., Balk, E. M., Murphy, M., Nonantimuscarinic treatment for overactive bladder: A systematic review, <i>American Journal of Obstetrics and Gynecology</i> , 215, 34-57, 2016 | Systematic Review - all included studies checked for relevance |
| Pomian, A., Lisik, W., Kosieradzki, M., Barcz, E., Obesity and Pelvic Floor Disorders: A Review of the Literature, <i>Medical Science Monitor</i> , 22, 1880-6, 2016 | Literature review |
| Qaseem, A., Dallas, P., Forciea, M. A., Starkey, M., Denberg, T. D., Shekelle, P., Clinical Guidelines Committee of the American College of Physicians, Nonsurgical management of urinary incontinence in women: a clinical practice guideline from the American College of Physicians, <i>Annals of internal medicine</i> , 161, 429-440, 2014 | Guidance based on a systematic review - included studies checked for relevance |
| Schumacher, L., Wing, R., Thomas, J. G., Pavlovic, J., Digre, K., Farris, S., Steffen, K., Sarwer, D., Bond, D., Does sexual functioning improve with migraine improvements and/or weight loss?-A post hoc analysis in the Women's Health and Migraine (WHAM) trial, <i>Obesity Science and Practice</i> ., 2020 | Incorrect population, incorrect comparison group |
| Tantawy, S. A., Kamel, D. M., Abdelbasset, W. K., Elgohary, H. M., Effects of a proposed physical activity and diet control to manage constipation in middle-aged obese women, <i>Diabetes, Metabolic Syndrome and Obesity Targets and Therapy</i> Diabetes Metab Syndr Obes, 10, 513-519, 2017 | Constipation is not relevant |
| Vissers, D., Neels, H., Vermandel, A., De Wachter, S., Tjalma, W. A., Wyndaele, J. J., Taeymans, J., The effect of non-surgical weight loss interventions on urinary incontinence in overweight women: a systematic review and meta-analysis, <i>Obesity Reviews</i> , 15, 610-7, 2014 | Systematic review - included studies checked for relevance |
| Wing, R. R., Creasman, J. M., West, D. S., Richter, H. E., Myers, D., Burgio, K. L., Franklin, F., Gorin, A. A., Vittinghoff, E., Macer, J., Kusek, J. W., Subak, L. L., Program to Reduce Incontinence by, Diet, Exercise,, Improving urinary incontinence in overweight and obese women through modest weight loss, <i>Obstetrics & Gynecology</i> Obstet Gynecol, 116, 284-92, 2010 | No relevant data |
| Yamada, B.S., Govier, F.E., Does weight loss improve urinary incontinence in overweight and obese women?, <i>Nature Clinical Practice Urology</i> , 3, 16-17, 2006 | Commentary paper |
| Yazdany, T., Jakus-Waldman, S., Jeppson, P. C., Schimpf, M. O., Yurteri-Kaplan, L. A., Ferzandi, T. R., Weber-LeBrun, E., Knoepp, L., Mamik, M. M., Viswanathan, M., Ward, R. M., American Urogynecologic Society, American Urogynecologic Society Systematic Review: The Impact of Weight Loss Intervention on Lower Urinary Tract Symptoms and Urinary Incontinence in Overweight and Obese Women [Erratum 2020; 26(&): 466], <i>Female Pelvic Medicine & Reconstructive Surgery</i> Female pelvic med, 26, 16-29, 2020 | Review, checked for references |

Economic studies

No economic evidence was identified for this review.

Appendix L – Research recommendations

Research recommendations for review question: What is the effectiveness of weight loss interventions for improving symptoms of pelvic floor dysfunction?

Research question

Can weight loss reduce symptoms of pelvic floor dysfunction in women who are overweight or who have obesity?

Why this is important

In women with a BMI >30, weight loss is effective at improving urinary incontinence and urgency. It is unclear as to the role of weight loss in improving other symptoms (and signs) of pelvic floor dysfunction (PFD), such as prolapse. If weight loss were an effective intervention, some women may be able to avoid surgery and other invasive interventions.

Table 7: Research recommendation rationale

| Research question | Does weight loss reduce symptoms of PFD in women who are overweight or have obesity? |
|---|---|
| Why is this needed | |
| Importance to ‘patients’ or the population | Weight loss is often suggested to people with pelvic floor dysfunction. However, there is very limited evidence to guide whether weight loss is associated with symptomatic improvement for Pelvic Organ Prolapse (POP) and whether this advice would benefit particular groups of individuals. Without this information, people may modify their weight in a manner which serves no useful purpose for the management of pelvic floor dysfunction. |
| Relevance to NICE guidance | The relative absence of evidence regarding this topic currently restricts NICE guidance from making recommendations regarding weight loss for POP in pelvic floor dysfunction. The outcome of this research would allow such recommendations to be developed and become part of NICE guidance. |
| Relevance to the NHS | Weight loss is an intervention with relatively low cost and may reduce the need for interventions with higher cost impacts on the NHS. It may be that the recommendations could be combined with existing advice. |
| National priorities | Being overweight is a key predictor of ill health is a key national priority. |
| Current evidence base | There is current evidence regarding weight loss for PFD, but limited evidence regarding its effectiveness for POP, or for whether weight loss advice can be followed by all groups of individuals (for example those with comorbid psychological issues may struggle with such advice). |
| Equality | Can weight loss advice be followed by all groups of individuals (for example those with co-morbid psychological issues, those with learning disabilities)? |
| Feasibility | Can the appropriate weight loss advice be routinely offered as part of primary and secondary care consultations regarding PFD? Or does it require extra training/resources? |
| Other comments | None |

PFD, pelvic floor dysfunction; POP, pelvic organ prolapse

Table 8: Research recommendation modified PICO table

| Criterion | Explanation |
|-------------------|--|
| Population | Women with a BMI of 25-30 and women with a BMI >30 |

| Criterion | Explanation |
|-------------------------------|---|
| Intervention | Weight loss program |
| Comparator | Structured education program |
| Outcomes | <ul style="list-style-type: none"> • Weight loss • POP symptoms: <ul style="list-style-type: none"> ○ change in POP-Q ○ change in other symptoms of pelvic floor dysfunction • Anxiety and depression |
| Study design | RCT |
| Timeframe | 2-5 years |
| Additional information | Include analysis of any predictors of the effectiveness such as psychological conditions. |

BMI, body mass index; POP, pelvic organ prolapse; POP-Q, pelvic organ prolapse quantification system; RCT, randomised controlled trial