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Nutritional interventions for survivors of childhood cancer

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Publication Details
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Abstract

**Background**  Childhood cancer survivors are at a higher risk of developing health conditions such as osteoporosis, and cardiovascular disease than their peers. Health-promoting behaviour, such as consuming a healthy diet, could lessen the impact of these chronic issues, yet the prevalence rate of health-protecting behaviour amongst survivors of childhood cancer is similar to that of the general population. Targeted nutritional interventions may prevent or reduce the incidence of these chronic diseases. **Objectives** The primary aim of this review was to assess the efficacy of a range of nutritional interventions designed to improve the nutritional intake of childhood cancer survivors, as compared to a control group of childhood cancer survivors who did not receive the intervention. Secondary objectives were to assess metabolic and cardiovascular risk factors, measures of weight and body fat distribution, behavioural change, changes in knowledge regarding disease risk and nutritional intake, participants’ views of the intervention, measures of health status and quality of life, measures of harm associated with the process or outcomes of the intervention, and cost-effectiveness of the intervention. **Search methods** We searched the electronic databases of the Cochrane Central Register of Controlled Trials (CENTRAL; 2013, Issue 3), MEDLINE/PubMed (from 1945 to April 2013), and Embase/Ovid (from 1980 to April 2013). We ran the search again in August 2015; we have not yet fully assessed these results, but we have identified one ongoing trial. We conducted additional searching of ongoing trial registers - the International Standard Randomised Controlled Trial Number register and the National Institutes of Health register (both screened in the first half of 2013) - reference lists of relevant articles and reviews, and conference proceedings of the International Society for Paediatric Oncology and the International Conference on Long-Term Complications of Treatment of Children and Adolescents for Cancer (both 2008 to 2012). **Selection criteria** We included all randomised controlled trials (RCTs) that compared the effects of a nutritional intervention with a control group which did not receive the intervention in this review. Participants were childhood cancer survivors of any age, diagnosed with any type of cancer when less than 18 years of age. Participating childhood cancer survivors had completed their treatment with curative intent prior to the intervention. **Data collection and analysis** Two review authors independently selected and extracted data from each identified study, using a standardised form. We assessed the validity of each identified study using the criteria outlined in the Cochrane Handbook for Systematic Reviews of Interventions. We used the GRADE criteria to assess the quality of each trial. **Main results** Three RCTs were eligible for review. A total of 616 participants were included in the analysis. One study included participants who had been treated for acute lymphoblastic leukaemia (ALL) (275 participants). Two studies included participants who had all forms of paediatric malignancies (266 and 75 participants). All participants were less than 21 years of age at study entry. The follow-up ranged from one month to 36 months from the initial assessment. All intended outcomes were not evaluated by each included study. All studies looked at different interventions, and so we were unable to pool results. We could not rule out the presence of bias in any of the studies.

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Nutritional interventions for survivors of childhood cancer

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A B S T R A C T

Background
Childhood cancer survivors are at a higher risk of developing health conditions such as osteoporosis, and cardiovascular disease than their peers. Health-promoting behaviour, such as consuming a healthy diet, could lessen the impact of these chronic issues, yet the prevalence rate of health-protecting behaviour amongst survivors of childhood cancer is similar to that of the general population. Targeted nutritional interventions may prevent or reduce the incidence of these chronic diseases.

Objectives
The primary aim of this review was to assess the efficacy of a range of nutritional interventions designed to improve the nutritional intake of childhood cancer survivors, as compared to a control group of childhood cancer survivors who did not receive the intervention. Secondary objectives were to assess metabolic and cardiovascular risk factors, measures of weight and body fat distribution, behavioural change, changes in knowledge regarding disease risk and nutritional intake, participants’ views of the intervention, measures of health status and quality of life, measures of harm associated with the process or outcomes of the intervention, and cost-effectiveness of the intervention.

Search methods
We searched the electronic databases of the Cochrane Central Register of Controlled Trials (CENTRAL; 2013, Issue 3), MEDLINE/PubMed (from 1945 to April 2013), and Embase/Ovid (from 1980 to April 2013). We ran the search again in August 2015; we have not yet fully assessed these results, but we have identified one ongoing trial. We conducted additional searching of ongoing trial registers - the International Standard Randomised Controlled Trial Number register and the National Institutes of Health register (both screened in the first half of 2013) - reference lists of relevant articles and reviews, and conference proceedings of the International Society for Paediatric Oncology and the International Conference on Long-Term Complications of Treatment of Children and Adolescents for Cancer (both 2008 to 2012).

Selection criteria
We included all randomised controlled trials (RCTs) that compared the effects of a nutritional intervention with a control group which did not receive the intervention in this review. Participants were childhood cancer survivors of any age, diagnosed with any type of cancer when less than 18 years of age. Participating childhood cancer survivors had completed their treatment with curative intent prior to the intervention.
Data collection and analysis

Two review authors independently selected and extracted data from each identified study, using a standardised form. We assessed the validity of each identified study using the criteria outlined in the Cochrane Handbook for Systematic Reviews of Interventions. We used the GRADE criteria to assess the quality of each trial.

Main results

Three RCTs were eligible for review. A total of 616 participants were included in the analysis. One study included participants who had been treated for acute lymphoblastic leukaemia (ALL) (275 participants). Two studies included participants who had all forms of paediatric malignancies (266 and 75 participants). All participants were less than 21 years of age at study entry. The follow-up ranged from one month to 36 months from the initial assessment. All intended outcomes were not evaluated by each included study. All studies looked at different interventions, and so we were unable to pool results. We could not rule out the presence of bias in any of the studies.

There was no clear evidence of a difference in calcium intake at one month between those who received the single, half-day, group-based education that focused on bone health, and those who received standard care (mean difference (MD) 111.60, 95% confidence interval (CI) -258.97 to 482.17; P = 0.56, low quality evidence). A regression analysis, adjusting for baseline calcium intake and changes in knowledge and self-efficacy, showed a significantly greater calcium intake for the intervention as compared with the control group at the one-month follow-up (beta coefficient 4.92, 95% CI 0.33 to 9.52; P = 0.04). There was statistically significant higher, self-reported milk consumption (MD 0.43, 95% CI 0.07 to 0.79; P = 0.02, low quality evidence), number of days on calcium supplementation (MD 11.42, 95% CI 7.11 to 15.73; P < 0.00001, low quality evidence), and use of any calcium supplementation (risk ratio (RR) 3.35, 95% CI 1.86 to 6.04; P < 0.0001, low quality evidence), with those who received this single, face-to-face, group-based, health behaviour session.

There was no clear evidence of a difference in bone density Z-scores measured with a dual-energy X-ray absorptiometry (DEXA) scan at 36 months follow-up (MD -0.05, 95% CI -0.26 to 0.16; P = 0.64, moderate quality evidence) between those who received calcium and vitamin D supplementation combined with nutrition education and those who received nutrition education alone. There was also no clear evidence of a difference in bone mineral density between the intervention and the control group at the 12-month (median difference -0.17, P = 0.99) and 24-month follow-up (median difference -0.04, P = 0.54).

A single multi-component health behaviour change intervention, focusing on general healthy eating principles, with two telephone follow-ups brought about a 0.17 lower score on the four-point Likert scale of self-reported junk food intake compared with the control group (MD -0.17, 95% CI -0.33 to -0.01; P = 0.04, low quality evidence); this result was statistically significant. There was no clear evidence of a difference between the groups in the self-reported use of nutrition as a health protective behaviour (MD -0.05, 95% CI -0.24 to 0.14; P = 0.60, low quality evidence).

Authors’ conclusions

Due to a paucity of studies, and the heterogeneity of the studies included in this review, we are unable to draw conclusions regarding the effectiveness of nutritional interventions for use with childhood cancer survivors. Although there is low quality evidence for the improvement in health behaviours using health behaviour change interventions, there remains no evidence as to whether this translates into an improvement in dietary intake. There was also no evidence that the studies reduced the risk of cardiovascular and metabolic disorders in childhood cancer survivors, although no evidence of effect is not the same as evidence of no effect. This review highlights the need for further well designed trials to be implemented in this population.

Plain Language Summary

Nutritional interventions for survivors of childhood cancer

Background

Survivors of childhood cancer are at a higher risk of chronic health conditions such as, osteoporosis, metabolic syndrome (including obesity and type II diabetes), and cardiovascular disease. These diseases have the potential to be reduced or prevented with targeted nutritional interventions.

Objective
This review looks at three randomised controlled trials that studied the effects of interventions designed to improve the dietary intake of children who have completed treatment for cancer.

**Study Characteristics**

The three studies included 616 participants who had completed their therapy for childhood cancer. All of the participants were less than 21 years of age at study entry. The interventions ranged from the promotion of health behaviours to vitamin and mineral supplementation. The follow-up ranged from one month to 36 months from the initial assessment.

**Key results**

There was low quality evidence that those who received a health behaviour intervention decreased their self-reported intake of “junk food”. They also increased their intake of dairy foods, as well as increasing their calcium supplementation. The interventions did not appear to translate to an improvement in their dietary intake, body composition, or bone mineral density.

**Quality of the evidence**

The results from this review do not provide enough evidence regarding the effectiveness of nutritional interventions for childhood cancer survivors. There was low quality evidence overall. Further well designed research is needed in this area.

**BACKGROUND**

**Description of the condition**

In the last thirty years, detection and treatment methods for childhood cancer have improved to such an extent that up to 80% of paediatric patients now survive their cancer (Cox 2009; Jemal 2009). This has resulted in a growing number of child cancer survivors and an increased clinical and research interest in the survivorship issues as a consequence of treatment, in particular treatment-related morbidity and quality of life (Cox 2009). Childhood cancer survivors have a relative risk of developing a chronic condition of 3.3 and a relative risk of a severe or life-threatening condition of 8.2 when compared with their siblings (Oeffinger 2006). Female sex and older age at diagnosis are independent risk factors for developing chronic conditions (Oeffinger 2009). These chronic health conditions include (but are not limited to) secondary cancers, endocrine disorders, renal dysfunction, and severe musculoskeletal problems (Dickerman 2007; Diller 2009; Ness 2007; Oeffinger 2006). However, it may be many years before patients display these conditions which tend to worsen over time (Oeffinger 2006).

There is now much focus in the literature on the importance of long-term monitoring of these patients (Friedman 2006; Hudson 2009; Landier 2006), and increasing recognition of the need for both secondary and tertiary interventions that may lessen the burden of these chronic conditions (Oeffinger 2009; Steinberger 2012; Stolley 2010). It may be possible to reduce the incidence of these chronic conditions with focused prevention strategies (Nathan 2009; Oeffinger 2006), aiming for quality of life similar to peers (Skinner 2006). Specific chronic health conditions of long-term survivors that have the potential to be managed by lifestyle factors include osteoporosis, metabolic syndrome, endocrine disorders, and cardiovascular disease (Nathan 2009). An individual’s risk of these conditions varies depending on factors such as disease and treatment type, age, and sex. For example, survivors of acute lymphoblastic leukaemia (ALL) who were treated with radiotherapy are at a greater risk of obesity, whereas those who received treatment for brain tumours are at risk of inadequate growth hormone (Hudson 2009). Those who received chemotherapy agents such as anthracycline are at risk of cardiovascular disease (Mulrooney 2009).

