Efficacy and safety of low and very low carbohydrate diets for type 2 diabetes remission: systematic review and meta-analysis of published and unpublished randomized trial data

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ABSTRACT

OBJECTIVE
To determine the efficacy and safety of low carbohydrate diets (LCDs) and very low carbohydrate diets (VLCDs) for people with type 2 diabetes.

DESIGN
Systematic review and meta-analysis.

DATA SOURCES
Searches of CENTRAL, Medline, Embase, CINAHL, CAB, and grey literature sources from inception to 25 August 2020.

STUDY SELECTION
Randomized clinical trials evaluating LCDs (<130 g/day or <26% of a 2000 kcal/day diet) and VLCDs (<10% calories from carbohydrates) for at least 12 weeks in adults with type 2 diabetes were eligible.

DATA EXTRACTION
Primary outcomes were remission of diabetes (HbA1c <6.5% or fasting glucose <7.0 mmol/L, with or without the use of diabetes medication), weight loss, HbA1c, fasting glucose, and adverse events. Secondary outcomes included health related quality of life and biochemical laboratory data. All articles and outcomes were independently screened, extracted, and assessed for risk of bias and GRADE certainty of evidence at six and 12 month follow-up. Risk estimates and 95% confidence intervals were calculated using random effects meta-analysis.

RESULTS
Outcomes were assessed according to a priori determined minimal important differences to determine clinical importance, and heterogeneity was investigated on the basis of risk of bias and seven a priori subgroups. Any subgroup effects with a statistically significant test of interaction were subjected to a five point credibility checklist.

WHAT THIS STUDY ADDS
This systematic review of the effect of LCDs on remission of type 2 diabetes included 23 trials, including unpublished HbA1c, and medication use data from five trials.

WHAT IS ALREADY KNOWN ON THIS TOPIC
Previous systematic reviews have used broad definitions of low carbohydrate (eg, <45% of calories from carbohydrates) and have not systematically assessed remission of diabetes.

Results from reviews based on a subgroup of 10 randomized trials assessing low carbohydrate diets (LCDs) (<26-45% of daily calories from carbohydrate) have been encouraging.

CONCLUSIONS
On the basis of moderate to low certainty evidence, patients adhering to an LCD for six months may experience remission of diabetes without adverse consequences. Limitations include continued debate around what constitutes remission of diabetes, as well as the efficacy, safety, and dietary satisfaction of longer term LCDs.

PROSPERO CRD42020161795.
Introduction
Diabetes is a common, deadly, and expensive medical condition. It is estimated that 1 in 11 adults worldwide have diabetes and that it is responsible for 11% of deaths annually, costing $760bn (£570bn; €626bn) in direct costs alone. Type 2 diabetes is the most common form of diabetes, accounting for 90-95% of cases, and for decades has been a rapidly growing international concern. Type 2 diabetes is characterized by insulin resistance driven by chronic hyperglycemia and is commonly diagnosed by measures of glycemia such as fasting blood glucose concentrations of 7.0 mmol/L or above or glycated hemoglobin (HbA1c) values of 6.5% (48 mmol/mol) or above. It is associated with several risk factors including genetics and lifestyle influences, but by far the most common risk factor is obesity.

Structured dietary interventions are commonly recommended for patients with diabetes, with varied recommendations from authoritative organizations. Before the discovery of insulin, diets emphasizing carbohydrate restriction had been used extensively in the management of diabetes, but more recently they have fallen out of favor. Because a key underlying mechanism of type 2 diabetes is insulin resistance driven in part by chronic hyperglycemia, lowering dietary intake of carbohydrate, most of which is absorbed as glucose or fructose, has been suggested to improve blood glucose control and outcomes of type 2 diabetes. Structured diets with carbohydrate restriction have been variably described in the research literature but have been commonly grouped into three categories: 20-50 g/day carbohydrates or less than 10% of the 2000 kcal/day diet that is generally sufficient to induce ketosis; less than 130 g/day or less than 26% of the 2000 kcal/day diet; and less than 45% of the 2000 kcal/day diet. For the purposes of this review, we refer to diets with less than 130 g/day or less than 26% of calories from carbohydrates based on 2000 kcal/day as a low carbohydrate diet (LCD).

Type 2 diabetes remains a significant and worsening problem worldwide, despite many pharmaceutical developments and a global emphasis on glycemic control. Structured diets are recognized as an essential component of treating diabetes, but confusion remains about which diet to choose. Systematic reviews and meta-analyses to date have attempted to pool carbohydrate restricted diets for diabetic populations, reporting mixed results.

Type 2 diabetes remains significant and worsening problem worldwide, despite many pharmaceutical developments and a global emphasis on glycemic control. Structured diets are recognized as an essential component of treating diabetes, but confusion remains about which diet to choose. Systematic reviews and meta-analyses to date have attempted to pool carbohydrate restricted diets for diabetic populations, reporting mixed results. }

Methods

Search strategy and selection criteria
On the basis of an a priori and publicly available protocol (PROSPERO CRD42020161795), we did a systematic review with meta-analysis of randomized controlled trials assessing the efficacy and safety of LCDs among adult patients with a diagnosis of type 2 diabetes. We included people with or without cardiovascular conditions regardless of medication use or glucose concentration and HbA1c level.

