Carbohydrate quality and human health

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Carbohydrate quality and human health: a series of systematic reviews and meta analyses

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Summary

Background Previous systematic reviews and meta analyses explaining the relationship between carbohydrate quality and health have usually examined a single marker and a limited number of clinical outcomes. We have considered the impact of carbohydrate quality as measured by intakes of dietary fibre, whole grains or pulses, and dietary glycaemic index or glycaemic load on non-communicable disease (NCD) incidence, mortality, and risk factors to more precisely quantify the predictive potential of the markers, to determine which are most useful, and to establish an evidence base for quantitative recommendations for intakes of dietary fibre.

Methods Prospective studies published prior to April 2017 and randomised controlled trials published prior to February 2018, which reported on indicators of carbohydrate quality and NCD incidence, mortality and risk factors were systematically reviewed and meta analysed. Studies were identified by searches in PubMed, Ovid Medline, Embase, and the Cochrane Central Register of Controlled Trials and by hand searching of previous publications. Searches, data extraction, and bias assessment were duplicated independently. Robustness of pooled estimates from random effects models was considered with sensitivity analyses, meta regression, dose response testing, and subgroup analyses. The GRADE approach was used to assess quality of evidence.

Findings 135 million person years of data from prospective studies and 58 clinical trials with a total of 4,635 adult participants were included in the analyses. Observational data suggest a 15-30% decrease in all-cause and cardiovascular related mortality, and incidence of coronary heart disease, stroke, type 2 diabetes, and colorectal cancer when comparing the highest dietary fibre consumers with the lowest. Clinical trials show significantly lower body weight, systolic blood pressure and total cholesterol when comparing higher with lower intakes. Risk reduction associated with a range of critical outcomes was greatest when daily intake of dietary fibre was between 25-29 grams. Dose response curves suggested that higher intakes may confer even greater benefit with regard to protection against cardiovascular diseases, type 2 diabetes and colorectal and breast cancer. Comparable findings for wholegrain intake were observed. Smaller or no risk reductions were found with the observational data when comparing the effects of diets characterised by low rather than higher glycaemic index or load. Overall the certainty of evidence regarding the relationships between carbohydrate quality and critical outcomes was graded as moderate for dietary fibre, low to moderate for whole grains, and low to very-low for glycaemic index. Data relating to other dietary exposures was limited.

Interpretation The complementary findings from prospective studies and clinical trials relating to the reduction in mortality, NCD incidence, and their risk factors associated with relatively high intakes of dietary fibre and whole grains as well as striking dose response evidence indicate that the relationships may be causal. Benefit to individuals and populations may be expected from implementation of dietary recommendations to increase dietary fibre intake and to replace refined grains with whole grains.

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**Research in Context**

**Evidence before this study**
Carbohydrate-containing foods consisting principally of sugars, starches and dietary fibre (non starch polysaccharide) provide the major source of energy worldwide. The role of free sugars as a determinant of adverse health outcomes has been clarified and clear guidelines relating to their restriction issued. Dietary fibre and some starches are associated with health benefits and dietary guidelines typically encourage regular consumption of vegetables, cereals, pulses and whole fruit which are rich sources of these and other health promoting nutrients. However, previous systematic reviews and meta analyses examining the relationship between starches and dietary fibre, and health outcomes have usually examined a single indicator of carbohydrate quality and a limited number of disease outcomes. Thus it has not been possible to determine the extent to which the predictive potential of these indicators applies across the spectrum of non-communicable disease (NCDs) nor which are most useful in nutrition guidelines or when recommending food choices. Quantitative recommendations relating to dietary fibre have not had a strong evidence base.

**Added value of this study**
We have undertaken systematic reviews and meta analyses of prospective studies and clinical trials that have reported on the relationship between the most widely studied indicators of carbohydrate quality (dietary fibre, whole grains or pulses, dietary glycaemic index or glycaemic load) and mortality and incidence of a wide range of NCDs and their risk factors. Parallel consideration of prospective studies and clinical trials has enabled an exploration of the extent to which changes in cardiometabolic risk factors associated with altering intake of dietary carbohydrate align with the effect of carbohydrate quality on disease risk observed in the prospective studies. Dose response curves were generated and the benefits from different amounts of total dietary fibre were calculated. The approach recommended by the GRADE Working Group has been used to assess the quality of evidence and the magnitude and importance of the observed associations which influence the confidence in nutrition recommendations.

**Implications of all the available evidence**
The complementary findings from prospective studies and clinical trials, which show that higher intakes of dietary fibre or whole grains are related to a reduction in the risk of a wide range of NCDs and their risk factors, provide convincing evidence for nutrition recommendations to replace refined grains with whole grains and increase dietary fibre to at least 25-29g per day, with additional benefits likely to accrue with greater intakes. In the light of current evidence, dietary glycaemic index or glycaemic load may be less useful as overall measures of carbohydrate quality than dietary fibre and wholegrain content.
Introduction

Prior to the mid twentieth century carbohydrates were principally regarded as an energy source and nutrition recommendations suggested that carbohydrates should contribute the energy deficit remaining after intakes of fat and protein had been specified. From the mid 1950s there was increasing awareness of the potential of “sugar” (principally sucrose) to increase the risk of dental caries and in the 1960s Yudkin\(^1\) popularised the view that sugar was a major contributing cause of obesity, type 2 diabetes and cardiovascular disease, an opinion shared by Cleave and Campbell who described these and other chronic conditions as saccharine diseases.\(^2\) A substantial body of experimental, epidemiological and clinical trial data has accumulated since these early observations and based on extensive systematic reviews and meta analyses, the World Health Organization (WHO) has recently issued a strong recommendation, based on the association between free sugars and dental caries and obesity, for individuals to reduce intake to less than 10% total energy and a conditional recommendation suggesting that even greater benefit may accrue if intakes are below 5%.\(^3\) Comparable recommendations have been made by national governments and professional organisations worldwide.