**Description of the intervention**

Despite the fact that health-promoting behaviour, such as consuming a healthy diet or maintaining adequate physical activity, could lessen the impact of these chronic issues (Stolley 2009), the prevalence of health-protecting behaviour in adults who have survived childhood cancer is similar to that of the general population (Mulhern 1995; Nathan 2009). There is a strong association in the general population between inadequate physical activity combined with a diet high in saturated fat and sugar and low in fruit and vegetable intake, and symptoms associated with the metabolic syndrome (Pereira 2009). This is of concern, since many adult survivors of childhood cancer do not meet guidelines for fruit and vegetable intake, consume excessive fat, and have an inadequate calcium intake (Demark-Wahnefried 2005; Robien 2008). These
poor eating habits appear to be manifesting themselves early af-
fter treatment completion. Young childhood cancer survivors have 
been shown to have an excessive energy intake and an inadequa-
te calcium and folate intake (Cohen 2012). Long-term survivors re-
port barriers to consuming a healthy diet that include taste pre-
fences for higher fat foods and the lack of availability of healthier 
foods (Arroyave 2008). They may also be unaware of their risk of 
chronic disease (Nathan 2009), lessening the motivation to change 
their lifestyle. As childhood cancer survivors are already at a higher 
risk of long-term metabolic complications as a result of their can-
ter therapy, poor nutritional intake may be exacerbating this risk. 
Interventions may need to be age-specific and differ between the 
older and younger childhood cancer survivor cohorts. Interven-
tions may also need to target specific conditions and high risk 
groups or may target the general paediatric population. For ex-
ample, childhood cancer survivors treated for ALL using cranial ir-
radiation are at a higher risk for obesity and subsequently metabolic 
syndrome (Oeffinger 2008), and therefore, they could be targeted 
with specific nutritional interventions to reduce obesity rates. In 
contrast, patients treated with anthracycline are at risk of car-
diovascular sequelae (Oeffinger 2008), and therefore, interven-
tions may target not only weight reduction but also aim to reduce 
cardiovascular risk (Siviero-Miachon 2008). Strategies to manage 
these chronic conditions may involve prevention interventions for 
younger cancer survivors or treatment interventions for older can-
cer survivors. Due to these variations in risk, a “one-size fits all” 
approach may not be indicated.

How the intervention might work

There is clear evidence that lifestyle changes, including improved 
diet and physical activity, are effective in the prevention or reduc-
tion of metabolic and cardiovascular risk factors in the general 
adult population (Lakka 2007). A range of nutritional interven-
tions have been reported to be effective in preventing or reduc-
ing risk factors associated with the metabolic syndrome. These 
include: low glycaemic index/high protein diets, increased fruit, 
vegetable and fibre intake, reduced salt diets and a Mediterrane-
style diet (Brunner 2009; Tota-Maharaj 2010). A recent Cochrane 
review assessing nutritional interventions for reducing or prevent-
ing cardiovascular risk found that interventions were more likely 
to be effective in participants who were told of their higher risk of 
disease (Brunner 2009). In the general paediatric population, little research has focused on 
the prevention of metabolic syndrome. Rather, there is a focus on 
prevention and treatment of childhood obesity. The literature sug-
gests that family-targeted behavioural lifestyle interventions, us-
ing a combination of nutrition, physical activity, and behavioural 
components are effective for bringing about change in overweight 
children (Oude Luttikhuis 2009). There does not appear to be 
research focusing on the efficacy of specific types of nutritional 
interventions. As the mechanisms for the increased incidence of 
these chronic diseases may be different in the general population to 
the oncology population, the results and recommendations from 
these studies may not be able to be extrapolated to childhood can-
cer survivors. Interventions focusing on older and adult survivors 
of childhood cancer may not be appropriate for the younger sur-

Why it is important to do this review

As this is a new area of study, there are minimal data in the lit-

erature with regard to the most effective nutritional interventions 
available to reduce the incidence of chronic disease after childhood 
cancer, despite the ongoing focus on long-term follow-up of these 
patients. The purpose of this Cochrane review was to assess the 
literature regarding nutritional interventions developed for child-
hood cancer survivors, to facilitate the production of best-evidence 
management guidelines.

OBJECTIVES

The primary aim of this review was to assess the efficacy of a range 
of nutritional interventions designed to improve the nutritional 
intake of childhood cancer survivors, as compared to a control 
group of childhood cancer survivors who did not receive the in-
tervention. Secondary objectives were to assess metabolic and car-
diovascular risk factors, measures of weight and body fat distribu-
tion, behavioural change, changes in knowledge regarding disease 
risk and nutritional intake, participants' views of the intervention, 
measures of health status and quality of life, measures of harm 
associated with the process or outcomes of the intervention, and 
cost- effectiveness of the intervention.

METHODS

Criteria for considering studies for this review

Types of studies
We included all randomised controlled trials (RCTs) that studied 
the effects of nutritional interventions in this review. There was no 
limit to length of the intervention, type of intervention, or length 
of follow-up.
**Types of participants**

Studies that involved childhood cancer survivors of any age, who were diagnosed with any type of cancer when less than 18 years of age were eligible for the review. Participating childhood cancer survivors had completed their treatment with curative intent prior to the intervention. We also included studies including parents and/or carers of this participant group if the parents/caregivers were involved in the intervention or reported on the participant outcomes. Treatment included chemotherapy and/or radiotherapy. We excluded studies which included participants with a co-morbidity that may have affected eating, such as autism (Emond 2010), developmental delay (Kuhn 2004), and Down’s syndrome (Lewis 2004).

**Types of interventions**

**Strategies**

We included interventions that included educational and counselling strategies, health promotion or behavioural interventions with either individual or family-based interventions in this review.

**Topics**

We captured nutritional interventions involving cancer survivors, with or without their family members. We excluded physical activity interventions for cancer survivors and nutritional interventions for childhood cancer patients receiving active treatment as these have been targeted by alternate Cochrane reviews (Braam 2013a; Jones 2010).

**Settings**

We did not impose any restriction on the settings for the interventions; settings may have included community, home-based or hospital-based interventions.

**Delivery**

All methods of delivery of the intervention were eligible, including face-to-face, telephone and online interventions. There were no restrictions regarding the interventionist. That is, eligible interventions were those that were delivered by specialist and non-specialist medical and allied health professionals, as well as by other non-health professionals.

**Types of comparison**

We included studies which compared nutrition interventions to a non-intervention control group that received usual care or another intervention.

**Types of outcome measures**

**Primary outcomes**

A change in nutritional intake which was measured by one or more of the following.

1. Weighed food diaries.
2. Self-reported food diaries.
3. Single or multiple 24-hour recalls.
4. Food frequency questionnaires.

The nutrients may include but are not limited to:

1. energy;  
2. protein;  
3. fat;  
4. carbohydrate;  
5. calcium;  
6. iron;  
7. folate;  
8. vitamin(s);  
9. mineral(s).

**Secondary outcomes**

1. Metabolic risk factors, i.e. glucose and insulin metabolism.  
2. Cardiovascular risk factors, i.e. resting blood pressure, blood lipids, and cholesterol.  
3. Measures of weight and body fat distribution, i.e. body mass index (BMI), Dual-energy X-ray Absorptiometry (DEXA) and weight/height percentiles.  
4. Behavioural change, i.e. changes in nutritional intake.  
5. Changes in knowledge regarding disease risk and nutritional intake.  
6. Participant views of the intervention.  
7. Measures of health status and quality of life.  
8. Measures of harm associated with the process or outcomes of the intervention.  

**Search methods for identification of studies**

See: Cochrane Childhood Cancer methods used in reviews (Module CCG 2014).

**Electronic searches**

We searched the following electronic databases: the Cochrane Central Register of Controlled Trials (CENTRAL; 2013, Issue 3), MEDLINE/PubMed (from 1945 to 6 April 2013), and Embase/Ovid (from 1980 to 6 April 2013). The search strategies for the different electronic databases (using a combination of controlled vocabulary and text words) are shown in the appendices (Appendix 1; Appendix 2; Appendix 3).
We did run the search again in August 2015; we have not yet fully assessed these results, but we will fully incorporate them in the review at the next update.

Searching other resources
We located information about trials not registered in CENTRAL, MEDLINE/PubMED, Embase/OVID, either published or unpublished, by searching the reference lists of relevant articles and review articles. We handsearched the conference proceedings of the International Society for Paediatric Oncology (SIOP) (from 2008 to 2012) and the International Conference on Long-Term Complications of Treatment of Children and Adolescents for Cancer (2008 to 2012). We scanned the ISRCTN register (www.isrctn.com) and the register of the National Institute of Health (NIH) (clinicaltrials.gov) for ongoing trials in the first half of 2013. We did not impose language restrictions on the search.

Data collection and analysis

Selection of studies
Two review authors (JC, CW), worked independently, screening all the titles and abstracts resulting from the searches and excluded articles that were clearly irrelevant. The same review authors retrieved full-text copies of all relevant articles and using the defined eligibility criteria, determined their eligibility for inclusion. We resolved any disagreement between review authors on classification of an article between the review authors. Third party arbitration was not necessary. There was a need for clarification of detail of one trial. One of the review authors (JC) contacted the study authors from Rai 2008 to obtain clarification for a complete assessment of the trial’s relevance for the review. The reasons for exclusion of any study considered for review are summarised below (Characteristics of excluded studies).

Data extraction and management
Two review authors (JC and CW) independently extracted data, using a standardised form, from each article. For each trial, they extracted the following data.

1. Characteristics of the studies, including the study sponsors and the authors’ affiliations, study design, risk of bias items, duration of study, loss to follow-up, and compliance.

2. Characteristics of study population, including country where participants enrolled, inclusion and exclusion criteria, number randomised in each arm, information on the control group, demographic characteristics, type of cancer, age at diagnosis, cancer treatment, time since diagnosis, and time beyond active treatment.

3. Characteristics of the intervention, including type of nutritional intervention, details of the intervention, frequency, duration, intensity, number of sessions, intervention format (i.e. individual or group, professionally led or not, home- or facility-based), description of control intervention, adherence and contaminations, as well as cointerventions (i.e. physical activity, medication use).

4. Characteristics of the outcomes, as stated previously.
We entered and combined the trial data using Review Manager 5 (RevMan 2014). One review author entered the data into RevMan 5 (JC), and another review author worked independently to verify the data entry (CW). We resolved any disagreement between review authors on classification of an article between the review authors. Third party arbitration was not necessary.

Assessment of risk of bias in included studies
Two independent review authors (JC, CW) assessed the validity of each study using the risk of bias items, as described in the module of Cochrane Childhood Cancer (Module CCG 2014), which are based on the risk of bias domains from the Cochrane Handbook for Systematic Reviews of Interventions (Higgins 2011). We reported the following criteria for each trial: adequate sequence generation and allocation concealment (selection bias), masking or blinding of personnel, participants, and outcome assessors (performance or detection bias), incomplete data (attrition bias), and selective outcome reporting (reporting bias). We also assessed baseline imbalance (gender, ethnicity, diagnosis, age, and health behaviour or nutritional intake) and differential diagnostic activity as other potential sources of bias. We assessed these issues as ‘low risk of bias’, ‘high risk of bias’, or ‘unclear’. We resolved any disagreement between review authors. Third party arbitration was not necessary.

Measures of treatment effect
For continuous outcomes, we assessed the mean difference between groups. For dichotomous outcomes, we assessed relative risk.

Unit of analysis issues
We aimed to include cluster-randomised, cross-over, and repeated measures trials in this analysis, though none of the eligible studies used these methodologies.

Dealing with missing data
It was necessary to contact the authors of the Rai 2008 study to gather further detail on the nutrition intervention. We performed intention-to-treat analysis for all studies.
**Assessment of heterogeneity**

As we were unable to pool any of the data due to the different outcome measures and interventions between the trials, we were unable to assess heterogeneity using the I$^2$ analysis.

**Assessment of reporting biases**

We had planned to assess reporting bias by constructing funnel plots. However, as there were less than 10 studies included in this review, the power of the tests was too low to distinguish chance from real asymmetry (Higgins 2011), and so we did not carry this out.

**Data synthesis**

We entered the data of the included studies into Review Manager 5 (RevMan 2014). We performed data analysis according to the Cochrane Handbook for Systematic Reviews of Interventions (Higgins 2011). As we were unable to pool the data for a meta-analysis, we provided a narrative summary of the trial findings according to the review objectives. For data that was provided as medians and ranges, we converted the mean difference to mean and standard deviation based on the methodology of Hozo 2005.