We included trials comparing LCDs with any wait list controls or any active controls including competing dietary programs higher in carbohydrates (≥26%), with or without exercise, lifestyle, and behavioral recommendations. No language, date, or publication restrictions were applied. We sought unpublished data from investigators of published and unpublished trials.

To meet inclusion criteria, studies had to investigate allocation to an LCD (<26% calories from carbohydrates or <130 g/day) for a defined period (12 weeks or longer), with or without exercise (for example, walking, jogging, strength training) or lifestyle and behavioral recommendations (for example, cognitive therapy, group support). Primary outcomes of interest,
based on our a priori protocol,\textsuperscript{16} were remission of type 2 diabetes (dichotomously defined as HbA\textsubscript{1c} <6.5\% or fasting glucose <7.0 mmol/L), with or without the use of diabetes medication. Additional primary outcomes were weight loss, HbA\textsubscript{1c}, fasting glucose, and adverse events (total and serious adverse events). Secondary outcomes were health related quality of life, reduction of medication, and biochemical laboratory data including total cholesterol, low density lipoprotein cholesterol, high density lipoprotein cholesterol, triglycerides, homeostasis model assessment of insulin resistance (HOMA-IR), and inflammatory markers (C reactive protein).

We searched the following databases from inception to 25 August 2020 to identify studies: Cochrane Central Register of Controlled Trials (CENTRAL), Medline via PubMed, Embase, the Cumulative Index to Nursing and Allied Health Literature (CINAHL), and Commonwealth Agricultural Bureaux (CAB) abstracts. With the assistance of an expert clinical librarian, search strategies were customized, including the use of a Cochrane recommended filter for the identification of randomized controlled trials in PubMed.\textsuperscript{19} The Medline search strategy is reported in supplementary table A. On the basis of our study protocol, we also searched three trial registries (for example, clinicaltrials.gov) and four additional grey literature sources (for example, BIOSIS Citation Index, ProQuest Dissertations & Theses Global).\textsuperscript{16}

Two authors, independently and in duplicate, screened titles and abstracts and subsequently full text articles. Disagreements were resolved by consensus.

Data analysis
Data extraction was done independently and in duplicate using a pilot tested extraction form. Domains for extraction included study design factors, population, intervention, comparator, and surrogate and health outcomes (variables listed in supplementary table B). All outcomes were extracted and reported at six months (±3 months) and 12 months (±3 months). We used version 2.0 of the Cochrane Risk-of-Bias (RoB) instrument for randomized trials and assessed each of the RoB domains as “high,” “low,” or “some concern” using the Excel file provided by the RoB 2.0 development team.\textsuperscript{20}

We used Revman software (version 5.3) and the “meta” package in R (version 3.6.1) to do meta-analyses. For dichotomous outcomes, we calculated the pooled risk difference, risk ratio, and number needed to treat for an additional beneficial outcome (NNT) with 95\% confidence intervals. For continuous outcomes, we combined endpoint or change data; when both endpoint and change data were reported, we prioritized endpoint data.\textsuperscript{21} We calculated the pooled mean difference and/or standardized mean difference with corresponding 95\% confidence intervals. We pooled studies that measured continuous health related quality of life with different instruments if the underlying construct was the same or similar. To improve interpretability for readers, we followed published guidance and presented effect estimates in two ways.\textsuperscript{22} Firstly, we pooled the effect estimates as standardized mean differences. Secondly, we converted scores of the different health related quality of life instruments to units of the most commonly used reference instrument and presented the mean difference.\textsuperscript{22} Where possible, we presented the effect size on the basis of known or estimated minimal clinically important difference (MCID) thresholds for all outcomes (supplementary table C). We rated the overall certainty (quality) of evidence for each of our outcomes by using the GRADE (Grading of Recommendations Assessment, Development and Evaluation) approach, wherein randomized trials began as high certainty evidence but could be rated down by one or more levels on the basis of five categories of limitations: risk of bias, inconsistency, indirectness, imprecision, and publication bias.\textsuperscript{25} 26 We assessed the RoB and GRADE independently and in duplicate, with disagreement resolved by consensus. After a request from referees, we also did a sensitivity analysis comparing the certainty of evidence using GRADE versus NutriGRADE.\textsuperscript{27}

Following published guidance, we chose to use data from complete cases for our primary analysis.\textsuperscript{28} When studies had missing outcome data and reported a complete case analysis, we did sensitivity analyses and applied increasingly stringent but plausible assumptions to this data,\textsuperscript{24} 28 29 using Excel files made available from the authors of the GRADE guidance on missing outcome data.\textsuperscript{24} For assessing the effect of missing outcome data on risk of bias, we did these sensitivity assessments at the study level to best integrate with Cochrane RoB 2.0.\textsuperscript{20}

We assessed and reported heterogeneity quantitatively using the I\textsuperscript{2} statistic and did a χ\textsuperscript{2} test for homogeneity according to guidelines from the Cochrane Handbook (for example, 50\% to 90\% may represent substantial heterogeneity; 75\% to 100\% may represent considerable heterogeneity).\textsuperscript{30}

We investigated heterogeneity and the possibility of effect modification for our primary outcomes on the basis of risk of bias and seven a priori subgroups,\textsuperscript{16} with any subgroup effects with a statistically significant test of interaction subjected to a five point credibility checklist.\textsuperscript{31} Subgroups were very low carbohydrate diets (VLCD) (<10\% calories from carbohydrates) versus diets with between 10\% and 26\% of calories from carbohydrates; trials that provided behavioral support versus those that did not; LCDs versus comparator diets (for example, low fat diets, Mediterranean diets); trials in which caloric intake did not significantly differ between groups (iso-caloric) versus those in which it did; LCD trials that used caloric restriction versus those that did not; trials that included patients who used insulin versus those that did not; trials in which the intervention group showed adequate adherence (determined by three a priori criteria: 3-β-hydroxybutyrate, measured carbohydrate intake, and author definitions) versus those that did not. Furthermore, for each outcome, we investigated the effect on the point estimate when we
restricted the analysis to studies at low risk of bias; if the risk of bias sensitivity analysis was credible, we focused our results on those studies at low risk.