It is more than half a century since Burkitt, Trowell and Painter, based largely on epidemiological observations in Africa, suggested that processing of cereal based foods (grains) with removal of what came to be called dietary fibre, rather than excessive intakes of sugar, were key determinants of both cardiometabolic and large bowel diseases.\(^4,5\) Nevertheless, until relatively recently rather less attention has been given to starches and dietary fibre, the other major components of dietary carbohydrate. While nutrition guidelines issued by many governments and professional organisations encourage increased consumption of vegetables, fruit and whole grains, there are fewer quantitative guidelines for sources and intakes of dietary fibre and starch. We report here on a series of systematic reviews and meta analyses on indicators of carbohydrate quality and non-communicable disease (NCD) incidence, mortality, and risk factors. The research was commissioned by WHO to inform the development of updated recommendations regarding carbohydrate intake.

Methods

We followed reporting standards for systematic reviews and meta analyses.\(^6\) Literature searches, identification of eligible studies, data extraction and bias assessment were undertaken independently by at least two researchers, with discrepancies resolved with an additional reviewer.

PICO tables and eligibility criteria

PICO tables (Appendix A) were agreed by the WHO Nutrition Guidance Expert Advisory Group (NUGAG). We report here on markers of carbohydrate quality that have been measured in an appreciable number of studies and trials (dietary fibre, dietary glycaemic index or glycaemic load, and wholegrain intake) and outcomes specified in the PICO tables. For prospective studies critical outcomes included all-cause, coronary heart disease (CHD) and stroke mortality and incidence of CHD, stroke, type 2 diabetes, and colorectal cancer. Important outcomes included cardiovascular disease (CVD) incidence and mortality and incidence of adiposity-related cancers (breast, endometrial, oesophageal, and prostate). Prospective studies which included only cohorts with specified pre-existing conditions were excluded.
For clinical trials we have reported on adiposity, fasting glucose, fasting insulin, insulin sensitivity, HbA1c, triglycerides, cholesterol, and blood pressure. We included parallel and crossover randomised clinical trials of at least four weeks duration that reported on higher compared with lower intakes of the dietary markers. Eligible trials could include diets with test foods provided, dietary advice, ad libitum diets or controlled feeding trials on free living individuals. Weight loss trials and trials involving provision of dietary fibre supplements in the forms of powders were excluded. Comparison diets were required to be matched for macronutrient composition and lifestyle modifications such as exercise.

Participants of eligible trials were adults and children free from acute or chronic disease but could include those with prediabetes, mild-moderate hypercholesterolemia, mild-moderate hypertension or metabolic syndrome. Trials including people on medications known to effect outcomes of interest or who were pregnant or in situations where regular eating habits were likely to change e.g. those suffering from eating disorders or who were breast feeding were excluded.

**Literature search**

Prospective observational studies were initially identified from systematic reviews and meta analyses that reported associations between carbohydrate intake or one of the specified measures of carbohydrate quality, and one or more of the key outcome measures. These systematic reviews were found through online searches using Ovid Medline, Embase, PubMed, Web of Science, and Scopus. This strategy was augmented by searches with low-risk-of-bias search terms for individual prospective studies and run up to the end of April 2017 to ensure identification of relevant published studies. No date or language restrictions were applied. A validation of the search procedure is provided in Appendix A.

For clinical trials, highly sensitive Cochrane search strategies were used to identify trials examining the effects of carbohydrate intakes on obesity, blood pressure, and cardiometabolic risk factors. OVID Medline, Embase, Cochrane Central Register of Controlled Trials (CENTRAL), Cochrane Database of Systematic Reviews (CDSR), and Food Science and Technology Abstracts (FSTA) databases were searched for trials published up to February 2018. Hand searching of references of systematic reviews, prospective studies, and clinical trials were completed to identify any studies that may have been missed. Search strategies are shown in Appendix A.

**Study selection**

Reviewers identified eligible studies by screening titles, abstracts and where appropriate full texts of articles. Where there were multiple publications from the same cohort, we used data for the longest follow-up period.

**Data extraction**

Data were extracted using pre-tested forms. For prospective studies the most adjusted values for effect size were extracted, where that value did not also specifically include adjustment for other carbohydrates. For clinical trials involving multiple interventions we extracted data from all relevant interventions. For crossover trials involving multiple interventions we extracted data only from the most relevant intervention and either the control group or the most relevant comparator intervention.
Risk of bias assessment

We used the ROBIS assessment tool\(^8\) to assess systematic reviews and meta analyses for quality and risk of bias, and the Newcastle Ottawa Scale (NOS)\(^9\) to assess risk of bias of each prospective study. For clinical trials we used Cochrane criteria.\(^{10}\)

Data analysis

For prospective studies we pooled the reported odds ratios or risk ratios with the DerSimonian and Laird random effects model\(^{11}\) in a high quantile versus low quantile analysis. When individual studies reported results separately by sex, we first combined these effect size estimates with a fixed effects model before including them within the pooled estimate. When eligible studies were based on and reported combined results from multiple cohort studies we extracted results for each cohort to include in the meta-analysis. Prospective studies reporting incidence or mortality were analysed separately. Where data were reported in a suitable format, we considered dose response relationships with the Greenland and Longnecker method\(^{12}\) assuming linearity with a two-stage dose response random-effects analysis. The average or mid-point of each defined quantile was used for the dose amount. Where the quantile dose range was open-ended, half the range of the adjacent quantile was used to determine the average intake. We used 30g to represent one serve of whole grains when a value for weight was not stated.\(^{13}\) Non-linear dose-response was assessed using restricted cubic splines with three knots at 10%, 50%, and 90% of distribution combined with multivariate meta analyses.\(^{14}\) We imputed the number of cases per quantile from the RR value when necessary. Linear and spline (with 95%CI) models are shown with each data point overlaid as circles. Circle size indicates the weighting of each data point with bigger circles indicating greater influence. Absolute risk values were calculated with GRADE Pro software.\(^{15}\)

To help establish optimal intakes of dietary fibre we considered the dose-response curves for total dietary fibre intake and critical health outcomes. We also compared the lowest consumers of dietary fibre with those consuming between 15-19, 20-24, 25-29, 30-34, and 35-39 grams of fibre per day with a random effects model. When studies reported more than one quantile of data within the pre-specified intake ranges, we first combined these quantiles with a fixed effects model before including them within the pooled estimate. We did this to measure the number of critical outcomes where an improvement in relative risk was observed in the higher intake categories.

