Effect of fructose on markers of non-alcoholic fatty liver disease (NAFLD): a systematic review and meta-analysis of controlled feeding trials

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BACKGROUND/OBJECTIVES: In the absence of consistent clinical evidence, there are concerns that fructose contributes to non-alcoholic fatty liver disease (NAFLD). To determine the effect of fructose on markers of NAFLD, we conducted a systematic review and meta-analysis of controlled feeding trials.

SUBJECTS/METHODS: We searched MEDLINE, EMBASE, CINAHL and the Cochrane Library (through 3 September 2013). We included relevant trials that involved a follow-up of ≥ 7 days. Two reviewers independently extracted relevant data. Data were pooled by the generic inverse variance method using random effects models and expressed as standardized mean difference (SMD) for intrahepatocellular lipids (IHCL) and mean difference (MD) for alanine aminotransferase (ALT). Inter-study heterogeneity was assessed (Cochran Q statistic) and quantified (I² statistic).

RESULTS: Eligibility criteria were met by eight reports containing 13 trials in 260 healthy participants: seven isocaloric trials, in which fructose was exchanged isocalorically for other carbohydrates, and six hypercaloric trials, in which the diet was supplemented with excess energy (+ 21–35% energy) from high-dose fructose (+ 104–220 g/day). Although there was no effect of fructose in isocaloric trials, fructose in hypercaloric trials increased both IHCL (SMD = 0.45 (95% confidence interval (CI): 0.18, 0.72)) and ALT (MD = 4.94 U/l (95% CI: 0.03, 9.85)).

LIMITATIONS: Few trials were available for inclusion, most of which were small, short (< 4 weeks), and of poor quality.

CONCLUSIONS: Isocaloric exchange of fructose for other carbohydrates does not induce NAFLD changes in healthy participants. Fructose providing excess energy at extreme doses, however, does raise IHCL and ALT, an effect that may be more attributable to excess energy than fructose. Larger, longer and higher-quality trials of the effect of fructose on histopathological NAFLD changes are required.

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INTRODUCTION

Non-alcoholic fatty liver disease (NAFLD) is the most prevalent chronic liver disease and a cause of raised liver enzymes in developed countries, affecting 10–30% of people in developed countries.1 The increasing prevalence of NAFLD, which is closely linked with the increasing prevalence of obesity and type 2 diabetes mellitus (T2DM),2 has been associated with increased cardiovascular morbidity and mortality.1

Dietary factors that influence NAFLD have become a focus of attention. In particular, recent concerns have been raised regarding the role of dietary fructose in inducing NAFLD.4–7 Animal models featuring extreme levels of fructose exposure8–11...
and poor-quality observational studies have been used to underpin this hypothesis. In the absence of consistent clinical evidence, it is unclear whether fructose at typical levels of exposure induces NAFLD. To determine the effect of fructose on markers of NAFLD in humans, we conducted a systematic review and meta-analysis of available controlled feeding trials.

**SUBJECTS AND METHODS**

**Design**

We followed the Cochrane Handbook for Systematic Reviews of Interventions and the Preferred Reporting Items for Systematic Reviews and Meta-Analysis guidelines. The review protocol is available at ClinicalTrials.gov (registration number: NCT01363791).

**Study selection**

We searched MEDLINE, EMBASE, CINAHL and the Cochrane Library databases through 3 September 2013 for relevant articles. Supplementary Appendix Table 1 shows the full search term used in this study. Manual searches supplemented the electronic search strategy. No restrictions were placed on language. We included controlled trials investigating the effect of oral fructose on markers of NAFLD. A control arm was considered isocaloric when the carbohydrate comparator was exchanged for an equal amount of fructose. If the trial involved overfeeding of fructose so that the fructose provided excess energy resulting in a positive energy balance, then the comparison was still considered isocaloric as long as the carbohydrate comparator was matched for the excess energy resulting in the same positive energy balance. A comparison was considered hypercaloric when a control diet was supplemented with excess energy from fructose compared with the same control diet alone without the excess energy. Trials that involved a follow-up of < 7 days follow-up, administered intravenous fructose, lacked a control diet or did not provide suitable endpoint data were excluded.