To assess for the possibility of publication bias, we visually inspected funnel plots when 10 or more trials were included. We further assessed for publication bias by using Egger’s regression test for continuous outcomes and the Harbord score for dichotomous outcomes.32 33

**Patient and public involvement**

Given the nature of secondary data capture and analysis, patients and the public were not involved in the design or interpretation of this study.

**Results**

Our search yielded 14 759 records, of which 23 studies (1357 participants) met the inclusion criteria (fig 1). Table 1 shows characteristics of the clinical trials. In short, trials primarily included overweight and obese patients with type 2 diabetes, with 14/23 (61%) studies including participants using insulin. Trial size ranged from 12 to 144 participants with a mean age range of 47 to 67 years. Studies used various carbohydrate restriction thresholds with 12/23 (52%) meeting our criteria for very low carbohydrate diets (<10% daily calories from carbohydrates or <50 g/d). Trials primarily used low fat diets as control comparators (18/23; 78%). Duration of treatment ranged from three months to two years. Dropouts were common in the included studies. Eighteen (78%) of 23 studies reported missing participant outcome data, with 10 reporting more than 20% of data being missing. In studies with reported missing data, we assessed the robustness of reported effect estimates by using increasingly stringent assumptions about the missing data and incorporated this into the overall assessment for risk of bias.24 Overall, 59.4% of outcomes were rated as having some concern or high risk of bias, and 40.6% of outcomes were rated as having low risk of bias (fig 2). The randomization process was the risk of bias domain that had the poorest reporting, with just over 40% of trials having “some concerns.”

Eight studies reported on remission of diabetes at six months.34-41 Pooled analysis showed that when remission was defined by an HbA1c level below 6.5% independent of medication use, LCDs increased remissions by an additional 32 per 100 patients followed (risk difference 0.32, 95% confidence interval 0.17 to 0.47; 8 studies, n=264; GRADE=moderate) (fig 3; table 2). When remission was defined by an HbA1c level below 6.5% and the absence of diabetes medication, LCDs increased remissions at a lower rate (risk difference 0.05, –0.05 to 0.14; 5 studies, n=199; GRADE=low) (table 2). Three studies reported on remission at 12 months.35 39 41 When remission was defined independently of medication use, LCDs increased remission (risk difference 0.10, –0.02 to 0.21; 3 studies, n=171; GRADE=moderate), but they lowered the remission rate when the definition of remission included absence of diabetes medication (risk difference –0.04, –0.16 to 0.09; 2 studies, n=126; GRADE=low) (table 2).

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**Fig 1 | PRISMA study flow**
Table 1 | Characteristics of included trials