For clinical trials high-versus-low analyses were undertaken with generic inverse models and random-effects. For outcomes that could be measured by different units, reported effects were presented as standardised mean differences. For studies reporting multiple follow-ups over time, the most recent, appropriately reported published data were used in the meta analyses. When crossover (paired data) studies did not report the mean difference between treatments and its standard error or other relevant statistics, end of treatment values were analysed as independent samples. Subgroup analyses by fibre amount or principal starch source were conducted when there were enough studies for subgroupings including more than one trial. For example high fibre interventions (0-25, 25-30, 30-35, >35 g/day) were considered to determine whether there were threshold effects or a possible dose response.
For all analyses heterogeneity was assessed with the I² statistic, and the Cochrane Q test. Sensitivity analyses were conducted when a I² statistic was found to be more than 50% or a p for heterogeneity of <0·10. Publication bias was assessed with Egger’s and Begg’s tests, and the trim and fill method. The effect of each individual study’s findings was considered with an influence analysis. For prospective studies analyses excluding those that scored less than six out of a possible nine with the NOS were conducted. If there was still unexplained heterogeneity we considered the impact of small studies reporting less than 200 cases or less than 2000 participants. For clinical trials, analyses excluding trials with a high risk of bias for at least one criterion were conducted to examine the influence of potential bias on outcomes. Meta regression analyses further examined effects of potential explanatory factors including trial design (crossover or parallel), study or trial duration, global region, differences in fibre intake achieved, source of fibre or starch and nutrition status of participants. Analyses were performed using the Cochrane Collaboration software and Stata statistical software. We used GRADE protocols, to judge the quality of the body of evidence as either high, moderate, low, or very-low. More detail on this approach is provided in Appendix A. Quality of the evidence was assessed by the research team and revised if required after discussion with the NUGAG Subgroup on Diet and Health.

Role of the funding source

With the exception of WHO, the funders of the study had no role in study design, data collection, data analysis, data interpretation, or writing the report. The corresponding author had full access to all the data in the study and had final responsibility for the decision to submit for publication.

Results

Data from 185 publications of prospective studies involving just under 135 million person years and 58 clinical trials with a total of 4,635 adult participants were included in the meta analyses. A flow chart of identified studies is shown in Figure 1, with details of these studies in Supplement 10. Critical outcome data for total fibre, wholegrain intake, and dietary glycaemic index are summarised in Tables 1-3 and shown in full in Appendices B-D for observational studies and Supplements 1 and 2 for trials. Dose response data are shown in Figures 2-4 and the supplementary material. Summary forest plots from clinical trial data are shown in Figure 5. Data and GRADE tables relating to all other indicators and outcomes are in Supplement 1-9.

Dietary fibre

The observational data in Table 1 show that higher intakes of total dietary fibre are associated with a 15-31% reduction in the risk of specified critical outcomes. For all-cause mortality and coronary heart disease incidence this translates into 13 fewer deaths (95%CI 8 to 18) and 6 fewer cases of CHD (95%CI 4 to 7) per 1000 participants over the duration of the studies. Sensitivity analyses of the tested associations did not change the direction or significance of any observed result. The quality of evidence contributing to the meta analyses of the cohort studies was, with the exception of the data relating to stroke, considered to be moderate.
Figure 2 shows dose response relationships for total fibre intake and total mortality, incidence of coronary heart disease, type 2 diabetes and colorectal cancer, many of which are linear with no sign of a plateau within the available data. When comparing the lowest fibre intakes with pre-specified ranges the greatest benefits were observed for those consuming 25-29g per day (improvement in 6 of the 7 critical outcomes), more so than those consuming between 15-19g per day (improvement in 3 of the 7 critical outcomes), or 20-24g per day (improvement in 4 of the 7 critical outcomes). These analyses are shown in full in Appendix B.

Mean differences between higher versus lower fibre intakes for a range of cardiometabolic risk factors are shown in Table 1 and the summary forest plots in Figure 5a. Dose response or threshold effects could not be determined from the clinical trial data. The quality of evidence contributing to the meta analyses of the trial data relating to body weight is high and total cholesterol and systolic blood pressure moderate because of unexplained heterogeneity between the trials.

Broadly similar effects were apparent in both the prospective studies and clinical trials, when examining fibre from different food groups or fibre described as soluble or insoluble, though limited data were available, other than for cereal fibre, the largest contributor to total dietary fibre (Supplements 1-9).

**Whole grains**

Cohort data showing the relationship between whole grains and the effect of increasing wholegrain intake on critical outcomes are shown in Table 2. Higher intakes of whole grains were associated with a 13-33% reduction in the risk. For all-cause mortality and coronary heart disease incidence this translates into 26 fewer deaths (95%CI 14 to 39) and 7 fewer cases (95%CI 3 to 10) per 1000 participants over the duration of the studies. Sensitivity analyses did not typically change the direction or significance of any pooled effect. The quality of evidence relating to colorectal cancer incidence is moderate, whilst for other critical outcomes it is low due to high heterogeneity not fully explained by sensitivity analysis. Dose response curves showing clear associations with increasing wholegrain intake and all-cause mortality or risk of coronary heart disease, type 2 diabetes, and colorectal cancer incidence are shown in Figure 3. Mean differences in cardiometabolic risk factors between higher and lower wholegrain consumption are shown in Table 2 and summary forest plots in Figure 5b. Evidence relating to body weight, cholesterol, and blood pressure is graded as moderate, downgraded due to unexplained heterogeneity.

**Glycaemic index**

Cohort data showing the relationship between dietary glycaemic index and the effect of decreasing the dietary glycaemic index on critical outcomes as demonstrated in the trials are shown in Table 3, dose responses are shown in Figure 4, and summary forest plot in Figure 5c. Data relating to the cohort studies which examined the effects of glycaemic load are presented in Supplement 3.

An 11% (95%CI 3% to 18%) relative risk reduction of type 2 diabetes was observed for those consuming low glycaemic index diets. However sensitivity analysis due to high heterogeneity attenuated the relative risk reduction to 5% (95%CI 13% less to 4% more). Stroke mortality was lower amongst those consuming lower
glycaemic index diets. The prospective studies generated evidence that is graded as low or very-low quality as a
result of high risk of bias, imprecision, and inconsistencies. Key outcome markers from the clinical trials on
decreasing the glycaemic index of a diet are shown in forest plots in Figure 5c. Trial data were usually of
moderate quality.