**GRADE**

Two independent review authors used the GRADE system to rate the overall quality of the evidence for each of the following outcomes (Guyatt 2008a; Guyatt 2008b): calcium intake, bone mineral density, use of nutrition as health protective behaviour, junk food intake, milk consumption, and calcium supplementation. The GRADE approach defines the quality of a body of evidence as ‘high’, ‘moderate’, ‘low’ or ‘very low’. (Higgins 2011). Factors that may have resulted in a decrease in the quality of evidence included: 1) study limitations; 2) inconsistency; 3) indirectness; 4) imprecision; and 5) publication bias.

**Subgroup analysis and investigation of heterogeneity**

We had planned to perform subgroup analyses based on the following categories: 1) age at intervention (< 13 years; 13 to 18 years; > 18 years); 2) forms of intervention (face-to-face; phone etc); 3) duration of intervention; 4) childhood cancer type; and 5) type of treatment received. Due to insufficient trials and lack of data in the included studies, we were unable to conduct such analyses.

**Sensitivity analysis**

As pooling of the results was not possible, we were unable to use sensitivity analyses to explore the impact of the inclusion of studies with a high risk of bias and studies with an unclear risk of bias.

**RESULTS**

**Description of studies**

See: Characteristics of included studies; Characteristics of excluded studies; Characteristics of ongoing studies.

**Results of the search**

We identified a total of 3607 studies from running the search through three electronic databases: CENTRAL, MEDLINE/PubMED, and Embase/OVID in April 2013. We identified an additional study from searching the ongoing trial registries. We did not identify any studies upon screening reference lists of relevant articles and reviews. We did not identify any studies from the conference proceedings from the International Pediatric Oncology Society (SIOP) or the International Conference on Long-Term Complications of Treatment of Children and Adolescents for Cancer. Initial screening of the title and abstracts of each study allowed the exclusion of 3598 publications. We obtained ten full-text articles, of which three studies, described in 6 full-text articles, met the inclusion criteria. Three studies did not meet the inclusion criteria and we classified one of the studies as ongoing. In August 2015 we identified an additional study, which we assessed in full-text and classified as an ongoing study (see Figure 1).
Figure 1. Study flow diagram.

- 3607 records identified through database searching
- 1 additional record identified through other sources
- 1 additional record identified through top-up search (results not yet completely assessed)

3609 articles identified

3598 studies were excluded based on screening of title and/or abstract

11 full-text articles assessed for eligibility

3 studies excluded (see table of excluded studies for reasons)

2 ongoing trials

3 studies included, described in 6 full-text articles
Included studies

We included three studies in this review. All three studies were randomised controlled trials (RCTs). For further details on the studies see Characteristics of included studies.

Participants

A total of 616 participants from the three studies were included in the analysis. One of the studies included participants who had been treated for ALL (Rai 2008). Cox 2005 and Mays 2011 included participants with all forms of paediatric cancer.

The number of participants in each study varied. The smallest study included a total of 38 participants in the intervention and 37 in the control group (Mays 2011). It was unclear whether any participants were lost to follow-up. The Cox 2005 study included a total of 266 participants (131 in the intervention and 135 in the control group). Four and one participant(s), respectively were lost to follow-up. The largest study included a total of 275 participants (141 in the intervention and 134 in the control group) (Rai 2008). Ninety-four participants (45 in the intervention and 49 in the control group) did not complete the study.

The ages of the participants varied among the three studies. Two studies recruited adolescent childhood cancer survivors (ages 11 to 21 years (Mays 2011) and 12 to 18 years (Cox 2005)). The third study included childhood cancer survivors of all ages up to 18 years (Rai 2008). None of the included studies had participants older than 21 years at study entry. The participants were a mean of seven years (Rai 2008), and a mean of 15 years since their cancer treatment had been completed (Cox 2005). Information on time since diagnosis was not clear for the study of Mays 2011.

Intervention

The timing of the interventions after the childhood cancer therapy varied among the studies. Mays 2012 included participants within two years of diagnosis, Cox 2005 included patients who had completed treatment at least two years prior, and Rai 2008 included participants who had completed treatment at least five years prior. The intervention and timing also varied among the three studies included in this analysis. Two of the studies included interventions that consisted of an initial, single, face-to-face health education session focusing on health behaviour change (Cox 2005; Mays 2011). One of these studies focused on general health behaviours (Cox 2005), such as reducing junk food intake. The individual education session was provided by a clinician or nurse practitioner during a routine visit to the hospital. These participants were giving education reinforcement, via the telephone, at three and six months after the intervention. The other intervention focused on bone health, calcium, and dairy intake, and the final assessment was done one month after the intervention (Mays 2011). The education session was provided in a group setting by a registered Dietitian.

The final study had a 36-month follow-up, with the focus of the intervention being on bone health (Rai 2008). The intervention consisted of calcium and vitamin D supplementation. Nutrition education was provided at baseline and every six months for 24 months. At baseline and 12 months postbaseline, the education was given face-to-face by a registered dietitian. At six months and 18 months the nutrition education was in the form of mailed information. For further information on these studies, see Characteristics of included studies.

The study of Cox 2005 also included a co-intervention of changing the health behaviour practices of smoking cessation, sun protection, and exercise. This study did not have any contraindications. The studies of Mays 2011 and Rai 2008 did not include any co-interventions or contraindications.

Control

Of the three studies included in this review, the control groups of two of those studies received standard care (Cox 2005; Mays 2011). The standard care between these groups did vary. The standard care of the control group for the study of Cox 2005 included late-effects screening and education on their risk factors which was provided during routine clinic visits. The standard care of the control group for the Mays 2011 study was no education on nutrition related risk factors. The control group of the final study received an identical nutrition education component as the intervention group in combination with placebo tablets (Rai 2008).

Outcomes

The primary outcomes of the studies in this review, were dietary/nutrient intake. The secondary outcomes measured by the included studies were body composition (bone mineral density) and health behaviours. The control group measurements were assessed at the same time points as the intervention groups for all three of the studies. The time points for the outcome measures differed between the studies. The study of Mays 2011 measured their outcomes (milk consumption frequency, calcium supplementation, dietary calcium intake) at baseline and one-month postintervention. The study of Cox 2005 measured their outcomes (frequency of nutrition as a health protective behaviour, frequency of junk food consumption as a health risk behaviour) at baseline and 12 months postintervention. The final study of Rai 2008 measured their outcomes (bone mineral density) at baseline, 12 months, 24 months and 36 months postintervention.
The other secondary outcomes were not addressed in any of the three included studies. These secondary outcomes were: metabolic risk factors, cardiovascular risk factors, changes in knowledge, participant views of the intervention, health status and quality of life, measures of harm, or cost-effectiveness of the intervention. All three studies had different methodologies and different outcomes being measured, and for this reason we were unable to pool the data. For further information on these studies see Characteristics of included studies and Data and analyses.

**Excluded studies**

We analysed the full-text publications of three studies but subsequently excluded them. *Mays 2012* was a validation study and did not include an intervention. *Nathan 2009* was a review of the literature and the results of a smoking cessation intervention. The final study included participants on maintenance therapy who had not completed their cancer therapy (*Moyer-Mileur 2009*). For information on the excluded studies, see Characteristics of excluded studies.

**Ongoing studies**

We could not include two studies in this review as the data collection is ongoing (*NCT01473342; Stern 2015*). For further information about these studies, see Characteristics of ongoing studies.

**Risk of bias in included studies**

See the risk of bias section in Characteristics of included studies and Figure 2 and Figure 3 for detailed information on the risk of bias assessment.
Figure 2. Risk of bias summary: review authors' judgements about each risk of bias item for each included study.

<table>
<thead>
<tr>
<th></th>
<th>Cox 2005</th>
<th>Mays 2011</th>
<th>Rai 2008</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>+</td>
<td>?</td>
<td>+</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td></td>
<td>?</td>
<td>+</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
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<td>+</td>
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<tr>
<td>Selective reporting (reporting bias)</td>
<td></td>
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</tr>
<tr>
<td>Other bias</td>
<td>+</td>
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<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
</tbody>
</table>
Figure 3. Risk of bias graph: review authors’ judgements about each risk of bias item presented as percentages across all included studies.

<table>
<thead>
<tr>
<th>Random sequence generation (selection bias)</th>
<th>Allocation concealment (selection bias)</th>
<th>Incomplete outcome data (attrition bias)</th>
<th>Selective reporting (reporting bias)</th>
<th>Other bias</th>
</tr>
</thead>
<tbody>
<tr>
<td>![Bias Graph]</td>
<td>![Bias Graph]</td>
<td>![Bias Graph]</td>
<td>![Bias Graph]</td>
<td>![Bias Graph]</td>
</tr>
</tbody>
</table>

Allocation

Two of the studies described an adequate random sequence generation and we assessed them at low risk of bias (Cox 2005; Rai 2008). In the study by Rai 2008, randomisation was completed by the pharmacy after participants had been stratified into sex, race, age, and bone mineral density. The Cox 2005 study used a randomisation procedure that was stratified by gender and age. We assessed the final study as unclear in the use of random sequence generation (Mays 2011). Mays 2011 reported that the participants were randomised, but no further information was provided on the procedure. We assessed two of the studies as having an unclear allocation concealment as there was no mention of the procedures used in the study methodologies (Cox 2005; Mays 2011). The Cox 2005 study referred to the methodology used by another author, though the methods used were still not clear. The Rai 2008 study used a well described randomisation procedure and we assessed this as having a low risk of allocation concealment.

Blinding

Performance bias

Due to the nature of the interventions, blinding of personnel or participants was impossible with two of the three studies (Cox 2005; Mays 2011); we assessed both studies as having a high risk of performance bias. In the final study (Rai 2008), participants and personnel were blinded to the intervention, as participants were given a vitamin supplement or a placebo; we assessed this study as having a low risk of performance bias.

Detection Bias

Although personnel cannot be blinded when delivering nutrition interventions such as these, it is possible for detection bias to be minimised by blinding the outcome assessment. The Cox 2005 study did not provide any information regarding blinding of the outcome assessment and the outcome was subjective (a self-reported outcome) and therefore we assessed the blinding of outcome assessment as high risk. In the remaining two studies (Mays 2011; Rai 2008), we assessed detection bias as low risk because the assessors were blinded to the study groups.

Incomplete outcome data

Two of the three studies reported dropouts during the study (Cox 2005; Rai 2008). No further information was provided on how the missing data were handled and we assessed these studies as having an unclear risk of attrition bias. Although the third study had a short follow-up time of one month and was less likely to have dropouts, no information was provided on study attrition; we assessed this study as having an unclear risk (Mays 2011).
Selective reporting

We assessed Mays 2011 as having a low risk of reporting bias. This study reported data at baseline and follow-up on all outcomes cited in the protocol or methodology section. Cox 2005 presented the results of a secondary analysis, not mentioned in the original protocol (Hudson 2002) and Rai 2008 did not publish all outcomes that were reported on the clinical trials registry. We assessed these two studies to be at high risk of reporting bias.

Other potential sources of bias

We assessed all studies for baseline imbalances and differential diagnostic activity as other potential sources of bias. In regards to baseline imbalances, there was no significant difference between the baseline data between the intervention and the control group for all studies (Cox 2005; Mays 2011; Rai 2008). We assessed all three studies as being at low risk for baseline imbalances. We classified all three studies at low risk of bias for differential diagnostic activity because the studies performed the same assessments in the intervention and the control group at all time points (Cox 2005; Mays 2011; Rai 2008).

Effects of interventions

The three studies included in this review focused on different outcomes. We were unable to pool the data and the findings reported were from individual studies only.

Primary Outcome

Change in nutritional intake

Calcium intake was the only nutrient that was assessed across any of the studies (Mays 2011). Use of a single, group-based behaviour change intervention showed no statistically significant difference in the calcium intake (as measured by a 24-hour recall) between the intervention (n = 38) and control group (n = 37) at the one-month follow-up (mean difference (MD) 111.60, 95% confidence interval (CI) -258.97 to 482.17; P = 0.56, low quality evidence) (Mays 2011; Figure 4). We downgraded the quality of the evidence for study limitations and imprecision (Table 1). After regression analysis, adjusting for baseline calcium intake and changes in knowledge and self-efficacy, there was a significantly greater calcium intake for the intervention as compared with the control group at the one-month follow-up (beta coefficient 4.92, 95% CI 0.33 to 9.52; P = 0.04) Mays 2011.