<table>
<thead>
<tr>
<th>Trial</th>
<th>BMI; HbA1c range; insulin dependence inclusive</th>
<th>Mean age (years)</th>
<th>Carbohydrate intake; caloric restriction; intense behavioral support</th>
<th>Comparator (diet, iso-caloric)</th>
<th>Adherence</th>
<th>Missing participant outcome data</th>
<th>Duration of intervention (months)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dyson et al (2010)54</td>
<td>BMI &gt;25; did not include patients receiving insulin (n=25)</td>
<td>54</td>
<td>&lt;40 g/d; caloronically restricted; intense behavioral support</td>
<td>Low fat Adequate</td>
<td>17%</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>Morris et al (2019)55</td>
<td>BMI &gt;30; did not include patients receiving insulin (n=33)</td>
<td>69/64, 55% female</td>
<td>&lt;26% carbohydrate/d; calorically restricted</td>
<td>Low fat Adequate</td>
<td>0%</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>Saslow et al (2014)56</td>
<td>BMI ≥25; did not include patients receiving insulin (n=34)</td>
<td>65/55, 74% female</td>
<td>“A very low carbohydrate, high fat, non caloric-restricted diet whose goal is to induce nutritional ketosis”, intense behavioral support</td>
<td>Low fat, iso-caloric (per intake not per goal) Adequate</td>
<td>11%</td>
<td>12</td>
<td></td>
</tr>
<tr>
<td>Saslow et al (2017)57</td>
<td>BMI≥25; did not include patients receiving insulin (n=25)</td>
<td>53/58, 56% female</td>
<td>20-50 g/d</td>
<td>Low fat, iso-caloric Adequate 24%</td>
<td>8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yamada et al (2014)58</td>
<td>Did include patients receiving insulin (n=24)</td>
<td>63/63, 50% female</td>
<td>70-130 g/d</td>
<td>Low fat, iso-caloric (per intake not per goal) Adequate</td>
<td>0%</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>Tay et al (2014)59</td>
<td>BMI 26-45; HbA1c ≥7.0% or taking diabetes medication; did include patients receiving insulin (n=131)</td>
<td>58/58, 37% female</td>
<td>50-70 g/d target; calorically restricted; intense behavioral support</td>
<td>Low glycemic index; iso-caloric Adequate</td>
<td>29%</td>
<td>24</td>
<td></td>
</tr>
<tr>
<td>Jonsson et al (2009)60</td>
<td>HbA1c &gt;5.5%; did not include patients receiving insulin (n=17)</td>
<td>66/63, 24% female</td>
<td>&lt;130 g/d</td>
<td>Low fat Adequate 18%</td>
<td>6 (crossover study; first 3 month comparison used in analysis)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sato et al (2017)61</td>
<td>BMI &gt;23; HbA1c &gt;7.5%; did include patients receiving insulin (n=66)</td>
<td>61/58, 23% female</td>
<td>130 g/d target</td>
<td>Low fat Adequate</td>
<td>6%</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>Goldstein et al (2011)62</td>
<td>BMI 30-40; HbA1c &gt;7.0%; did not include patients receiving insulin (n=52)</td>
<td>57/55, 48% female</td>
<td>25-40 g/d; intense behavioral support</td>
<td>Low fat (ADA 2000); iso-caloric Adherent at VLCD level</td>
<td>42%</td>
<td>12</td>
<td></td>
</tr>
<tr>
<td>Gulbrandsen et al (2012)63</td>
<td>Did include patients receiving insulin (n=91)</td>
<td>61/62, 56% female</td>
<td>20%; calorically restricted</td>
<td>Low fat, iso-caloric Adequate</td>
<td>0%</td>
<td>24</td>
<td></td>
</tr>
<tr>
<td>Iqbal et al (2010)64</td>
<td>BMI≥30; did include patients receiving insulin (n=144)</td>
<td>60/60, 11% female</td>
<td>&lt;30 g/d; intense behavioral support</td>
<td>Low fat Not adequate</td>
<td>47%</td>
<td>24</td>
<td></td>
</tr>
<tr>
<td>Nishimori et al (2018)65</td>
<td>NAFLD in addition; did include patients receiving insulin (n=28)</td>
<td>49/50, 36% female</td>
<td>70-130 g/d; goal was for restriction but was not seen with intake</td>
<td>Low fat; iso-caloric (per intake not per goal) Adequate</td>
<td>0%</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>Vlachos et al (2013)66</td>
<td>BMI &gt;30; did include patients receiving insulin (n=79)</td>
<td>NA</td>
<td>“Low-carbohydrate and protein sparingmodified fast”, calorically restricted</td>
<td>Low glycemic NA</td>
<td>22%</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>Westman et al (2008)67</td>
<td>BMI 27-50; did include patients receiving insulin (n=97)</td>
<td>52/52, 78% female</td>
<td>&lt;20 g/d; intense behavioral support</td>
<td>Low glycemic iso-caloric (per intake not per goal) Adequate</td>
<td>48%</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>Zadeh et al (2018)68</td>
<td>BMI 30-38; did include patients receiving insulin (n=42)</td>
<td>46.5</td>
<td>20%; intense behavioral support; calorically restricted</td>
<td>Low fat; high fat; probably iso-caloric (three control arms) NA</td>
<td>7%</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>Daly et al (2006)69</td>
<td>Obese, poorly controlled 2D; did include patients receiving insulin (n=102)</td>
<td>58/59, 52% female</td>
<td>&lt;7 g/d</td>
<td>Low fat, iso-caloric Adequate 23%</td>
<td>3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Davis et al (2009)70</td>
<td>BMI≥25; A1c ≤9.7%; did include patients receiving insulin (n=105)</td>
<td>54/53, 50% female</td>
<td>20-25 g/d + 2 weeks + 5 g/wk; calorically restricted</td>
<td>Low fat, iso-caloric Adherent at VLCD level</td>
<td>20%</td>
<td>12</td>
<td></td>
</tr>
<tr>
<td>Yancy et al (2010)71</td>
<td>BMI 27-30 plus obesity related disease, or BMI≥30; did include patients receiving insulin (n=46)</td>
<td>57/55, 13% female</td>
<td>≤20 g/d; intense behavioral support</td>
<td>Low fat + orlistat Adequate</td>
<td>11%</td>
<td>12</td>
<td></td>
</tr>
<tr>
<td>Samaha et al (2003)72</td>
<td>BMI ≥35; did include patients receiving insulin (n=52)</td>
<td>NA</td>
<td>&lt;30 g/d; intense behavioral support</td>
<td>Low fat NA</td>
<td>42%</td>
<td>12</td>
<td></td>
</tr>
<tr>
<td>Shiai et al (2008)73</td>
<td>BMI≥27 or coronary heart disease; did include patients receiving insulin (n=46)</td>
<td>NA</td>
<td>20 g/d × 2 months with gradual increase to maximum 120 g/d</td>
<td>Low fat; Mediterrane an (two control arms) NA</td>
<td>22%</td>
<td>24</td>
<td></td>
</tr>
<tr>
<td>Lee et al (2013)74</td>
<td>BMI 30-65 (n=105)</td>
<td>NA</td>
<td>&lt;70 g/d</td>
<td>Low fat NA</td>
<td>25%</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>Breukelman et al (2019)75</td>
<td>BMI 38.9 (mean) (n=39)</td>
<td>55/56, 60% female</td>
<td>&lt;50 g/d</td>
<td>No treatment control NA</td>
<td>10%</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>Perna et al (2019)76</td>
<td>BMI 24.9-34.9; A1c ≤7.8%; taking metformin; did not include patients receiving insulin (n=17)</td>
<td>59.5/67.8, 65% female</td>
<td>≤125 g/d; calorically restricted</td>
<td>“Standard” calorically restricted + metformin NA</td>
<td>0%</td>
<td>3</td>
<td></td>
</tr>
</tbody>
</table>

ADA = American Diabetes Association; BMI = body mass index; HbA1c = glycated hemoglobin; NA = not available; NAFLD = non-alcoholic fatty liver disease; T2D = type 2 diabetes; VLCD = very low calorie diet.