Discussion

Higher intakes of total dietary fibre or whole grains result in reduced incidence and mortality from several
NCDs. Less useful markers of carbohydrate quality are glycaemic index, glycaemic load, and sources of dietary
fibre where inconsistent findings or insufficient data provide low or very low quality evidence. In randomised
trials higher intakes of dietary fibre reduce body weight, lower blood cholesterol and systolic blood pressure.
These findings are supported by cohort studies that report reduced risk of coronary heart disease incidence and
mortality and diabetes incidence. The consistency between the trial and prospective study results together with
the dose response relationships are evidence that the effect on cardiometabolic diseases are likely to be causal
and not a consequence of confounding. In addition, prospective studies show striking reductions in and dose
response relationships with all-cause mortality, total cancer deaths, total cardiovascular disease, stroke
incidence, and colorectal, breast, and oesophageal cancer. For several of these outcomes the dose response is
linear. These findings together with the comparisons of clinical outcomes amongst those with different intakes
of dietary fibre suggest that individual adult intakes of total dietary fibre should be no less than 25 to 29 grams
per day with additional benefits likely to accrue with higher intakes. Population intakes in this range are
reported in some countries, but the majority consume less than 20 grams a day. Broadly similar trends were
apparent in the prospective studies that examined cereal fibre, typically the largest contributor to total dietary
fibre. Limited data were available regarding specific sources (legume, fruit, vegetable) or subcategories
(‘soluble’, ‘insoluble’, or extracted) of dietary fibre.

The results for wholegrain foods reflect those for dietary fibre. Prospective studies showed a reduction in all-
cause mortality, coronary heart disease, cancer deaths, and incidence of type 2 diabetes. As with dietary fibre
the observed reductions in risk are considerable, typically around 20% with significant dose response
relationships. The randomised controlled trials involving an increase in the amount of whole grains showed
improvements in body weight and lipids. The similar protective effects of higher intakes of wholegrain foods
and of dietary fibre suggest that the beneficial effects of whole grains may be due to their typically high dietary
fibre content. The GRADE criteria categorise the evidence linking most clinical outcomes with dietary fibre as
moderate, and with whole grains as low quality.

Dietary starch can be divided into several categories although rarely are measurements made of these
individual components in either prospective studies or randomised controlled trials. However the glycaemic
index of starch-containing foods or the overall glycaemic load of meals or diets including starchy foods provide
measures of starch quality and are widely reported. We have found that diets with a lower overall glycaemic
index appear to be associated with a reduced risk of stroke and type 2 diabetes. However the risk estimates,
other than for stroke mortality, are modest when compared with those for dietary fibre and following sensitivity
analyses were reduced and associated with confidence intervals which included 1. The findings from
prospective studies of glycaemic load are inconsistent. The results from trials show no consistent benefits on the clinical outcomes when changing the glycaemic index of a diet.

A major strength of the present study is that it has related key markers of carbohydrate quality to total mortality and mortality and incidence of the major nutrition-related non-communicable diseases and that prospective studies have been considered alongside randomised controlled trials. Other reviews and meta analyses have reported on a single indicator of carbohydrate quality and one or more outcomes. Our approach has enabled us to use these indicators of carbohydrate quality to provide a stronger justification than had previously been available for a quantitative recommendation relating to dietary fibre intake. That the evidence for the associations between the quality markers and outcomes was most frequently rated as ‘moderate’ or ‘low’ rather than ‘high’ may be regarded as a limitation. However this is an inevitable consequence of the use of GRADE criteria for assessment which typically require evidence from randomised controlled trials with disease endpoints in order to be rated as being of ‘high’. Furthermore when using the GRADE approach downgrading frequently occurs as a consequence of unexplained heterogeneity amongst the results of the different studies, even when all trend in a similar direction. This may be a consequence of studies being carried out in diverse populations or as a result of different methods of measuring dietary intake. With regard to the associations we have reported here between dietary fibre and whole grains and a wide range of clinical outcomes, the consistency of the findings, the striking dose response relationships and the substantial body of mechanistic evidence all contribute to the totality of evidence and increases our confidence in the findings.

Our findings are broadly comparable with other reviews and meta analyses that have reported on the association between dietary fibre and whole grains and one or more disease outcomes. However there is less consistency in our findings than in earlier reports with regard to the potential benefit of low glycaemic index or glycaemic load diets. Three systematic reviews have shown a reduced incidence of type 2 diabetes associated with the consumption of diets of lower glycaemic index or glycaemic load, though the effect was modest when compared with the protective effect of total dietary fibre or wholegrains. In the present study sensitivity analyses due to high heterogeneity reduced risk reduction and confidence intervals included 1. A review of prospective studies by Turati et al. suggested a small but significant increase in colorectal cancer incidence associated with high glycaemic index or glycaemic load. This finding was subject to high unexplained heterogeneity and included retrospective case-control studies which may be subject to dietary recall bias. Other studies have reported a lower incidence of stroke and CHD amongst those consuming low glycaemic index or glycaemic load diets, whilst we found a reduced risk of stroke only. We were unable to confirm an effect of low glycaemic index or glycaemic load diets on haemoglobin A1c or blood cholesterol which have been reported in many short and medium term trials. However we excluded trials which involved only people with diabetes or marked hyperlipidaemia who were the participants in the majority of trials reporting reduction in these important risk indicators. Our study does not exclude the value of these indicators of carbohydrate quality in this clinical context.

Whole grains offer a useful means of increasing dietary fibre intake and reducing risk of NCDs. However fruit and vegetables are also important contributors to dietary fibre intake. We did not specifically explore the
relationship between fruit and vegetable consumption and NCDs given the recent systematic review and meta analyses by Aune et al. They report risk reductions of around 10% per 200g fruit and vegetables combined for CHD, stroke and total mortality and smaller but still significant reductions for total cardiovascular disease and cancer. Appreciable dose response effects were apparent for most outcomes up to 800g/day. Inverse associations were observed between the intake of apples, pears, citrus fruits, green leafy vegetables, cruciferous vegetables and salad and cardiovascular disease and all-cause mortality. Intake of green yellow vegetables and cruciferous vegetables were inversely associated with total cancer risk. In addition to fibre, fruits and vegetables contain many other nutrients that are potentially protective and confer some risk reduction.