Secondary Outcome

1. Metabolic risk factors

This outcome was not assessed in any of the included studies.

2. Cardiovascular risk factors

This outcome was not assessed in any of the included studies.

3. Measures of weight and body fat distribution

Body composition was used as an outcome measure in one study (Rai 2008). The data were provided as medians and ranges. We converted the data to mean and standard deviation based on the methodology of Hozo 2005. There was no statistically significant difference in bone mineral density (measured with a DEXA scan) at the 36-month follow-up (MD -0.05, 95% CI -0.26 to 0.16; P = 0.64, moderate quality evidence) between those who received the calcium and vitamin D supplementation in conjunction with nutrition education (n = 141) and those participants who received nutrition education alone (n = 134) (Rai 2008; Figure 5). We downgraded the quality of the evidence due to imprecision (Table 1). There was no statistically significant difference in bone mineral density at the one-month follow-up (MD -0.05, 95% CI -0.26 to 0.16; P = 0.64).
density between the intervention and the control group at the 12-month (median difference -0.17, \( P = 0.99 \)) and 24-month follow-up (median difference -0.04, \( P = 0.54 \)).

Figure 5. Forest plot of comparison: 2 Comparison of calcium and vitamin D supplementation and nutrition education with nutrition education alone, outcome: 2.1 Body composition (bone mineral density).

4. Behavioural Change

The behaviour change outcome was assessed in two studies. In the first study, health behaviour was measured using single questions on a four-point Likert scale (Cox 2005). The participants were asked how often they engaged in health practising behaviours and rated this from 1 = never to 4 = always. A single, face-to-face, multi-component health behaviour change intervention with two telephone follow-ups brought about no statistically significant difference in the use of nutrition as a health protective behaviour (\( n = 131 \)) compared with those who received standard care (\( n = 135 \)) (MD -0.05, 95% CI -0.24 to 0.14; \( P = 0.60 \), low quality evidence) (Cox 2005; Figure 6). We downgraded the quality of the evidence due to study limitations and imprecision (Table 1). The same intervention brought about a statistically significant reduction in self-reported junk food intake (measured on a four-point Likert scale: 1 = never to 4 = always) in the intervention (\( n = 131 \)) compared with the control group (\( n = 135 \)) (MD -0.17, 95% CI -0.33 to -0.01; \( P = 0.04 \), low quality evidence) (Figure 7). We downgraded the quality of the evidence due to study limitations and imprecision (Table 1).
A single, face-to-face, group-based health behaviour session focusing on bone health brought about a statistically significant increase in the intervention group’s self-reported milk consumption (measured in number of days) (MD 0.43, 95% CI 0.07 to 0.79; P = 0.02, low quality evidence) as compared with those who received standard care (Mays 2011; Figure 8). We downgraded the quality of the evidence due to study limitations and imprecision (Table 1).

The intervention was also effective in increasing the participants days on calcium supplementation (MD 11.42, 95% CI 7.11 to 15.73; P < 0.00001) (Figure 9). There was a statistically significant increase in calcium supplementation in the group that received the education sessions compared with those who received standard care (risk ratio (RR) 3.35, 95% CI 1.86 to 6.04; P < 0.0001, low quality evidence) (Figure 10). We downgraded the quality of the evidence due to study limitations and imprecision (Table 1). A total of 31 participants took some form of calcium supplementation after the intervention and nine participants took some form of calcium supplementation in the standard care group.

5. Changes in knowledge regarding disease risk and nutritional intake

This outcome was not assessed in any of the included studies.
6. Participant views of the intervention
This outcome was not assessed in any of the included studies.

7. Measures of health status and quality of life
This outcome was not assessed in any of the included studies.

8. Measures of harm associated with the process or outcomes of the intervention
This outcome was not assessed in any of the included studies.

9. Cost-effectiveness of the intervention
This outcome was not assessed in any of the included studies.

DISCUSSION

Summary of main results
Childhood cancer survivors are at higher risk of health conditions such as osteoporosis, metabolic syndrome, endocrine disorders, and cardiovascular disease than their peers (Nathan 2009). Targeted nutritional interventions may prevent (Steinberger 2012; Stolley 2010), or reduce the incidence of these chronic diseases (Nathan 2009; Oeffinger 2006). This systematic review included three trials that have studied the efficacy of a nutritional intervention, in a randomised manner, in childhood cancer survivors (Cox 2005; Mays 2011; Rai 2008). These studies utilised differing methodologies, and as a consequence, we were unable to pool results.

The interventions that appeared to bring about a significant positive change were those that focused on health behaviour change. A single, group-based health behaviour education session significantly increased self-reported milk intake (mean difference (MD) 0.43, 95% confidence interval (CI) 0.07 to 0.79; P = 0.02), use of calcium supplementation (risk ratio (RR) 3.35, 95% CI 1.86 to 6.04; P < 0.0001), and the number of days on calcium supplementation (MD 11.42, 95% CI 7.11 to 15.73; P < 0.0001) as compared with standard care (Mays 2011). The intervention did not improve calcium intake (MD 111.60, 95% CI -258.97 to 482.17; P = 0.56), though a regression analysis, adjusting for baseline calcium intake and changes in knowledge and self-efficacy, found a significantly greater calcium intake for the intervention as compared with the control group at the one month follow-up (beta coefficient 4.92, 95% CI 0.33 to 9.52; P = 0.04). This study had a short follow-up time of one month and the effect of the intervention long-term was not assessed.

A face-to-face, multi-component health behaviour session with two telephone follow-ups with education reinforcement, over a 12-month period, reduced self-reported junk food intake (MD -0.17, 95% CI -0.33 to -0.01; P = 0.04) but did not improve childhood cancer survivors’ use of nutrition as a health-protecting behaviour (MD -0.05, 95% CI -0.24 to 0.14; P = 0.60) as compared with standard care (Cox 2005).

The Rai 2008 study was the only study to assess the efficacy of nutritional supplementation on childhood cancer survivors’ body composition. This study was a randomised, double-blind randomised controlled trial (RCT) of calcium and vitamin D supplementation versus placebo. Both the intervention and control group received nutrition education by a registered dietitian. There was no statistically significant difference in bone mineral density as measured by dual-energy X-ray absorptiometry (DEXA) between the intervention and the control group at the 36-month follow-up (MD -0.05, 95% CI -0.26 to 0.16; P = 0.64). There was also no statistically significant difference in bone mineral density between the intervention and the control group at the 12-month (median difference -0.17, P = 0.99) and 24-month follow-up (median difference -0.04, P = 0.54).

Overall completeness and applicability of evidence
This review does not provide evidence that the nutritional interventions used in these studies improved dietary intake or body composition in childhood cancer survivors. The Mays 2011 study was the only included study that assessed the primary outcome of a change in nutritional intake. Mays 2011 found no statistically significant improvement in calcium intake with a single, group-based, education session. Although a regression analysis, adjusting for baseline calcium intake and changes in knowledge and self-efficacy, found a significantly greater calcium intake for the intervention as compared with the control group at the one month follow-up. The study had a short follow-up time of one month and long-term compliance with the nutritional changes were not assessed.

There was a modest, positive effect for health behaviour change interventions on improving self-reported health behaviours such as junk food consumption (Cox 2005), and milk intake (Mays 2011). As the results of the Cox 2005 study were based on a secondary analysis, these results do need to be interpreted with caution. Although no statistically significant differences were found for many of the outcomes, this could be the result of low power in the studies. It should be noted that no evidence of effect is not the same as evidence of no effect.

The following outcomes were not assessed in any of the included studies: metabolic and cardiovascular markers, changes in knowledge, participant views of the intervention, health status and quality of life, measures of harm, or the cost-effectiveness of the intervention.

The two studies that did show a positive change in health behaviours may not be applicable in all settings. The intervention required an initial face-to-face information session. This type of
intervention may not be possible for survivors of childhood cancer who come from geographically diverse regions who may not travel to the primary care centre for long-term follow-up. An efficacy of interventions utilising computer and other technologies may need to be assessed. The ongoing study that was not included in this review is assessing the use of Smartphone applications and other virtual technologies to provide nutritional education for survivors of childhood cancer (NCT01473342). This type of intervention may be more applicable across a variety of clinical settings. Many of this systematic review’s predetermined outcomes (e.g. metabolic risk factors, cardiovascular risk factors, changes in knowledge, and measures of harm) were not assessed in the included studies. Only one of the studies assessed the primary outcome of dietary intake. Although two of the interventions found a significant positive change in health behaviours, there is no evidence to suggest that this translates to the prevention of risk factors such as cardiovascular disease, metabolic syndrome, or obesity. Future interventions should consider assessing outcomes such as body composition and blood lipids in combination with dietary intake and changes in health behaviours. All three of the captured studies were from paediatric oncology units in the USA. The findings therefore may not be generalisable to childhood cancer survivors from other countries, especially low-income countries.

Quality of the evidence
By applying the GRADE criteria (Guyatt 2008a; Guyatt 2008b), the quality of findings varied between moderate (bone mineral density) and low (all other outcomes). We downgraded all outcomes one level for imprecision. Due to lack of blinding of participants, personnel, and outcome assessors, we further downgraded the quality of evidence for the outcomes ‘self-reported nutrition’ and ‘junk food’. Due to lack of details regarding the randomisation procedure and lack of blinding of participants and personnel, we also downgraded the outcomes ‘calcium intake’, ‘milk consumption’, and ‘calcium supplementation’ to low quality. The Cox 2005 study had a high risk of reporting bias (results were from a secondary analysis), performance bias (inadequate blinding of personnel) and detection bias (inadequate blinding of outcome assessors). The Cox 2005 study had unclear selection bias and attrition bias. Results from this study therefore need to be interpreted with caution. The Rai 2008 study was the only study that we assessed as having a low risk of performance bias, as both the participants and personnel were blinded; the other two studies were at high risk of performance bias. Although it is difficult to blind participants to the intervention due to the nature of many nutritional trials, two studies blinded the assessors (Mays 2011; Rai 2008). Adequate allocation concealment would be possible for all nutritional intervention trials, though the Rai 2008 study was the only study that we assessed at low risk of selection bias; Mays 2011 had an unclear risk. The Mays 2011 study had a low risk of reporting bias and the other studies were at high risk of reporting bias. We assessed all three studies as unclear in their attrition bias and at low risk of other bias (Cox 2005; Mays 2011; Rai 2008). The studies had minimal baseline imbalance and no differential diagnostic activity.

Potential biases in the review process
We developed the search strategies for the electronic databases (CENTRAL, MEDLINE/PubMED, Embase/OVID) in collaboration with Cochrane Childhood Cancer. We undertook additional searching of clinical trial databases, reference lists, and proceedings from conferences. Although it is always possible that we have not identified all studies, an earlier published review did not identify any different additional interventions prior to 2010 (Stolley 2010).

Agreements and disagreements with other studies or reviews

Only one other review paper was identified in the literature systematically reviewing diet (and exercise) in childhood cancer survivors (Stolley 2010). This review included studies that focused on diet in childhood cancer survivors, though the majority of these were observational studies and unable to be included in the current review. They identified one nutritional intervention in childhood cancer survivors which was also included in our review (Cox 2005). Stolley 2010 concluded that the literature on the dietary intake of childhood cancer survivors is methodologically weak. There were very limited intervention studies and use of control groups in the observational studies was rare. Stolley 2010, highlights the minimal use of validated methods of dietary assessment. Since the Stolley 2010 review was published, the three trials included in this review have been completed, though the use of validated dietary methods remains poor.

AUTHORS’ CONCLUSIONS
Implications for practice
Due to a paucity of research and the heterogeneity of the studies included in this review, the review authors are unable to draw conclusions regarding the effectiveness of nutritional interventions for childhood cancer survivors. Although there is weak evidence for the improvement in health behaviours using health behaviour change interventions, there remains no evidence as to whether this translates into an improvement in dietary intake.