*Saslow 2014 met inclusion criteria but included 4/34 randomized participants who had pre-diabetes, not type 2 diabetes (>88% diabetic population). Contact with authors for diabetes specific data was unsuccessful. Although this situation was not considered a priori, research team decided to include this study on basis that any study with >80% of population having diabetes would be eligible; this scenario was not relevant in any other cases. This decision was made before results of study were reviewed.
Eighteen studies reported on weight loss at six months.\textsuperscript{34-50, 56} Pooled analysis showed that patients on LCDs achieved greater weight loss compared with control (mean difference $-3.46$, 95% confidence interval $-5.25$ to $-1.67$; $n=882$; GRADE=moderate) (table 2). On the basis of subgroup credibility testing, we found that in studies at low risk of bias, LCDs achieved 7.41 kg greater weight loss compared with controls (mean difference $-7.41$, $-9.75$ to $-5.08$; 6 studies, $n=171$; test for subgroup differences $P<0.001$) (fig 4). Seven studies reported on weight loss at 12 months,\textsuperscript{36, 39, 42-44, 50, 51} with our pooled analysis showing that any benefit over control diets was trivial and non-significant (mean difference 0.29 (–1.02 to 1.60) kg; $n=499$; GRADE=moderate) (table 2).

Seventeen studies reported on HbA\textsubscript{1c} levels at six months.\textsuperscript{34-38, 40-47, 49, 50, 52, 56} LCDs achieved greater reductions in HbA\textsubscript{1c} than did control diets (mean difference $-0.47\%$, $-0.60$ to $-0.34\%$; $n=747$; GRADE=high) (table 2). At 12 months, eight studies reported on HbA\textsubscript{1c} levels, showing that the effect size had decreased by around half (mean difference $-0.23\%$, $-0.46\%$ to $-0.00\%$; $n=489$; GRADE=moderate) (table 2).

Fourteen studies reported on fasting glucose at six months.\textsuperscript{35, 36, 38-40, 42, 44-48, 52, 53, 56} Pooled analysis showed that LCDs achieved an average 0.73 mmol/L greater reduction in glucose concentrations compared with control diets (mean difference $-0.73$, $-1.19$ to $-0.27$; $n=611$; GRADE=moderate) (table 2). Seven studies reported on fasting glucose at 12 months,\textsuperscript{39, 42, 44, 51-53} with little or no difference observed between the comparator diets (mean difference 0.06, $-0.37$ to 0.48; $n=365$; GRADE=moderate) (table 2).

Eleven studies reported total adverse events or serious adverse events at six months.\textsuperscript{34, 35, 37-39, 41-45, 47, 52} Pooled analysis suggested a trivial and non-significant increase in total adverse events among patients on LCDs (risk difference 0.04, $-0.01$ to 0.08; 9 studies, $n=423$; GRADE=very low) and similarly little or no effect on serious adverse events (risk difference 0.00, $-0.03$ to 0.02; 8 studies, $n=448$; GRADE=low) (table 2). Three studies reported on total adverse events or serious adverse events at 12 months,\textsuperscript{39, 43, 44} with pooled estimates showing that LCDs were associated with a small, non-significant decrease in total adverse events (risk difference $-0.05$, $-0.24$ to 0.14; 2 studies, $n=156$; GRADE=very low) and a trivial, non-significant decrease in serious adverse events (risk difference $-0.01$, $-0.06$ to 0.04; 3 studies, $n=217$; GRADE=low) (table 2).

Table 3 shows secondary outcomes. Briefly, pooled analyses showed that LCDs led to greater reductions in diabetes medication and clinically important benefits threefold greater than the MCID estimate for triglycerides and insulin resistance (HOMA-IR) at six and 12 months. LCDs had clinically important harms on quality of life and low density lipoprotein cholesterol at 12 months, with little to no effect observed at six months. LCDs had little or no effect on total and high density lipoprotein cholesterol concentrations or C