The benefits of fibre reported in the present and other papers are supported by over 100 years of research into its chemistry, physical properties, physiology and metabolic effects. Fibre containing foods must be chewed before passing through the stomach and into small bowel where they affect satiety, glucose and insulin responses and lipid absorption. Whilst more recent systematic reviews have shown only small effects on appetite, satiety or blood lipids these studies have been conducted largely using defined fibre supplements rather than whole foods. Whole foods that require chewing and retain much of their structure in the gut are more likely to increase satiety through a variety of mechanisms leading to weight loss and to modulation of carbohydrate and lipid metabolism. In the large bowel fibre is almost completely broken down by the resident microflora in a series of anaerobic reactions known as fermentation. The gut microbiota play a number of important roles in human health including protecting against pathogens, development of the gut immune system, vitamin synthesis, metabolism of xenobiotics and may be involved in complex gut-brain communication. However the principal function of the microbiome is digestion of fibre and other carbohydrates that escape breakdown in the small bowel and it is the availability of fibre in the diet that dominates the metabolism of the gut microbiome and leads to protection from conditions such as colorectal cancer. This coming together of the epidemiological and experimental work on fibre allows conclusions to be drawn that increased fibre intakes should result in improvements in population health.

While we have not considered the evidence regarding total carbohydrate intake, epidemiological evidence and relatively long term clinical trials suggest that a wide range of intakes is acceptable, a finding which is endorsed by authoritative dietary guidelines. Our study contributes to the growing body of evidence that carbohydrate quality rather than quantity determines major health outcomes. Translating these findings regarding dietary fibre and whole grains into dietary advice for individuals and populations should be accompanied by a caveat. Dietary fibre as defined by Codex Alimentarius is naturally occurring in foods, but may be extracted from foods or synthesised and added into manufactured foods. The large body of literature which contributed to this and other systematic reviews and meta analyses relate principally to fibre rich foods since most of the studies were undertaken before synthetic and extracted fibre were widely used. The concept of wholegrain foods has also changed appreciably. Wholegrain foods are required to have a nutrient composition similar to that of the original grain, without regard to the degree of processing. Many breakfast cereals and other manufactured “whole grain” products are more highly processed than in the past. There is limited, but quite striking evidence that increased processing of whole grains can result in a deterioration of a several biomarkers of cardiometabolic disease. As these are relatively recent developments there is no epidemiological evidence
of the consequences of these changes in the food supply on clinical outcomes and mortality. Until such evidence is available it seems appropriate that dietary advice should emphasise the benefits of naturally occurring dietary fibre in whole grains, vegetables and fruits that have been minimally processed.

There is a considerable body of evidence relating to the adverse consequences of high intakes of sugar sweetened beverages and strong recommendations to reduce the intake of free sugars. Our findings based on a series of systematic reviews and meta analyses provide strong evidence for the importance of including advice regarding the nature and source of other carbohydrates in dietary guidelines aimed at reducing the risk of NCDs. Diets high in total dietary fibre especially from whole grains and, on the basis of another recent systematic review, vegetables and fruits, are associated with a significant reduction in a range of NCDs when compared with lower dietary fibre and refined rather than whole grain intakes. The types of studies we have considered did not identify risks associated with dietary fibre. However high intakes may be associated with deleterious effects in populations with borderline iron/mineral status, amongst whom very high whole grain intakes may further compromise iron status. High intakes of dietary fibre and whole grains are more clearly associated with good health outcomes than measures of glycaemic index or glycaemic load. Whilst glycaemic index provides a measure of the glycaemic potential of the carbohydrate content of foods, some low glycaemic index foods may have other attributes, which are not health promoting. Foods containing added fructose or sucrose and composite foods containing both saturated fat and carbohydrate (e.g. confectionary products) may have a low glycaemic index. Our complementary findings from randomised controlled trials and prospective studies together with the dose response effects, supported by much experimental work, are evidence that diets characterised by a low content of dietary fibre contribute to the cause of a number of NCDs and that benefit will accrue from implementation of quantitative recommendations regarding dietary fibre intake. Intakes in the range of 25 to 29g daily are optimal whilst the dose response data show that amounts greater than 30g per day confer additional benefits.

**Contributors**

AR was responsible for the systematic reviews and meta analyses of prospective studies, wrote the first draft of the manuscript, was involved with the interpretation of results, and approved the submission of the final manuscript. LTM was responsible for the systematic reviews and meta analyses of clinical trials, was involved with the interpretation of results, and approved the submission of the final manuscript. JC was involved with the interpretation of results, and approved the submission of the final manuscript. NW was involved with the systematic reviews and meta analyses of clinical trials and approved the submission of the final manuscript. EM was involved with the systematic reviews and meta analyses of prospective studies and approved the submission of the final manuscript. JM was involved with the interpretation of results, had full access to all the data in the study and had final responsibility for the decision to submit for publication.

**Declarations of interest**

We declare no competing interests

**Acknowledgments**
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References


Figure 1 Flow chart indicating the process by which eligible prospective studies and randomised controlled trials were identified.

Prospective Observational Studies (POS)

1,458 systematic review and meta analyses (SRMA) identified up to February 2017

- 1,384 SRMA excluded based on title or abstract

- 74 SRMA for full assessment

- 56 SRMA eligible for this review

- 246 POS identified from SRMA and 10,653 publications identified in searches up to April 2017

- 10,221 publications excluded on title/abstract

- 678 for full assessment

- 493 publications excluded not eligible

- 185 POS eligible for this review

Randomised controlled trials (RCT)