It is important to note that no evidence of effect is not the same as evidence of no effect. Many outcomes were not assessed in the
Implications for research

This review highlights the need for further intervention trials to be implemented in this population. More robust research methodology is required to determine whether dietary changes can occur in survivors of childhood cancer and whether these can reduce their risk of long-term health issues. The use of a randomised design with blinding of personnel to the outcome measures is possible with this type of nutritional intervention and is recommended in future studies. It is also suggested that future studies utilise validated measures of dietary intake. Objective measures of body composition, cardiovascular, and metabolic risk should also be included as outcome measures in these studies.

Acknowledgements

The authors would like to acknowledge the editorial base of Cochrane Childhood Cancer for their advice and support. The authors would also like to thank Susan Kaste for providing additional data for the study of Rai 2008 and Jodie Bartle for input into the initial protocol. We also thank Dr A Spinola-Castro and an undisclosed person who kindly agreed to peer review our manuscript. The editorial base of Cochrane Childhood Cancer is funded by Stichting Kinderen Kankervrij (KiKa), the Netherlands.

References

References to studies included in this review

Cox 2005 [published data only]

Mays 2011 [published data only]

Rai 2008 [published data only]

References to ongoing studies

NCT01473342 [published data only]

Stern 2015 [published data only]

Nutritional interventions for survivors of childhood cancer (Review)

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Additional references

Arroyave 2008

Braam 2013a

Diller 2009

Hozo 2005

Hudson 2002

Hudson 2009

Guyatt 2008b

Higgins 2011

Kuhn 2004

Lakka 2007

Landier 2006

Lewis 2004

Module CCG 2014
Kremers LCM, Leclercq E, van Dalen EC. Childhood Cancer Group. About The Cochrane Collaboration

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Nutritional interventions for survivors of childhood cancer (Review)

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(Cochrane Review Groups (CRGs)) 2014, Issue 11. Art. No.: CHILDCA.

**Mulhern 1995**

**Mulrooney 2009**

**Ness 2007**

**Oeffinger 2006**

**Oeffinger 2008**

**Oeffinger 2009**

**Oude Luttikhuis 2009**

**Pereira 2009**

**RevMan 2014 [Computer program]**

**Rubien 2008**

**Siviero-Miachon 2008**

**Skinner 2006**

**Steinberger 2012**

**Stolley 2009**

**Stolley 2010**

**Tota-Maharaj 2010**

**Zelen 1974**

**References to other published versions of this review**

Cohen 2012a
## Characteristics of included studies [ordered by study ID]

### Cox 2005

| Methods | Design: Parallel RCT  
Setting: Single site paediatric oncology unit, USA |
|---------|--------------------------------------------------------------------|
| Participants | **Number**  
Intervention: n = 131 (4 lost to follow-up)  
Control: n = 135 (1 lost to follow-up)  
**Age at study entry**  
Group: 12-18 years  
Intervention (mean ± SD): 15.09 ± 1.90 years  
Control (mean ± SD): 14.96 ± 1.97 years  
**Sex**  
Intervention: 57 males: 74 females  
Control: 61 males: 74 females  
**Diagnosis**  
Leukaemia/lymphoma:  
Intervention: 73  
Control: 72  
Solid tumour:  
Intervention: 58  
Control: 63  
**Treatment**  
Information not available  
**Age at diagnosis**  
Information not available  
**Time since diagnosis**  
Intervention (mean ± SD): 15.09 (1.90) years  
Control (mean ± SD): 10.31 (2.94) years  
**Inclusion criteria**  
1. 12-18 years  
2. In remission 2+ years from completion of therapy  
3. Adequate cognitive functioning  
4. English as a primary language  
**Exclusion criteria**  
1. Not USA residents  
2. English not their primary language |
| Interventions | **Intervention**  
The intervention consisted of standard care plus a single multi-behavioural intervention provided by a clinical physician or nurse practitioner during a routine visit to the long-term follow-up clinic. The multi-behavioural intervention consisted of:  
1. discussion of after therapy clinical summary  
2. health behaviour training of health goal  
3. health goal commitment to practice  
TelephoneNumber reinforcement of the education was provided at 3 and 6 months after their }
initial clinic visit
Cointerventions:
Changing the health behaviour practices of smoking cessation, sun protection and exercise
Contraindications:
None

**Control Group**
Standard care consists of:
1. breast or testicular self-examination
2. targeted late effects screening
3. clinical assessment
4. late effects risk counselling

**Cointerventions**
Other health behaviour practices were targeted during the intervention. These included; smoking cessation, sun protection and exercise

**Outcomes**
The outcomes were measured at baseline and 12 months postintervention for both the intervention and control groups

**Outcome measure:** Behavioural change
1) Frequency of nutrition as a health protective behaviour
2) Frequency of junk food consumption as a health risk behaviour

**Notes**

**Study sponsors**
Oncology Nursing Society Foundation (2003-2005)
American Lebanese Syrian Associated Charities (ALSAC)

### Risk of bias

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<tr>
<th>Bias</th>
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<th>Support for judgement</th>
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<td>Low risk</td>
<td>Quote: &quot;The randomisation was stratified by gender and age because of the clinical impression that risk perception could carry by gender or age&quot;</td>
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<td>Unclear risk</td>
<td>Comment: The study reported that five participants (four in the intervention group and one in the control group) were lost to follow-up. There was no discussion on how this data were handled. We were unable to assess how this would influence the outcome or whether this would have a clinically relevant effect</td>
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### Cox 2005 (Continued)

<table>
<thead>
<tr>
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<th>Risk Level</th>
<th>Comment</th>
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<tbody>
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<td>High risk</td>
<td>Comment: This study presented a secondary analysis of data. This analysis was not in the original publication of the results</td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>Comment: Minimal baseline imbalance: At baseline, there was no significant difference between the intervention and the control group for demographic and other reported characteristics. No differential diagnostic activity: All assessments were performed at baseline and follow-up for both the intervention and the control group.</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>High risk</td>
<td>Comment: This study does not discuss whether participants or personnel were blinded. Due to the nature of the study and the form of the intervention, it would be impossible for the participants and personnel to be blinded.</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>High risk</td>
<td>Comment: The outcome is subjective (a self-reported outcome) and the participants are not blinded.</td>
</tr>
</tbody>
</table>

### Mays 2011

**Methods**  
Design: Parallel RCT  
Setting: Two sites, Paediatric oncology units, USA

**Participants**

<table>
<thead>
<tr>
<th>Number</th>
</tr>
</thead>
</table>
| Intervention: 38  
Control: 37  
No information on attrition was available |

**Age at study entry**

<table>
<thead>
<tr>
<th>Group: 11-21 years</th>
</tr>
</thead>
</table>
| Intervention (mean ± SD): 14.2 ± 2.0 years  
Control (mean ± SD): 14.2 ± 2.8 years |

**Sex**

| Intervention: 17 males: 21 females  
Control: 19 males: 18 females |

**Diagnosis**

| Intervention: 21 leukaemia: 17 others  
Control: 18 leukaemia: 19 others |

**Treatment**

| Information not available |

**Age at diagnosis**
Information not available

**Time since treatment completion**
Information not available

**Inclusion criteria**
1. Previously treated for any form of oncologic malignancy
2. One or more years off treatment
3. One or more years cancer-free
4. Able to comprehend and speak English

**Exclusion criteria**
1. Suffering from renal insufficiency or end stage renal disease
2. Currently taking a thiazide diuretic
3. Suffering from a pervasive developmental or other major psychiatric disorder precluding valid informed consent

### Interventions

**Intervention**
The intervention consisted of a single half-day, group workshop in addition to standard care. The workshop was given by a registered dietitian. The workshop included an interactive behavioural session and focused on risk reducing health promotion behaviours. The workshop had a focus on bone health

**Cointerventions:**
None

**Contraindications:**
None

**Control group**
The control group received standard care and were offered the intervention at the conclusion of the study

### Outcomes
The outcomes were measured at baseline and 1 month postintervention for both the intervention and control groups. These outcomes were:
A) change in nutritional intake:
1) dietary calcium intake measured with 24-hour recall
B) behaviour change:
1) milk consumption frequency
2) use of calcium supplementation

### Notes
**Study sponsors**
American Cancer Society
Lance Armstrong Foundation
National Cancer Institute (CA091831)

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Unclear risk</td>
<td>Comment: The paper states that the participants were randomly allocated but no further information on the methodology was provided</td>
</tr>
</tbody>
</table>
### Allocation concealment (selection bias)  
Unclear risk  
Comment: The paper states that the participants were randomly allocated but no further information on the methodology was provided.

### Incomplete outcome data (attrition bias)  
All outcomes  
Unclear risk  
Comment: There was no information provided on participant attrition.

### Selective reporting (reporting bias)  
Low risk  
Comment: This study reported data at baseline and follow-up on all outcomes cited in the protocol or methodology section.

### Other bias  
Low risk  
Comment: Minimal baseline imbalance: At baseline, there was no significant difference between the intervention and the control group for demographic and other reported characteristics.  
No differential diagnostic activity: All assessments were performed at baseline and follow-up for both the intervention and the control group.

### Blinding of participants and personnel (performance bias)  
All outcomes  
High risk  
Comments: This study does not discuss whether participants or personnel were blinded. Due to the nature of the study and the form of the intervention, it would be impossible for the participants and personnel to be blinded.

### Blinding of outcome assessment (detection bias)  
All outcomes  
Low risk  
Quote: “All telephone interviews were administered by a trained research assistant who was masked to the trial condition.”

### Rai 2008

#### Methods

| Design: Parallel RCT | Setting: Single site, paediatric oncology unit, USA |

#### Participants

<table>
<thead>
<tr>
<th>Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intervention: n = 141 (45 dropouts)</td>
</tr>
<tr>
<td>Control: n = 134 (49 dropouts)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Age at study entry</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intervention (mean; range): 16.6 (9.4 to 35.3) years</td>
</tr>
<tr>
<td>Control (mean; range): 17.2 (9.4 to 33.5) years</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sex</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intervention: 78 males: 63 females</td>
</tr>
<tr>
<td>Control: 78 males; 56 females</td>
</tr>
<tr>
<td>-----------------------------</td>
</tr>
</tbody>
</table>

**Diagnosis**

ALL

**Treatment**

Radiation:

Intervention: 53
Control: 34

Chemotherapy:

Intervention: 141
Control: 134

**Age at diagnosis**

Intervention (mean; range): 4.7 (0.7 to 17.4) years
Control (mean; range): 4.6 (1.0 to 16.39) years

**Time since treatment completion**

Intervention (mean; range): 7.1 (5.0 to 18.2) years
Control (mean; range): 7.2 (4.6 to 19.1) years

**Inclusion criteria**

1. Treated on St Judes Children's Research Hospitals total XI, XII or XIII treatment protocol
2. At least five years from completion of cancer therapy
3. In first remission

**Exclusion criteria**

1. Active disease
2. Pregnant or lactating females
3. Inability to chew or swallow pills
4. Currently consuming more than 800 mg of supplemental calcium or 800 IU of vitamin D
5. Anaemia

<table>
<thead>
<tr>
<th>Interventions</th>
</tr>
</thead>
</table>

**Intervention**

This study was a 24-month nutrition and supplementation intervention. The intervention group received nutrition education sessions every 6 months. At baseline and 12 months these were given face-to-face by a registered dietitian, and at 6 months and 18 months these were given in the form of mailed information. The education included information such as:

1. number of serves of dairy products
2. serve sizes of dairy foods
3. healthy diet

The intervention group was also given 24 months of calcium and vitamin D supplementation which were taken daily

**Cointerventions:**

None

**Contraindications:**

None

**Control group**

The control group received education sessions identical to the intervention group. They also received placebo tablets instead of calcium and vitamin D supplements.
Outcomes | The outcomes were measured at baseline, 12 months, 24 months and 36 months post-intervention for both the intervention and control groups  
Outcome measure: body composition  
1) Bone mineral density