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>No of events/total</th>
<th>Risk difference, IV, random (95% CI)</th>
<th>Weight (%)</th>
<th>Risk difference, IV, random (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dyson 2010</td>
<td>3/6</td>
<td>1/6</td>
<td>6.5</td>
<td>0.33 (0.17 to 0.83)</td>
</tr>
<tr>
<td>Jönsson 2009</td>
<td>6/6</td>
<td>3/6</td>
<td>8.3</td>
<td>0.50 (0.08 to 0.92)</td>
</tr>
<tr>
<td>Morris 2019</td>
<td>11/21</td>
<td>0/12</td>
<td>15.0</td>
<td>0.52 (0.29 to 0.76)</td>
</tr>
<tr>
<td>Saslow 2014</td>
<td>8/9</td>
<td>5/13</td>
<td>10.8</td>
<td>0.50 (0.17 to 0.84)</td>
</tr>
<tr>
<td>Saslow 2017</td>
<td>6/11</td>
<td>0/8</td>
<td>11.4</td>
<td>0.55 (0.23 to 0.86)</td>
</tr>
<tr>
<td>Sato 2017</td>
<td>4/22</td>
<td>0/27</td>
<td>18.2</td>
<td>0.18 (0.01 to 0.35)</td>
</tr>
<tr>
<td>Tay 2014</td>
<td>36/46</td>
<td>30/47</td>
<td>17.5</td>
<td>0.14 (0.04 to 0.33)</td>
</tr>
<tr>
<td>Yamada 2014</td>
<td>2/12</td>
<td>2/12</td>
<td>12.2</td>
<td>0.00 (0.30 to 0.30)</td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td>76/133</td>
<td>41/131</td>
<td>100.0</td>
<td>0.32 (0.17 to 0.47)</td>
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Test for heterogeneity: $\chi^2=0.02$; $\chi^2=16.50$, $df=7$, $P=0.02$; $I^2=58\%$
Test for overall effect: $Z=4.15$, $P<0.001$
We did subgroup assessments (level of carbohydrate restriction, behavioral support intensity, comparator diet, iso-caloric comparator, caloric restriction, inclusion of patients who used insulin, and adherence) for each of our five primary outcomes. Most subgroup observations were not deemed credible; however, three credible subgroups were identified on the basis of meeting four of five credibility criteria. Specifically, for these subgroups, statistical analysis suggested that chance could not explain the apparent subgroup effect, the effect was consistent across studies, the subgroup hypothesis was one of a small number of hypotheses developed a priori with direction specified, and strong pre-existing biological support existed (supplementary table D). Studies that included patients using insulin had fewer remissions for both definitions of diabetes remission (HbA1c <6.5% + no diabetes medication) at six months (risk difference 0.51, 0.36 to 0.65; 0.20, 0.03 to 0.38) (test for subgroup difference P<0.001; P=0.03). Diet with very low carbohydrates (<10% of daily calories from carbohydrates) led to smaller weight loss at six months (mean difference −1.05, −2.27 to 0.17) than did less restrictive diets (mean difference −5.22, −8.33 to −2.11) (test for subgroup difference P=0.01).

Discussion
Among 23 studies comparing LCDs with mostly low fat control diets in patients with type 2 diabetes, on the basis of moderate to low certainty evidence, patients on LCDs achieved higher diabetes remission rates at six months (HbA1c <6.5%; NNT=3; HbA1c <6.5% and no diabetes medication: NNT=20). On the basis of very low to high certainty evidence, no statistically
significant and clinically important detrimental effects on cardiovascular risk factors (for example, lipids, C reactive protein) or adverse events were detected with LCDs. However, we observed a trend for clinically important increases in low density lipoprotein cholesterol at 12 months. Additionally, LCDs increased weight loss, reduced medication use, and improved triglyceride concentrations at six months. In general, most benefits diminished at 12 months, a finding consistent with previous reviews.15 57

Sensitivity and subgroup analyses
We did sensitivity analyses based on risk of bias for all outcomes, but only one outcome, weight loss, showed a credible subgroup effect between studies with higher and lower risk of bias. Studies with lower risk of bias showed more dramatic increases in weight loss, findings that were both statistically and clinically significant, supporting our overall findings.

Subgroup analyses, based on credibility testing,16 27 suggested that patients not using insulin, compared with those that did, had increased diabetes remission rates at six months. For patients not using insulin, the NNT was 2 for remission defined as HbA1c below 6.5% and 5 for remission defined as HbA1c below 6.5% without diabetes medication. Furthermore, on the basis of our subgroup testing, VLCDs underperformed compared with less restrictive LCDs for weight loss at six months. However, this difference was negated when we considered patients highly adherent to VLCDs. Of note, the limited number of studies with 12 month outcome data providing differing levels of support and having highly adherent versus less adherent intervention arms precluded subgroup analyses that explicitly explored the effects of adherence at 12 months. Although improvements noted at six months diminished by 12 months, determining with any certainty whether this is related to intensity of intervention and/or dietary adherence beyond six months is difficult.

Strengths of study
Our systematic review has several important strengths. Firstly, we did a thorough literature search and contacted authors of all studies for any unpublished
<table>
<thead>
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<th>Table 3</th>
<th>Secondary outcomes</th>
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<tr>
<td>Outcome</td>
<td>Endpoint</td>
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<tr>
<td>Quality of life—diabetes specific overall score (DDS and PAID)</td>
<td>6 months</td>
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<td>12 months</td>
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<tr>
<td>Quality of life—diabetes specific overall score (PAID, converted)</td>
<td>6 months</td>
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<td>12 months</td>
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<tr>
<td>Medication reduction (No of participants who reduced diabetes medication)</td>
<td>6 months</td>
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<td>12 months</td>
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<td>Total cholesterol (mmol/L)</td>
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<td>LDL cholesterol (mmol/L)</td>
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<td>Triglycerides (mmol/L)</td>
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<tr>
<td>Insulin resistance (HOMA-IR)</td>
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<td>12 months</td>
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<tr>
<td>Inflammation (C reactive protein, mg/L)</td>
<td>6 months</td>
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<td>12 months</td>
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DDS=diabetes distress scale; HDL=high density lipoprotein; HOMA-IR=homeostasis model assessment of insulin resistance; LDL=low density lipoprotein; MCID=minimal clinically important difference; MD=mean difference; NA=not available; PAID=problem areas in diabetes; RD=risk difference; SMD=standardized mean difference.