10,245 publications identified up to February 2018

- 9,967 publications excluded on title/abstract

- 278 for full assessment

- 220 publications excluded not eligible

- 58 RCT eligible for this review
<table>
<thead>
<tr>
<th>Outcome</th>
<th>Number of studies</th>
<th>Type of study</th>
<th>Number of cases or N in intervention</th>
<th>Person years or N of controls</th>
<th>Effect size (95%CI)</th>
<th>GRADE quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>All-cause mortality</td>
<td>10</td>
<td>observational</td>
<td>80,139</td>
<td>12·3 million PY</td>
<td>RR 0·85 (0·79 to 0·91)</td>
<td>Moderate</td>
</tr>
<tr>
<td>CHD mortality</td>
<td>10</td>
<td>observational</td>
<td>7,243</td>
<td>6·9 million PY</td>
<td>RR 0·69 (0·60 to 0·81)*</td>
<td>Moderate</td>
</tr>
<tr>
<td>CHD incidence</td>
<td>9</td>
<td>observational</td>
<td>7,155</td>
<td>2·7 million PY</td>
<td>RR 0·76 (0·69 to 0·83)</td>
<td>Moderate</td>
</tr>
<tr>
<td>Stroke mortality</td>
<td>2</td>
<td>observational</td>
<td>1,103</td>
<td>1·3 million PY</td>
<td>RR 0·80 (0·56 to 1·14)</td>
<td>Very low</td>
</tr>
<tr>
<td>Stroke incidence</td>
<td>9</td>
<td>observational</td>
<td>13,134</td>
<td>4·6 million PY</td>
<td>RR 0·78 (0·69 to 0·88)**</td>
<td>Low</td>
</tr>
<tr>
<td>Cancer mortality</td>
<td>5</td>
<td>observational</td>
<td>29,593</td>
<td>11·2 million PY</td>
<td>RR 0·87 (0·79 to 0·95)</td>
<td>Moderate</td>
</tr>
<tr>
<td>Type 2 diabetes incidence</td>
<td>17</td>
<td>observational</td>
<td>48,468</td>
<td>6·9 million PY</td>
<td>RR 0·84 (0·78 to 0·90)</td>
<td>Moderate</td>
</tr>
<tr>
<td>Colorectal cancer incidence</td>
<td>22</td>
<td>observational</td>
<td>22,920</td>
<td>16·9 million PY</td>
<td>RR 0·84 (0·78 to 0·89)</td>
<td>Moderate</td>
</tr>
<tr>
<td>Change in body weight (kg)</td>
<td>27</td>
<td>randomised trials</td>
<td>1,294</td>
<td>1201</td>
<td>MD 0·37 lower (0·63 kg lower to 0·11 kg lower)</td>
<td>High</td>
</tr>
<tr>
<td>Change in HbA1c (%)</td>
<td>6</td>
<td>randomised trials</td>
<td>191</td>
<td>189</td>
<td>SMD 0·35 lower (0·73 lower to 0·03 higher)</td>
<td>Low</td>
</tr>
<tr>
<td>Change in total cholesterol (mmol/L)</td>
<td>36</td>
<td>randomised trials</td>
<td>1,832</td>
<td>1,671</td>
<td>MD 0·15 lower** (0·22 lower to 0·07 lower)</td>
<td>Moderate</td>
</tr>
<tr>
<td>Change in systolic blood pressure (mm Hg)</td>
<td>15</td>
<td>randomised trials</td>
<td>1,064</td>
<td>988</td>
<td>MD 1·27 lower** (2·50 lower to 0·04 lower)</td>
<td>Moderate</td>
</tr>
</tbody>
</table>

PY person years, RR relative risk, MD mean difference, SMD standardised mean difference
Egger’s test for bias p = 0·0040. Trim and fill analysis did not change the direction or significance of the pooled estimate.
The high heterogeneity of the pooled effect size (>50%) is unexplained by sensitivity analyses.
Detailed justification for the GRADE quality of evidence is given in Appendix B for observational studies and Supplement 1 for trials.
<table>
<thead>
<tr>
<th>Outcome</th>
<th>N of studies</th>
<th>Type of study</th>
<th>Number of cases or N in intervention</th>
<th>Person years or N of controls</th>
<th>Effect size (95%CI)</th>
<th>GRADE quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>All-cause mortality</td>
<td>9</td>
<td>observational</td>
<td>99,224</td>
<td>10·7 million PY</td>
<td>RR 0·81 (0·72 to 0·90)*</td>
<td>Low</td>
</tr>
<tr>
<td>CHD mortality</td>
<td>2</td>
<td>observational</td>
<td>1,588</td>
<td>2·0 million PY</td>
<td>RR 0·66 (0·56 to 0·77)</td>
<td>Low</td>
</tr>
<tr>
<td>CHD incidence</td>
<td>6</td>
<td>observational</td>
<td>7,697</td>
<td>2·8 million PY</td>
<td>RR 0·80 (0·70 to 0·91)*</td>
<td>Low</td>
</tr>
<tr>
<td>Stroke mortality</td>
<td>2</td>
<td>observational</td>
<td>694</td>
<td>2·0 million PY</td>
<td>RR 0·74 (0·58 to 0·94)</td>
<td>Low</td>
</tr>
<tr>
<td>Stroke incidence</td>
<td>3</td>
<td>observational</td>
<td>1,247</td>
<td>1·1 million PY</td>
<td>RR 0·86 (0·61 to 1·21)</td>
<td>Very low</td>
</tr>
<tr>
<td>Cancer mortality</td>
<td>5</td>
<td>observational</td>
<td>32,727</td>
<td>10·1 million PY</td>
<td>RR 0·84 (0·76 to 0·92)*</td>
<td>Low</td>
</tr>
<tr>
<td>Type 2 diabetes incidence</td>
<td>8</td>
<td>observational</td>
<td>14,686</td>
<td>3·9 million PY</td>
<td>RR 0·67 (0·58 to 0·78)*</td>
<td>Low</td>
</tr>
<tr>
<td>Colorectal cancer incidence</td>
<td>7</td>
<td>observational</td>
<td>8,803</td>
<td>6·8 million PY</td>
<td>RR 0·87 (0·79 to 0·96)</td>
<td>Moderate</td>
</tr>
<tr>
<td>Change in body weight (kg)</td>
<td>11</td>
<td>randomised trials</td>
<td>498</td>
<td>421</td>
<td>MD 0·62 lower (1·19 lower to 0·05 lower)</td>
<td>Moderate</td>
</tr>
<tr>
<td>Change in HbA1c (%)</td>
<td>3</td>
<td>randomised trials</td>
<td>141</td>
<td>141</td>
<td>SMD 0·54 lower (1·28 lower to 0·20 higher)</td>
<td>Low</td>
</tr>
<tr>
<td>Change in total cholesterol (mmol/L)</td>
<td>17</td>
<td>randomised trials</td>
<td>772</td>
<td>701</td>
<td>MD 0·09 lower (0·23 lower to 0·04 higher)</td>
<td>Moderate</td>
</tr>
<tr>
<td>Change in systolic blood pressure (mm Hg)</td>
<td>8</td>
<td>randomised trials</td>
<td>493</td>
<td>432</td>
<td>MD 1·01 lower (2·46 lower to 0·44 higher)</td>
<td>Moderate</td>
</tr>
</tbody>
</table>

PY person years, RR relative risk, MD mean difference, SMD standardised mean difference

* The high heterogeneity of the pooled effect size (>50%) is unexplained by sensitivity analyses.