Notes | Study sponsors  
National Institutes of Health; Grant number: P30 CA-21765  
Center of Excellence grant from the State of Tennessee  
Le Bonheur Foundation (Memphis TN)  
American Lebanese Syrian Associated Charities (ALSAC)  
NIH; Grant numbers: R21 HD059292; GM 92666  
Gabrielle’s Angel Foundation

Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>Comment: The participants were stratified when randomised into sex, race, age and BMD Z-score</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Quote: “Only the St Jude pharmacy had access to the randomisation system, which is maintained by the Department of Biostatistics at St Jude”</td>
</tr>
</tbody>
</table>
| Incomplete outcome data (attrition bias)  
All outcomes | Unclear risk | Comment: There were a large number of dropouts in both the intervention (45) and control groups (49). It is unclear how this data were treated |
| Selective reporting (reporting bias) | High risk | Comment: This study did not publish all outcomes that were reported on the clinical trials registry |
| Other bias | Low risk | Minimal baseline imbalance: At baseline, there was no significant difference between the intervention and the control group for demographic and other reported characteristics  
No differential diagnostic activity: All assessments were performed at baseline and follow-up for both the intervention and the control group |
| Blinding of participants and personnel (performance bias)  
All outcomes | Low risk | Comment: Both the participants and the research personnel were blinded |
Characteristics of excluded studies [ordered by study ID]

<table>
<thead>
<tr>
<th>Study</th>
<th>Reason for exclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mays 2012</td>
<td>This was a validation study and did not include an intervention</td>
</tr>
<tr>
<td>Moyer-Mileur 2009</td>
<td>The study included participants on maintenance therapy</td>
</tr>
<tr>
<td>Nathan 2009</td>
<td>This study contains a review of the literature and only reported on a smoking cessation intervention in childhood cancer survivors</td>
</tr>
</tbody>
</table>

Characteristics of ongoing studies [ordered by study ID]

NCT01473342

<table>
<thead>
<tr>
<th>Trial name or title</th>
<th>Mila Blooms intervention study: an App for promoting physical activity and health diet among adolescent survivors of childhood cancer</th>
</tr>
</thead>
</table>
| Methods             | Design: RCT (own controls)  
Setting: Single site paediatric oncology unit, USA                                                                             |

| Participants | Number  
Estimated enrolment: 30  
Age  
12-19 years  
Sex  
Both genders  
Diagnosis  
ALL  
Inclusion criteria  |
|--------------|-----------------------------------------------------------------------------------------------------------------------------------|
|              | 1. Off therapy for 2+ years  
2. Karnofsky score > 80  
3. Speak and read/write fluent English  
4. Working phone number  
5. Participants must live with parents |
**Exclusion criteria**

1. Physician reported deficits in neurocognitive functioning

### Interventions

**Control phase:**
All participants will take part in the 8-week control phase where they will complete initial data collection

**Intervention:**
The intervention will commence after the 8-week control phase has finished and will run for 9 weeks. The participants will receive a smart phone which contains the Mila Blooms gaming app. The app consists of:
1. tools for tracking health behaviours
2. avatar system to provide virtual rewards
3. social network of users for support

Participants will also receive weekly coaching telephone calls to help with goal-setting and compliance

### Outcomes

Healthy diet
Physical activity
Prevent weight gain

### Starting date

April 2011

### Contact information

Sharnail Bazemore: sharnail.bazemore@duke.edu

### Notes

Stern 2015

### Stern 2015

**Trial name or title**
NOURISH-T

**Methods**
Design: Feasability study RCT
Setting: Two paediatric oncology units, USA

**Participants**

**Number**
Estimated enrolment: 110

**Age**
Caregivers of 5-12 year olds

**Sex**
Both genders

**Diagnosis**
All childhood cancer survivors between 6 months to 4 years off treatment

**Interventions**

**Intervention**
6-week, 10 session study
6 sessions delivered face-to-face
4 sessions delivered via the telephone
Focus on behavioural change, healthy eating, and physical activity

**Control**
Enhanced usual care
1 x wellness session + written education materials on healthy eating
### Stern 2015  *(Continued)*

| Outcomes          | Feasibility  
|-------------------|--------------|
|                   | Dietary intake  
|                   | QoL  
|                   | BMI  
|                   | Physical activity  

| Starting date | Unknown  

| Contact information | Marilyn Stern: mstern1@usf.edu  

| Notes |  

BMI: body mass index  
QoL: quality of life  
RCT: randomised controlled trial
**DATA AND ANALYSES**

Comparison 1. Comparison of a single, group behaviour intervention, with standard care

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Change in nutritional intake (calcium)</td>
<td>1</td>
<td>75</td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>111.60 [-258.97, 482.17]</td>
</tr>
<tr>
<td>2 Behavioural change (milk consumption)</td>
<td>1</td>
<td>75</td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>0.43 [0.07, 0.79]</td>
</tr>
<tr>
<td>3 Behavioural change (days of calcium supplementation)</td>
<td>1</td>
<td>75</td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>11.42 [7.11, 15.73]</td>
</tr>
<tr>
<td>4 Behavioural change (any calcium supplementation)</td>
<td>1</td>
<td>75</td>
<td>Risk Ratio (M-H, Fixed, 95% CI)</td>
<td>3.35 [1.86, 6.04]</td>
</tr>
</tbody>
</table>

Comparison 2. Comparison of calcium and vitamin D supplementation combined with nutrition education with nutrition education only

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Body composition (bone mineral density)</td>
<td>1</td>
<td>275</td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>-0.05 [-0.26, 0.16]</td>
</tr>
</tbody>
</table>

Comparison 3. Comparison of a 12-month, face-to-face and telephone health behaviour intervention with standard care

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Behavioural change (nutrition)</td>
<td>1</td>
<td>266</td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>-0.05 [-0.24, 0.14]</td>
</tr>
<tr>
<td>2 Behavioural change (junk food)</td>
<td>1</td>
<td>266</td>
<td>Mean Difference (IV, Fixed, 95% CI)</td>
<td>-0.17 [-0.33, -0.01]</td>
</tr>
</tbody>
</table>
### Analysis 1.1. Comparison 1 Comparison of a single, group behaviour intervention, with standard care, Outcome 1 Change in nutritional intake (calcium).

Review: Nutritional interventions for survivors of childhood cancer

Comparison: 1 Comparison of a single, group behaviour intervention, with standard care

Outcome: 1 Change in nutritional intake (calcium)

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Single, education session</th>
<th>Standard care</th>
<th>Mean Difference</th>
<th>Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean(SD)</td>
<td>N</td>
<td>Mean(SD)</td>
<td>IV,Fixed, 95% CI</td>
</tr>
<tr>
<td>Mays 2011</td>
<td></td>
<td>38</td>
<td>1263.7 (736.2)</td>
<td>37</td>
</tr>
<tr>
<td><strong>Total (95% CI)</strong></td>
<td>38</td>
<td>37</td>
<td>0.43 [ 0.07, 0.79 ]</td>
<td></td>
</tr>
</tbody>
</table>

Heterogeneity: not applicable

Test for overall effect: Z = 0.59 (P = 0.56)

Test for subgroup differences: Not applicable

### Analysis 1.2. Comparison 1 Comparison of a single, group behaviour intervention, with standard care, Outcome 2 Behavioural change (milk consumption).

Review: Nutritional interventions for survivors of childhood cancer

Comparison: 1 Comparison of a single, group behaviour intervention, with standard care

Outcome: 2 Behavioural change (milk consumption)

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Single, education session</th>
<th>Standard care</th>
<th>Mean Difference</th>
<th>Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean(SD)</td>
<td>N</td>
<td>Mean(SD)</td>
<td>IV,Fixed, 95% CI</td>
</tr>
<tr>
<td>Mays 2011</td>
<td>3.36 (0.72)</td>
<td>38</td>
<td>2.93 (0.88)</td>
<td>37</td>
</tr>
<tr>
<td><strong>Total (95% CI)</strong></td>
<td>38</td>
<td>37</td>
<td>100.0 %</td>
<td>0.43 [ 0.07, 0.79 ]</td>
</tr>
</tbody>
</table>

Heterogeneity: not applicable

Test for overall effect: Z = 2.31 (P = 0.021)

Test for subgroup differences: Not applicable
Analysis 1.3. Comparison 1 Comparison of a single, group behaviour intervention, with standard care, Outcome 3 Behavioural change (days of calcium supplementation).

Review: Nutritional interventions for survivors of childhood cancer

Comparison: 1 Comparison of a single, group behaviour intervention, with standard care

Outcome: 3 Behavioural change (days of calcium supplementation)

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Single, education session</th>
<th>Standard care</th>
<th>Mean Difference</th>
<th>Weight</th>
<th>Mean Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N  Mean(SD)</td>
<td>N  Mean(SD)</td>
<td>IV,Fixed,95% CI</td>
<td></td>
<td>IV,Fixed,95% CI</td>
</tr>
<tr>
<td>Mays 2011</td>
<td>38 14.45 (10.97)</td>
<td>37 3.03 (7.86)</td>
<td>-11.42 [ 7.11, 15.73 ]</td>
<td>100.0 %</td>
<td></td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td>38</td>
<td>37</td>
<td>100.0 %</td>
<td>11.42 [ 7.11, 15.73 ]</td>
<td></td>
</tr>
</tbody>
</table>

Heterogeneity: not applicable
Test for overall effect: Z = 5.19 (P < 0.00001)
Test for subgroup differences: Not applicable

Analysis 1.4. Comparison 1 Comparison of a single, group behaviour intervention, with standard care, Outcome 4 Behavioural change (any calcium supplementation).

Review: Nutritional interventions for survivors of childhood cancer

Comparison: 1 Comparison of a single, group behaviour intervention, with standard care

Outcome: 4 Behavioural change (any calcium supplementation)

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Single, education session</th>
<th>Standard care</th>
<th>Risk Ratio</th>
<th>Weight</th>
<th>Risk Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n/N</td>
<td>n/N</td>
<td>M-H,Fixed,95% CI</td>
<td></td>
<td>M-H,Fixed,95% CI</td>
</tr>
<tr>
<td>Mays 2011</td>
<td>31/38</td>
<td>9/37</td>
<td>3.35 [ 1.86, 6.04 ]</td>
<td>100.0 %</td>
<td></td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td>38</td>
<td>37</td>
<td>100.0 %</td>
<td>3.35 [ 1.86, 6.04 ]</td>
<td></td>
</tr>
</tbody>
</table>

Total events: 31 (Single, education session), 9 (Standard care)
Heterogeneity: not applicable
Test for overall effect: Z = 4.03 (P = 0.000055)
Test for subgroup differences: Not applicable
### Analysis 2.1. Comparison 2
Comparison of calcium and vitamin D supplementation combined with nutrition education with nutrition education only, Outcome 1 Body composition (bone mineral density).

**Review:** Nutritional interventions for survivors of childhood cancer

**Comparison:** 2 Comparison of calcium and vitamin D supplementation combined with nutrition education with nutrition education only

**Outcome:** 1 Body composition (bone mineral density)

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Supplements + education</th>
<th>Education only</th>
<th>Mean Difference</th>
<th>Weight</th>
<th>Mean Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N Mean(SD)</td>
<td>N Mean(SD)</td>
<td>IV,Fixed,95% CI</td>
<td></td>
<td>IV,Fixed,95% CI</td>
</tr>
<tr>
<td>Rai 2008</td>
<td>141 -0.61 (1.042)</td>
<td>134 -0.56 (0.72)</td>
<td>100.0 %</td>
<td>-0.05</td>
<td>[ -0.26, 0.16 ]</td>
</tr>
<tr>
<td><strong>Total (95% CI)</strong></td>
<td>141</td>
<td>134</td>
<td>100.0 %</td>
<td>-0.05</td>
<td>[ -0.26, 0.16 ]</td>
</tr>
</tbody>
</table>

Heterogeneity: not applicable

Test for overall effect: $Z = 0.46$ ($P = 0.64$)

Test for subgroup differences: Not applicable

### Analysis 3.1. Comparison 3
Comparison of a 12-month, face-to-face and telephone health behaviour intervention with standard care, Outcome 1 Behavioural change (nutrition).