data on remission of diabetes. Although only three included studies previously published HbA1c threshold criteria and medication use to determine diabetes remission, our successful contact with authors yielded trial data from five additional studies to determine remission rates. Increasing the precision and overall certainty of the effect estimates, recent systematic reviews conducted by Sainsbury, van Zuuren, and Snorgaard have shown important reductions in mean HbA1c values with low and very low carbohydrate diets, but no previous review has summarized HbA1c as a dichotomous outcome informed by the suggested American Diabetes Association remission definitions (for example, <6.5% HbA1c threshold). We believe that our meta-analytic summary of published and unpublished data from eight randomized controlled trials using HbA1c thresholds, a first in the literature, will lead to more informed clinical decision making in the management of type 2 diabetes.

Secondly, on the basis of a publicly available protocol, we used robust evidence synthesis methods including the use of Cochrane’s Risk of Bias instrument and subgroup credibility assessments based on a priori stated effect modifiers. Missing data for participants is particularly important in nutrition research in general given the often dramatic losses to follow-up in diet based clinical trials (>20% among 10/23 (43%) of trials included in this analysis) and the corresponding risk of bias due to losses to follow-up. Subgroup credibility assessment is of particular interest to researchers in this field given that some have advocated for subgroup elucidation when considering LCDs for treating diabetes. Whereas previous reviews have focused on one or two potential modifiers—for example, Korsmo et al, who explored subgroups on length of follow-up and carbohydrate intake, and Naude et al, who explored calorically matched controls—in our protocol driven approach, we explored seven actively debated potential effect modifiers by using published, explicit subgroup credibility criteria.

Thirdly, the use of GRADE for rating the certainty of evidence in systematic reviews of nutrition studies has been questioned, with some calling for a methodological approach specific to nutrition studies. However, we believe the logic of scientific inquiry demands consistent standards for casual inference across health claims, preferably using GRADE, a more conservative rating approach than the alternative systems suggested by the nutrition community. Nevertheless, we did a sensitivity analysis comparing GRADE ratings with NutriGRADE ratings (supplementary table E). NutriGRADE analysis resulted in 16/30 (53%) outcomes with the same rating as GRADE; 10 (33%) of outcomes were judged to be of higher certainty using NutriGRADE, and 4 (13%) were judged to be of lower certainty using NutriGRADE. Overall, the certainty of evidence using NutriGRADE indicates, on average, a higher degree of confidence in the efficacy and safety of LCDs across outcomes, particularly our primary outcomes including diabetes remission and fasting glucose, and higher certainty in the evidence for little to no short term risk of adverse events with LCDs.

Fourthly, our interpretations of estimates for continuous outcomes were based on a priori estimates of the minimal clinically important differences (supplementary table C). To our knowledge, no previous review on this topic has attempted to present effect estimates while considering MCID thresholds, thresholds that will help clinicians and patients to better interpret the magnitude of treatment effect. Among 10 continuous outcomes, two showed improvements that met or surpassed the MCID at six months (triglycerides, insulin resistance) with no detrimental effects. At 12 months, two had improvements that surpassed the
MCID (triglycerides, insulin resistance) and two had a clinically important worsening (quality of life, low density lipoprotein cholesterol), although neither was statistically significant (P=0.24 and P=0.05).

Limitations of study
Our study is not without limitations. Firstly, the definition of remission of diabetes is the subject of considerable debate, specifically with regards to threshold levels of HbA1c/fasting glucose, use of diabetes medication, and the length of follow-up time meeting these criteria. We attempted to overcome this by using multiple a priori definitions of remission (both with and without the use of diabetes medication) at both of our predetermined endpoints (six months and 12 months).

Secondly, safety concerns have been raised with LCDs. Although no significant or clinically important increase in total or serious adverse events was identified, these outcomes were poorly reported among trials and the certainty of evidence for safety ranges from low to very low. By contrast, we have moderate to high certainty that surrogate markers for cardiovascular disease risk, such as blood lipids, do not worsen, whereas triglycerides significantly improved in a clinically meaningful way. One exception was low density lipoprotein cholesterol concentrations at 12 months’ follow-up, which seemed to worsen, surpassing the MCID. Thirdly, 18/23 (78%) studies used low fat diets as a comparator, limiting the applicability of our results to other dietary regimens such as a Mediterranean-style diet.

Fourthly, an important concern with LCDs is the potential confounding factor of caloric restriction. Restricting carbohydrates, which tends to reduce hunger, would mean that whether any purported benefit was due to carbohydrate restriction or caloric restriction was unclear. For this reason, as part of our a priori planned subgroup analysis, we investigated the effect of calorically matched controls (as assessed by follow-up dietary questionnaires). On the basis of 18 studies providing adequate data, we identified no evidence of credible effect modification based on caloric matching or lack thereof. However, self-reported dietary intake data are prone to measurement error, particularly in dietary trials in which participants are not blinded.