Detailed justification for the GRADE quality of evidence is given in Appendix C for observational studies and Supplement 2 for trials.
<table>
<thead>
<tr>
<th>Outcome</th>
<th>N of studies</th>
<th>Type of study</th>
<th>Number of cases or N in intervention</th>
<th>Person years or N of control</th>
<th>Effect size (95%CI)</th>
<th>GRADE quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>All-cause mortality</td>
<td>3</td>
<td>observational</td>
<td>7,698</td>
<td>0·6 million PY</td>
<td>RR 0·89 (0·70 to 1·13)*</td>
<td>Very low</td>
</tr>
<tr>
<td>CHD mortality</td>
<td>1</td>
<td>observational</td>
<td>incidence not stated</td>
<td>0·04 million PY</td>
<td>RR 1·10 (0·69 to 1·75)</td>
<td>Very low</td>
</tr>
<tr>
<td>CHD incidence</td>
<td>10</td>
<td>observational</td>
<td>8,456 + not reported in one study</td>
<td>2·4 million PY</td>
<td>RR 0·93 (0·83 to 1·04)</td>
<td>Low</td>
</tr>
<tr>
<td>Stroke mortality</td>
<td>3</td>
<td>observational</td>
<td>951</td>
<td>1·2 million PY</td>
<td>RR 0·63 (0·52 to 0·77)</td>
<td>Low</td>
</tr>
<tr>
<td>Stroke incidence**</td>
<td>5</td>
<td>observational</td>
<td>5,527</td>
<td>3·0 million PY</td>
<td>RR 0·84 (0·72 to 0·99)</td>
<td>Very low</td>
</tr>
<tr>
<td>Cancer mortality</td>
<td>1</td>
<td>observational</td>
<td>1,401</td>
<td>0·4 million PY</td>
<td>RR 1·11 (0·90 to 1·38)</td>
<td>Very low</td>
</tr>
<tr>
<td>Type 2 diabetes incidence**</td>
<td>14</td>
<td>observational</td>
<td>36,908</td>
<td>6·5 million PY</td>
<td>RR 0·89 (0·82 to 0·97)*</td>
<td>Very low</td>
</tr>
<tr>
<td>Colorectal cancer incidence</td>
<td>10</td>
<td>observational</td>
<td>11,245</td>
<td>8·8 million PY</td>
<td>RR 0·91 (0·82 to 0·97)*</td>
<td>Very low</td>
</tr>
<tr>
<td>Change in body weight (kg)</td>
<td>8</td>
<td>randomised trials</td>
<td>464</td>
<td>335</td>
<td>MD 0·29 lower (0·62 lower to 0·03 higher)</td>
<td>High</td>
</tr>
<tr>
<td>Change in HbA1c (%)</td>
<td>2</td>
<td>randomised trials</td>
<td>44</td>
<td>37</td>
<td>SMD 0·08 higher (0·35 lower to 0·52 higher)</td>
<td>Very low</td>
</tr>
<tr>
<td>Change in total cholesterol (mmol)</td>
<td>8</td>
<td>randomised trials</td>
<td>605</td>
<td>478</td>
<td>MD 0·02 lower (0·17 lower to 0·13 higher)</td>
<td>Moderate</td>
</tr>
<tr>
<td>Change in systolic blood pressure (mm Hg)</td>
<td>4</td>
<td>randomised trials</td>
<td>519</td>
<td>397</td>
<td>MD 0·17 lower (1·03 lower to 0·69 higher)</td>
<td>High</td>
</tr>
</tbody>
</table>

* PY person years, RR relative risk, MD mean difference, SMD standardised mean difference

* The high heterogeneity of the pooled effect size (>50%) is unexplained by sensitivity analyses.

** The pooled effect size did not maintain statistical significance during sensitivity analyses.

*** Only one eligible trial of children was identified in our systematic searches. Although the exposure was for diets of higher and lower GI, data from this trial has not been included with that of adults shown above.

Detailed justification for the GRADE quality of evidence is given in Appendix D for observational studies and Supplement 2 for trials.
**Figure 2** Dose response relationships between total dietary fibre and critical clinical outcomes based on data from prospective studies.

**Fig 2a** Total fibre and all-cause mortality. 68,183 deaths over 11.3 million person years. Assuming linearity a reduced risk ratio of 0.93 (0.90 to 0.95) was observed for every 8 grams more fibre consumed per day.

**Fig 2b** Total fibre and coronary heart disease incidence. 6,449 deaths over 2.5 million person years. Assuming linearity a reduced risk ratio of 0.81 (0.73 to 0.90) was observed for every 8 grams more fibre consumed per day.

**Fig 2c** Total fibre and incidence of type 2 diabetes. 22,450 cases over 3.2 million person years. Assuming linearity a reduced risk ratio of 0.85 (0.82 to 0.89) was observed for every 8 grams more fibre consumed per day.

**Fig 2d** Total fibre and incidence of colorectal cancer. 20,009 cases over 20.9 million person years. Assuming linearity a reduced risk ratio of 0.92 (0.89 to 0.95) was observed for every 8 grams more fibre consumed per day.
Figure 3 Dose response relationships between wholegrain intake and critical clinical outcomes based on data from prospective studies.

**Fig 3a** Wholegrain intake and all-cause mortality. 88,347 deaths over 8.2 million person years. Assuming linearity a reduced risk ratio of 0.94 (0.92 to 0.95) was observed for every 15 grams more whole grains consumed per day.

**Fig 3b** Wholegrain intake and incidence of coronary heart disease. 6,587 cases over 2.4 million person years. Assuming linearity a reduced risk ratio of 0.93 (0.89 to 0.98) was observed for every 15 grams more whole grains consumed per day.

**Fig 3c** Wholegrain intake and incidence of type 2 diabetes. 13,147 cases over 3.5 million person years. Assuming linearity a reduced risk ratio of 0.88 (0.81 to 0.95) was observed for every 15 grams more whole grains consumed per day.