**Review:** Nutritional interventions for survivors of childhood cancer

**Comparison:** 3 Comparison of a 12-month, face-to-face and telephone health behaviour intervention with standard care

**Outcome:** 1 Behavioural change (nutrition)

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Education</th>
<th>Standard care</th>
<th>Mean Difference</th>
<th>Weight</th>
<th>Mean Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N Mean(SD)</td>
<td>N Mean(SD)</td>
<td>IV,Fixed,95% CI</td>
<td></td>
<td>IV,Fixed,95% CI</td>
</tr>
<tr>
<td>Cox 2005</td>
<td>131 2.34 (0.75)</td>
<td>135 2.39 (0.794)</td>
<td>100.0 %</td>
<td>-0.05</td>
<td>[ -0.24, 0.14 ]</td>
</tr>
<tr>
<td><strong>Total (95% CI)</strong></td>
<td>131</td>
<td>135</td>
<td>100.0 %</td>
<td>-0.05</td>
<td>[ -0.24, 0.14 ]</td>
</tr>
</tbody>
</table>

Heterogeneity: not applicable

Test for overall effect: $Z = 0.53$ ($P = 0.60$)

Test for subgroup differences: Not applicable
Analysis 3.2. Comparison of a 12-month, face-to-face and telephone health behaviour intervention with standard care, Outcome 2 Behavioural change (junk food).

Review: Nutritional interventions for survivors of childhood cancer

Comparison: 3 Comparison of a 12-month, face-to-face and telephone health behaviour intervention with standard care

Outcome: 2 Behavioural change (junk food)

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Education</th>
<th>Standard care</th>
<th>Mean Difference</th>
<th>Weight</th>
<th>Mean Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N Mean(SD)</td>
<td>N Mean(SD)</td>
<td>IV,Fixed,95% CI</td>
<td></td>
<td>IV,Fixed,95% CI</td>
</tr>
<tr>
<td>Cox 2005</td>
<td>131 2.32 (0.643)</td>
<td>135 2.49 (0.71)</td>
<td>-0.17 [-0.33, -0.01]</td>
<td>100.0 %</td>
<td>-0.17 [-0.33, -0.01]</td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td>131 135</td>
<td></td>
<td>100.0 %</td>
<td>-0.17 [-0.33, -0.01]</td>
<td></td>
</tr>
</tbody>
</table>

Heterogeneity: not applicable

Test for overall effect: Z = 2.05 (P = 0.041)

Test for subgroup differences: Not applicable

ADDITIONAL TABLES

Table 1. GRADE Assessment

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Study limitations</th>
<th>Inconsistency</th>
<th>Indirectness</th>
<th>Imprecision</th>
<th>Publication bias</th>
<th>GRADE assessment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcium intake</td>
<td>Serious limitations (-1): lack of details on randomisation procedure, and lack of blinding of participants and personnel</td>
<td>n.a.</td>
<td>n.a.</td>
<td>Imprecision (-1): only one small study</td>
<td>n.a.</td>
<td>++, Low quality</td>
</tr>
<tr>
<td>Bone mineral density</td>
<td>No serious limitations</td>
<td>n.a.</td>
<td>n.a.</td>
<td>Imprecision (-1): only one study</td>
<td>n.a.</td>
<td>++++, Moderate quality</td>
</tr>
<tr>
<td>Use of nutrition as health protective behaviour</td>
<td>Serious limitations (-1): lack of blinding of participants, personnel,</td>
<td>n.a.</td>
<td>n.a.</td>
<td>Imprecision (-1): only one study</td>
<td>n.a.</td>
<td>++, Low quality</td>
</tr>
</tbody>
</table>
Table 1. GRADE Assessment  (Continued)

<table>
<thead>
<tr>
<th></th>
<th>and outcome assessors</th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Junk food intake</td>
<td>n.a.</td>
<td>n.a.</td>
<td>Imprecision (-1): only one study</td>
<td>n.a.</td>
<td>++, Low quality</td>
</tr>
<tr>
<td>1 (266)</td>
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<td>Calcium supplementation</td>
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n.a. = not applicable

APPENDICES

Appendix 1. Search strategy for Cochrane Central Register of Controlled Trials (CENTRAL)

1. For Population the following text words were used:
   (infant OR infant* OR newborn OR newborn* OR new-born* OR baby OR baby* OR babies OR neonat* OR child OR child* OR schoolchild* OR schoolchild OR school child OR school child* OR kid OR kids OR toddler* OR adolescent OR adole* OR teen* OR boy* OR girl* OR minors OR minors* OR underag* OR under ag* OR juvenil* OR youth* OR kindergar* OR puberty OR puber* OR pubescen* OR prepubescent* OR prepuberty* OR pediatrics OR pediatric* OR paediatric* OR pediatric* OR schools OR nursery school* OR preschool* OR pre school* OR primary school* OR secondary school* OR elementary school* OR high school* OR highschool* OR school age OR schoolage OR school age* OR schoolage* OR infancy OR young adult OR young adults OR young adult*)

   AND (post treatment OR off treatment OR treatment complet* OR treatment termin* OR follow up OR follow-up OR followup OR survivor OR survivors OR Long-Term Survivors OR Long Term Survivors OR Long-Term survivor OR survive* OR surviving)
2. For Nutrition the following text words were used:
patient education OR practice guideline OR practice guidelines OR dietary guideline OR dietary guidelines OR practice guideline* OR dietary guideline* OR diet OR diets OR diet* OR diets* OR dietetic OR dietetics OR diet therapy OR health diet OR healthy food OR health promoting behaviour OR health promoting behaviour OR (diet* AND intervent*) OR (diet* AND advic*) OR diet* AND counsel* OR (diet* AND therap*) OR (diet* AND treatment*) OR (diet* AND educat*) OR (nutriti* AND intervention*) OR (nutriti* AND advice*) OR (nutriti* AND counsel*) OR (nutriti* AND therap*) OR (nutriti* AND treatment*) OR (nutriti* AND educat*) OR (nutriti* AND support) OR supportive therapy

3. For Outcome the following text words were used:
food OR foods OR food* OR food intake OR eating OR ingestion OR nutrition OR nutrition* OR (health* AND diet*) OR (health* AND food*) OR energy intake OR caloric intake OR kilojoule OR kilojoules OR calorie OR calori* OR caloric restriction OR vitamin OR vitamins OR vitamin* OR minerals OR minerals* OR mineral OR mineral* OR micro-nutrient OR micro-nutrients OR macro-nutrient OR macro-nutrients OR nutrient OR nutrients OR calcium OR folate OR folic acid OR iron OR ferric OR ferrous OR protein OR proteins OR fat intake OR fat reduced OR dietary fat restriction OR low fat OR low calorie OR low energy OR reduced energy OR caloric controlled OR fatty foods OR high fat OR fruit OR fruits OR vegetable OR vegetables OR dietary composition OR carbohydrate intake OR obesity OR obese OR adiposity OR body weight OR overweight OR body mass index OR BMI OR body mass OR body fat distribution OR body composition OR "bioelectrical impedance analysis" OR health behavior OR health behaviors OR health behaviour OR health behaviours OR health behaviour* OR health behaviour* OR health promotion OR behaviour change OR health change OR behaviour change* OR behavior change* OR health behaviour change OR health behaviour change OR health behaviour change OR health behaviour change* OR health behaviour change* OR lifestyle OR life style* OR weight gain OR weight gains OR weight gain* OR body weight OR weight loss OR weight change OR weight changes OR weight change* OR overnutrition OR overeating OR hyperphagia OR Metabolic syndrome OR Waist hip ratio OR Waist height ratio OR Skinfold thickness OR Skinfold thicknesses OR Skinfold thickness* OR DEXA OR Diabetes OR type 2 diabetes OR glucose metabolism OR insulin metabolism OR insulin resistance OR hyperinsulinemia OR hyperinsulinaemia OR cardiomyopathy OR myocardial Infarction OR fat metabolism OR cardiovascular risk factor OR cardiovascular risk factors OR cardiovascular risk factor* OR cardiovascular disease OR cardiovascular diseases OR blood pressure OR hypertension OR blood lipid OR blood lipids OR blood lipid* OR hyperlipidemia OR hyperlipidaemia OR dyslipidemia OR dyslipidaemia OR cholesterol metabolism OR hypercholesterolemia OR osteoporosis OR bone mineral density OR dual energy x-ray absorptiometry OR malnutrition OR undernutrition OR Nutritional Deficiency OR Nutritional Deficiencies OR ideal body weight OR body image OR eating disorder OR eating disorders OR eating disorder* OR disordered eating OR fussy eating OR food refusal OR quality of life OR QoL

4. For Cancer the following text words were used:
cancer OR oncology OR oncolog* OR neoplasms OR neoplas* OR carcinoma OR carcinom* OR tumor OR tumour OR tumor* OR tumour* OR cancer* OR malignan* OR hematopoietical OR hematopoietical OR hematological neoplasms OR hematolo* OR bone marrow transplantation OR bone marrow transplant* OR leukemia OR leukaemia OR lymphoma

The search was performed in title, abstract or keywords
Final search 1 and 2 and 3 and 4
[* = zero to many characters]

Appendix 2. Search strategy for MEDLINE (PubMed)

1. For Population the following MeSH headings and text words were used:
(infant OR infant* OR newborn OR new born* OR new-born* OR baby OR baby* OR babies OR neonat* OR perinat* OR postnat* OR child OR child* OR schoolchild OR schoolchild OR school child OR school child* OR kid OR kids OR toddler* OR adolescent OR adole* OR teen* OR boy* OR girl* OR minors OR minors* OR underag* OR under ag* OR juvenil* OR youth* OR kindergar* OR puberty OR puber* OR pubescen* OR pubescen* OR prepuber* OR prepuberty* OR pediatrics OR pediatric* OR paediatric* OR paediatric* OR schools OR nursery school* OR preschool* OR pre school* OR primary school* OR secondary school* OR elementary school* OR elementary school OR high school* OR highschool* OR school age OR schoolage OR school age* OR schoolage* OR infancy OR schools, nursery OR infant, newborn OR young adult[mh] OR adult[mh] OR young adult)
AND (post treatment OR off treatment OR treatment complex* OR treatment termin* OR follow up OR follow-up OR followup OR survivor OR survivors OR Long-Term Survivors OR Long Term Survivors OR Long-Term survivor OR Survivor, Long-Term OR Survivors, Long-Term OR survivo* OR surviving)

2. For Nutrition the following MeSH headings and text words were used:
patient education OR practice guideline OR practice guidelines OR dietary guideline OR dietary guidelines OR practice guideline* OR dietary guideline* OR diet OR diets OR diet* OR diets* OR dietetic OR dietetics OR diet therapy OR health diet OR healthy food OR health promoting behaviour OR health promoting behaviour OR (diet* AND intervent*) OR (diet* AND advic*) OR diet* AND counsel* OR (diet* AND therap*) OR (diet* AND treatment*) OR (diet* AND educat*) OR (nutriti* AND intervent*) OR (nutriti* AND advice*) OR (nutriti* AND counsel*) OR (nutriti* AND therap*) OR (nutriti* AND treatment*) OR (nutriti* AND educat*) OR (nutriti* AND support) OR supportive therapy