Fifthly, we made a pragmatic a priori decision to assess our endpoints at six and 12 months (±3 months). Whereas trials informing our 12 month endpoint were all reported at this time point, those informing our six month endpoint varied between three months and eight months. Of the 14 trials informing our six month pooled estimates, 7/14 (50%) reported data at three to less than six months (3 months: 6 trials; 4 months: 1 trial), and 7/14 (50%) trials reported at six to nine months (6 months: 6 trials; 8 months: 1 trial). On the basis of comments from peer reviewers, we did a post hoc analysis on remission at six (±3) months. Evidence suggested larger treatment effects for LCDs in shorter term trials (3 to <6 months), suggesting that shorter term trials may be an effect modifier. For the definition of remission of HbA1c below 6.5%, the risk difference was 0.49 (95% confidence interval 0.30 to 0.68) for trials of three to less than six months in length compared with 0.25 (0.08 to 0.42) for trials of between six and nine months. Similarly, for the definition of remission of HbA1c below 6.5% and no diabetes medication use, the risk difference was 0.20 (0.03 to 0.38) for trials of three to less than six months compared with 0.00 (–0.07 to 0.07) for trials of between six and nine months.

Sixthly, our protocol driven results are limited to short term markers of remission of diabetes, adverse events, and related cardiometabolic outcomes. Future long term, well designed, calorie controlled randomized trials are needed to determine the effects of LCD on sustained weight loss and remission of diabetes, as well as cardiovascular mortality and major morbidity.

Seventhly, our review focused on studies defined by macronutrient quantity. Macronutrient quality may also be important, and, although we were unable to consider the characteristics of dietary quality given the lack of reporting in our 23 eligible trials, future trials should better document dietary quality (for example, processed versus unprocessed foods) using optimally validated questionnaires together with emerging objective biomarkers using microbiomics, metabolomics, or other high dimensional platforms.

Finally, the limited number of trials allowing patients to reduce their medication use impeded our ability to assess remission of diabetes when defined as HbA1c below 6.5% without diabetes medication. Only 7/23 (30%) of eligible trials permitted reduction of medication and reported usable medication data. Future trials should allow for, and adequately report on, reduction of medication while closely monitoring blood glucose concentrations. LCDs seem to promote important reductions in HbA1c, potentially increasing risk for hypoglycemic episodes, including severe syncope, if the dosage of diabetes medications is not adjusted accordingly. Because blinding is not possible in these studies, these adjustments should be applied using a priori algorithms that help to guide medication management. Reductions in medication may blunt the effect on mean HbA1c levels, biasing results towards the null and masking any effect; however, any improvement can still be captured if reduction of medication is included as an outcome of interest.

Conclusions
Moderate to low certainty evidence suggests that patients adhering to LCDs for six months may experience greater rates of remission of diabetes without adverse consequences compared with other diets commonly recommended for management of type 2 diabetes (for example, low fat diets). These benefits diminished at 12 months, and, although LCDs seem to improve triglycerides in a clinically meaningful way, some evidence shows clinical worsening of quality of life and low density lipoprotein cholesterol. Considering this and a recent systematic review of cohort studies suggesting...
that long term LCDs are associated with increased mortality, clinicians might consider short term LCDs for management of type 2 diabetes, while actively monitoring and adjusting diabetes medication as needed.

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**Contributors:** JZG and BCJ conceived the study. JZG, LT, and BCJ designed the study. JZG, JJ, and BCJ developed a priori estimates of the minimal clinically important difference. JB designed and executed the search. JJ and AD selected the articles and extracted the data. JZG, AD, and BCJ analyzed the data. JZG and BCJ wrote the first draft of the manuscript. GB, JS, SY, and Tj provided unpublished trial data and reviewed and interpreted the data of the draft manuscript. JZG, BCJ, AD, JB, LT, GB, JS, SY, Tj, and JJ interpreted the data and contributed to the writing of the final version of the manuscript. All authors agreed with the results and conclusions of this article. The corresponding author attests that all listed authors meet authorship criteria and that no others meeting the criteria have been omitted. JZG and BCJ are the guarantors.

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**Competing interests:** All authors have completed the ICJME uniform disclosure form at www.icjme.org/cgi disclose.pdf and declare: support from Texas A&M University; BCJ receives funds from Texas A&M AgriLife Research to support investigator initiated research related to saturated and polyunsaturated fats for a separate research project, as part of his recent recruitment to Texas A&M University; GB is author of the CSIRO Low Carb Diet Book that aims to promote a low carbohydrate dietary lifestyle and publicize the benefits; support from Texas A&M AgriLife institutional funds are from interest and investment earnings, not a sponsoring organization, industry, or company; GB does not publish any information regarding his status or financial support from the CSIRO. The other authors report no conflicts of interest.

**Ethical approval:** All the work was developed using aggregate level data.

**Data sharing:** No needed. All the work was developed using aggregate level data.

**Data sharing:** Future data are available on request through the corresponding author at bradley.johnston@jutalung.org.

The lead and senior authors (manuscript’s guarantors) affirm that this manuscript is an honest, accurate, and transparent account of the study being reported; that no important aspects of the study have been omitted; and that any discrepancies from the study as planned (and, if relevant, registered) have been explained.

**Dissemination to participants and related patient and public communities:** We plan to reach out to diabetes and obesity patient advocacy groups (eg, Obesity Canada) as well as professional medical, nutrition, and agricultural organizations (eg, Practice-based Evidence in Nutrition, Royal Australian College of General Practitioners, USDA) to help to disseminate this work. Additionally, all authors will work with their home institutions to leverage their unique dissemination platforms including social media communication and organizational websites.

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Web appendix: Supplementary tables