**Fig 3d** Wholegrain intake and incidence of colorectal cancer. 6,056 cases over 5.7 million person years. Assuming linearity a reduced risk ratio of 0.97 (0.95 to 0.99) was observed for every 15 grams more whole grains consumed per day.
**Figure 4** Dose response relationships between dietary glycaemic index and critical clinical outcomes based on data from prospective studies.

**Fig 4a** Glycaemic index and all-cause mortality. 7,699 deaths over 0.6 million person years. Assuming linearity a risk ratio of 1.16 (0.90 to 1.49) was observed for every 10 glycaemic index unit increase per day.

**Fig 4b** Glycaemic index and coronary heart disease incidence. 7,240 cases over 2.4 million person years. Assuming linearity a risk ratio of 1.09 (0.94 to 1.27) was observed for every 10 glycaemic index unit increase per day.

**Fig 4c** Glycaemic index and incidence of type 2 diabetes. 31,780 cases over 4.9 million person years. Assuming linearity a risk ratio of 1.10 (1.00 to 1.20) was observed for every 10 glycaemic index unit increase per day.

**Fig 4d** Glycaemic index and incidence of colorectal cancer. 10,390 cases over 6.5 million person years. Assuming linearity a risk ratio of 1.05 (1.00 to 1.10) was observed for every 10 glycaemic index unit increase per day.
**Figure 5** Summary forest plots of key outcomes from clinical trials.

**Fig 5a** Higher compared with lower total fibre intakes.

<table>
<thead>
<tr>
<th>Comparison</th>
<th>Total dietary fibre</th>
<th></th>
<th></th>
<th></th>
<th>Pooled Effect Estimates</th>
<th>Mean difference (95% CIs)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Studies</td>
<td>Intervention</td>
<td>Control</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Body weight (kg)</td>
<td>27</td>
<td>1294</td>
<td>1201</td>
<td>-0.37 (-0.63 to -0.11)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total cholesterol (mmol/L)</td>
<td>36</td>
<td>1832</td>
<td>1671</td>
<td>-0.15 (-0.22 to -0.07)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LDL cholesterol (mmol/L)</td>
<td>34</td>
<td>1801</td>
<td>1640</td>
<td>-0.09 (-0.15 to -0.04)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Triglycerides (mmol/L)</td>
<td>31</td>
<td>1700</td>
<td>1560</td>
<td>-0.06 (-0.11 to -0.01)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Systolic blood pressure (mm Hg)</td>
<td>15</td>
<td>1064</td>
<td>988</td>
<td>-1.27 (-2.59 to -0.04)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HbA1c (SMD)</td>
<td>6</td>
<td>191</td>
<td>189</td>
<td>-0.35 (-0.73 to 0.03)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fasting glucose (mmol/L)</td>
<td>39</td>
<td>1716</td>
<td>1547</td>
<td>-0.09 (-0.15 to -0.02)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Insulin resistance (SMD)</td>
<td>14</td>
<td>672</td>
<td>591</td>
<td>-0.12 (-0.26 to 0.03)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Figure 5b** Higher compared with lower wholegrain intakes.

<table>
<thead>
<tr>
<th>Comparison</th>
<th>Whole grains</th>
<th></th>
<th></th>
<th></th>
<th>Pooled Effect Estimates</th>
<th>Mean difference (95% CIs)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Studies</td>
<td>Intervention</td>
<td>Control</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Body weight (kg)</td>
<td>11</td>
<td>498</td>
<td>421</td>
<td>-0.62 (-1.19 to -0.05)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total cholesterol (mmol/L)</td>
<td>17</td>
<td>772</td>
<td>701</td>
<td>-0.09 (-0.23 to 0.04)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LDL cholesterol (mmol/L)</td>
<td>16</td>
<td>894</td>
<td>743</td>
<td>-0.09 (-0.19 to 0.02)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Triglycerides (mmol/L)</td>
<td>17</td>
<td>915</td>
<td>764</td>
<td>-0.01 (-0.05 to 0.03)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Systolic blood pressure (mm Hg)</td>
<td>8</td>
<td>413</td>
<td>342</td>
<td>-1.01 (-2.46 to 0.44)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HbA1c (SMD)</td>
<td>3</td>
<td>141</td>
<td>141</td>
<td>-0.54 (-1.38 to 0.20)</td>
<td></td>
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</tr>
<tr>
<td>Fasting glucose (mmol/L)</td>
<td>18</td>
<td>837</td>
<td>687</td>
<td>-0.05 (-0.12 to 0.02)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Insulin resistance (SMD)</td>
<td>7</td>
<td>284</td>
<td>283</td>
<td>-0.20 (-0.49 to 0.08)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Figure 5c** Comparison of diets characterised by lower compared with higher glycaemic index foods.

<table>
<thead>
<tr>
<th>Comparison</th>
<th>Glycaemic Index</th>
<th></th>
<th></th>
<th></th>
<th>Pooled Effect Estimates</th>
<th>Mean difference (95% CIs)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Studies</td>
<td>Intervention</td>
<td>Control</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Body weight (kg)</td>
<td>8</td>
<td>464</td>
<td>335</td>
<td>-0.20 (-0.62 to 0.03)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total cholesterol (mmol/L)</td>
<td>8</td>
<td>605</td>
<td>478</td>
<td>-0.02 (-0.17 to 0.13)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LDL cholesterol (mmol/L)</td>
<td>8</td>
<td>605</td>
<td>478</td>
<td>0.05 (+0.13 to 0.22)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Triglycerides (mmol/L)</td>
<td>8</td>
<td>605</td>
<td>478</td>
<td>-0.02 (-0.07 to 0.03)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Systolic blood pressure (mm Hg)</td>
<td>4</td>
<td>519</td>
<td>397</td>
<td>-0.17 (-1.03 to 0.69)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HbA1c (SMD)</td>
<td>2</td>
<td>44</td>
<td>37</td>
<td>0.08 (-0.35 to 0.52)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fasting glucose (mmol/L)</td>
<td>11</td>
<td>609</td>
<td>475</td>
<td>0 (+0.08 to 0.07)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Insulin resistance (SMD)</td>
<td>5</td>
<td>284</td>
<td>283</td>
<td>0.14 (+0.03 to 0.31)</td>
<td></td>
<td></td>
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</tbody>
</table>