3. For Outcome the following MeSH headings and text words were used:

food OR foods OR food* OR food intake OR eating OR ingestion OR nutrition OR nutrition* OR (health* AND diet*) OR (health* AND food*) OR energy intake OR caloric intake OR kilojoule OR kilojoules OR calorie OR calori* OR caloric restriction OR vitamin OR vitamins OR vitamin* OR minerals OR minerals* OR mineral OR mineral* OR micro-nutrient OR micro-nutrients OR macro-nutrient OR macro-nutrients OR nutrient OR nutrients OR calcium OR calcium OR folic acid OR iron OR ferric OR ferrous OR protein OR proteins OR fat intake OR fat reduced OR dietary fat restriction OR low fat OR low calorie OR low energy OR reduced energy OR caloric controlled OR fatty foods OR high fat OR fruit OR fruits OR vegetable OR vegetables OR dietary composition OR carbohydrate intake OR obesity OR obese OR adiposity OR body weight OR overweight OR body mass index OR BMI OR body mass OR body fat distribution OR body composition OR “bioelectrical impedance analysis” OR health behavior OR health behaviors OR health behaviour OR health behaviours OR health behaviour* OR health behaviour* OR health promotion OR behaviour change OR behaviour change OR behaviour change* OR health behaviour change OR health behaviour change OR health behaviour change* OR health behaviour change* OR life style OR life style* OR weight gain OR weight gains OR weight gain* OR body weight OR weight loss OR weight change OR weight change* OR overnutrition OR overeating OR hyperphagia OR Metabolic syndrome OR Waist hip ratio OR Waist height ratio OR Skinfold thickness OR Skinfold thicknesses OR Skinfold thickness* OR DXA OR Diabetes Type OR diabetes OR glucose metabolism OR insulin metabolism OR insulin resistance OR hyperinsulinemia OR hyperinsulinaemia OR cardiomyopathy OR myocardial Infarction OR fat metabolism OR cardiovascular risk factor OR cardiovascular risk factors OR cardiovascular risk factor* OR cardiovascular disease OR cardiovascular diseases OR blood pressure OR hypertension OR blood lipid OR blood lipids OR blood lipid* OR hyperlipidemia OR hyperlipidaemia OR dyslipidaemia OR dyslipidaemia OR cholesterol metabolism OR hypercholesterolemia OR osteoporosis OR bone mineral density OR dual energy x-ray absorptiometry OR malnutrition OR undernutrition OR Nutritional Deficiency OR Nutritional Deficiencies OR ideal body weight OR body image OR eating disorder OR eating disorders OR eating disorder* OR disordered eating OR fussy eating OR food refusal OR quality of life OR QoL.

4. For Cancer the following MeSH headings and text words were used:
cancer OR oncology OR oncolog* OR neoplasms OR neoplas* OR carcinoma OR carcinom* OR tumor OR tumour OR tumor* OR tumour* OR cancer* OR malignan* OR hematooncological OR hematologic neoplasms OR hematolo* OR bone marrow transplantation OR bone marrow transplant* OR leukemia OR leukaemia OR lymphoma

5. For RCTs and CCTs the following MeSH headings and text words were used:

(randomized controlled trial[pt]) OR (controlled clinical trial[pt]) OR (randomized[tiab]) OR (placebo[tiab]) OR (drug therapy[sh]) OR (randomly[tiab]) OR (trial[tiab]) OR (groups[tiab]) AND (humans[mh])

Appendix 3. Search strategy for EMBASE (OVID)

1. For Population the following Emtree terms and text words were used:

infant$ OR infancy/ OR newborn$ OR baby/ OR child/ OR preschool child/ OR school child/

2. adolescent/ OR juvenile/ OR boy/ OR girl/ OR puberty/ OR prepuberty/ OR pediatrics/

3. primary school/ OR high school/ OR kindergarten/ OR nursery school/ OR school/

4. ot/1-3

5. (infant$ OR newborn$ OR (new adj born$) OR baby OR baby$ OR babies OR neonate$ OR perinat$ OR postnat$).mp.

6. (child$ OR (school adj child$) OR schoolchild$ OR (school adj age$) OR schoolage$ OR (pre adj school$) OR preschool$).mp.

7. (kid OR kids OR toddler$ OR adolescent$ OR teen$ OR boy$ OR girl$).mp.

8. (minor$ OR (under adj age$) OR underage$ OR juvenile$ OR youth$ OR young adult OR young adults OR young adult$).mp.

9. (puber$ OR pubescen$ OR prepubescen$ OR prepubert$).mp.
(pediatric$ or paediatric$ or pediatric$).mp.
(school or schools or (high adj school$) or highschool$ or (primary adj school$) or (nursery adj school$) or (elementary adj school) or (secondary adj school$) or kindergar$.mp.
or/5-11
4 or 12
AND
(survivor or survivors or (long adj term survivor) or (long adj term survivors) or survivo$).mp.
survivor/ or cancer survivor/
survivi$.mp.
(post treatment or off treatment).mp.
(treatment complet* or treatment termin*).mp.
(follow up or followup or follow-up).mp. or exp follow up/
or/1-6
For Nutrition the following Emtree terms and text words were used:
patient education.mp. or exp patient education/
(practice guideline or practice guidelines or practice guideline$).mp.
exp practice guideline/
(dietary guideline or dietary guidelines or dietary guideline$).mp.
exp DIET/ or diet.mp.
(diets or diet$ or diets$ or dietetic or dietetics).mp.
diet therapy.mp. or exp diet therapy/
(health diet or healthy food).mp. or exp health food/
exp health behavior/
(health promoting behaviour or health promoting behavior).mp.
diet$ and intervent$.mp.
diet$ and advic$.mp.
diet$ and counsel$.mp.
diet$ and therap$.mp.
diet$ and treatment$.mp.
diet$ and educat$.mp.
(nutriti$ and intervent$.mp.
nutriti$ and advice$.mp.
nutriti$ and counsel$.mp.
nutriti$ and therap$.mp.
nutriti$ and treatment$.mp.
nutriti$ and educat$.mp.
nutriti$ and support).mp.
supportive therapy.mp.
or/1-24
For Outcome the following Emtree terms and text words were used:
(food or foods or food* or foods* or food intake).mp.
exp FOOD INTAKE/ or exp FOOD/
eting.mp. or exp EATING/
ingestion.mp. or exp INGESTION/
exp NUTRITION/
(nutrition or nutrition$).mp.
(health$ and diet$).mp.
(health$ and food$).mp.
(energy intake or carbohydrate intake or caloric intake).mp. or exp caloric intake/
(kilojoule or kilojoules or calorie or calori$ or caloric restriction).mp.
(vitamin/
(vitamin or vitamins or vitamin$).mp.
exp MINERAL/
14. (minerals or minerals$ or mineral or mineral$).mp.
15. exp trace element/
16. (micro-nutrient or micro-nutrients).mp.
17. exp MACRONUTRIENT/
18. (macro-nutrient or macro-nutrients or nutrient or nutrients).mp.
19. (calcium or 7440-70-2).mp.
20. (folic acid or 59-30-3).mp.
21. (iron or 7439-89-6 or ferric or ferrous).mp.
22. protein/
23. (protein or proteins).mp.
24. exp low fat diet/
25. (fat reduced or dietary fat restriction or low fat or fat intake).mp.
26. (low calorie or low energy or reduced energy or calorie controlled).mp.
27. (fatty foods or high fat).mp.
28. (fruit or fruits or vegetable or vegetables).mp.
29. exp dietary intake/ or dietary composition.mp.
30. exp OBESITY/
31. (obesity or obese).mp.
32. adiposity.mp.
33. body weight.mp. or exp body weight/
34. overweight.mp.
35. exp body mass/
36. (body mass index or BMI or body mass).mp.
37. body fat distribution.mp. or exp body fat distribution/
38. bioelectrical impedance analysis.mp.
39. body composition.mp. or exp body composition/
40. exp health behavior/
41. (health behavior or health behaviors or health behaviour or health behaviours or health behaviour$. or health behaviour$).mp.
42. health/
43. (health knowledge or health attitude$).mp.
44. health promotion.mp. or exp health promotion/
45. exp behavior change/
46. (behaviour change or behavior change or behaviour change$ or behavior change$ or health behaviour change or health behavior change or health behaviour change$ or health behavior change$).mp.
47. exp lifestyle/
48. (life style or life style$ or lifestyle or lifestyle$).mp.
49. (weight gain or weight gains or weight gain$).mp.
50. exp weight gain/
51. exp weight reduction/
52. (weight loss or weight change or weight changes or weight change$).mp.
53. exp OVERNUTRITION/
54. exp HYPERPHAGIA/
55. (overnutrition or overeating or hyperphagia).mp.
56. Metabolic syndrome.mp. or metabolic syndrome X/
57. Waist hip ratio.mp. or exp waist hip ratio/
58. Waist height ratio.mp.
59. exp skinfold thickness/
60. (Skinfold thickness or Skinfold thicknesses or Skinfold thickness$).mp.
61. DEXA.mp. or exp dual energy X ray absorptiometry/
62. (Diabetes or type 2 diabetes).mp. or exp diabetes mellitus/
63. glucose metabolism.mp. or exp glucose metabolism/
64. insulin metabolism.mp. or exp insulin metabolism/
65. exp hyperinsulinemia/ or (hyperinsulinemia or hyperinsulinaemia).mp.
66. exp CARDIOMYOPATHY/ or cardiomyopathy.mp.
67. myocardial Infarction.mp. or exp heart infarction/
68. fat metabolism.mp. or exp lipid metabolism/
69. exp cardiovascular risk/
70. (cardiovascular risk factor or cardiovascular risk factors or cardiovascular risk factor$).mp.
71. exp cardiovascular disease/ or (cardiovascular disease or cardiovascular diseases).mp.
72. blood pressure.mp. or exp blood pressure/
73. exp hypertension/ or hypertension.mp.
74. exp lipid blood level/
75. (blood lipid or blood lipids or blood lipid$).mp.
76. cholesterol metabolism.mp. or exp cholesterol metabolism/
77. exp hypercholesterolemia/ or hypercholesterolemia.mp.
78. exp hyperlipidemia/ or (hyperlipidemia or hyperlipidaemia).mp.
79. exp dyslipidemia/ or (dyslipidemia or dyslipidaemia).mp.
80. osteoporosis/co, dt, rt, si, th [Complication, Drug Therapy, Radiotherapy, Side Effect, Therapy]
81. Osteoporosis.mp.
82. bone mineral density.mp. or exp bone density/
83. malnutrition.mp. or exp MALNUTRITION/
84. undernutrition.mp.
85. exp nutritional deficiency/
86. (Nutritional Deficiency or Nutritional Deficiencies).mp.
87. ideal body weight.mp. or exp body weight/
88. body image.mp. or exp body image/
89. exp eating disorder/
90. (eating disorder or eating disorders or eating disorder$ or disordered eating or fussy eating).mp.
91. exp food refusal/ or food refusal.mp.
92. exp “quality of life”/ or (quality of life or QoL).mp.
93. or/1-92

4. For **Cancer** the following Emtree terms and text words were used:
1. (cancer or cancers or cancer$).mp.
2. (oncology or oncolog$).mp. or exp oncology/
3. (neoplasm or neoplasms or neoplasm$).mp. or exp neoplasm/
4. (carcinoma or carcinom$.mp. or exp carcinoma/
5. (tumor or tumour or tumor$ or tumour$ or tumors or tumours).mp. or exp tumor/
6. (malignan$ or malignant).mp.
7. (hematooncological or hemato oncological or hemato-oncological or hematologic neoplasms or hematolo$).mp. or exp hematologic
malignancy/
8. (leukemia or leukaemia).mp. or exp LEUKEMIA/
9. lymphoma.mp. or exp LYMPHOMA/
10. or/1-9

5. For **RCTs and CCTs** the following Emtree terms and text words were used:
1. Randomized Controlled Trial/
2. Controlled Clinical Trial/
3. randomized.ti,ab.
4. placebo.ti,ab.
5. randomly.ti,ab.
6. trial.ti,ab.
7. groups.ti,ab.
8. drug therapy.sh.
9. or/1-8
10. Human/
11. 9 and 10

Final search 1 AND 2 AND 3 AND 4 AND 5
CONTRIBUTIONS OF AUTHORS

JC and CW were the main contributors to the data extraction, management, and assessment of bias. RC reviewed the manuscript.

DECLARATIONS OF INTEREST

Jennifer Cohen: None known.

Claire Wakefield: None Known.

Richard Cohn: None Known.

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Internal sources

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External sources

- There are no external sources of support, Other.

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There has been a change to the author team.