**Data extraction**

Two reviewers (SC, AIC) independently reviewed and extracted relevant data from each report. The quality of each study was assessed using the Heyland methodological quality score (MQS). Disagreements were reconciled by consensus. Mean ± s.d. differences between fructose and control arms were extracted as the main end points. In those trials where the data were included in figures and not provided numerically, we used software program Plot Digitizer (http://plotdigitizer.sourceforge.net/) to extract the data. Additional information was requested from the authors of all included trials.

**Access to study**

All authors had access to the study data and reviewed and approved the final manuscript.

**Statistical analysis**

Data analyses were conducted using Review Manager (RevMan) version 5.1.6 (Copenhagen, Denmark) for primary analyses and Stata (version 12, College Station, TX, USA) for subgroup analyses. Separate analyses were conducted for the isocaloric and hypercaloric trials using the generic inverse variance method with random effects weighting. Data were expressed as standardized mean differences with 95% confidence intervals (CIs) for intrahepatic lipid (IHCL) and mean differences (MD) with 95% CIs for alanine aminotransferase (ALT).

Trials that did not report SE values had these computed from the available statistics using standard formulae. To generate SE for included crossover trials, we assumed a paired analyses as described by Elbourne. If insufficient data were available for computations in crossover trials, SE values were imputed using the pooled correlation coefficient between baseline and end-of-study values derived from a meta-analysis of trials reporting sufficient data or assuming a conservative correlation coefficient of 0.5 with sensitivity analyses at 0.25 and 0.75.

Inter-trial heterogeneity was assessed by the Cochran Q statistic with $\chi^2 < 0.10$ considered significant, and quantified by the $I^2$ statistic, where $I^2 > 50\%$ indicates substantial heterogeneity. Sources of heterogeneity were investigated by sensitivity analyses in which each individual study was removed from the analysis and through a priori subgroup analyses by comparator, baseline values, fructose form, follow-up, MQS, randomization, design and energy balance. Meta-regression analyses assessed the significance of subgroup effects. Publication bias was evaluated via visual inspection of funnel plots and Egger and Begg tests.

**RESULTS**

Search results

Figure 1 shows the trial selection process. We identified 1437 eligible reports. A total of eight reports (providing data for 13 trials) were selected for analyses.

**Trial characteristics**

Table 1 shows the trial characteristics. Only two markers of NAFLD were identified: IHCL and ALT. None of the available trials assessed NAFLD histologically from liver biopsies. There were a total of seven isocaloric trials (four for IHCL and six for ALT) in 184 healthy participants, and six hypercaloric trials (five for IHCL and four for ALT) in 76 participants ($n = 60$ healthy and $n = 16$ offspring of type 2 diabetes). The majority of both sets of trials were conducted in European countries in an outpatient setting and tended to be small (median (interquartile range) sample size, 29.0 (24.5–31.5) and 13.5 (10.5–15.8), in isocaloric and hypercaloric trials, respectively).

Participants tended to be healthy, young (median (IQR) age = 30.5 years (26.3–33.9 years) and 27.6 years (24.7–33.9 years)), male (median (IQR) percent male/female ratio = 100% (56–100) and 77.5–100), and overweight (median (IQR) body mass index = 25.9 kg/m² (22.4–29.4 kg/m²) and 24.3 kg/m² (22.2–28.5 kg/m²)) in isocaloric and hypercaloric trials, respectively. Median (IQR) baseline ALAT values (in U/l) were 26.0 (23–28.9) in isocaloric trials and 20.58 (18.5–25.8) in hypercaloric trials. Median values for baseline IHCL (in %) could not be computed from the data reported.

Crossover designs were used in 29% of isocaloric trials and all hypercaloric trials. The majority of isocaloric trials (86%) and 17% of hypercaloric trials were randomized. Glucose was the comparator in all isocaloric trials except in the two trials of Aeberli et al. where glucose and sucrose were the comparators in the high dose trial and glucose and starch were the comparators in the low dose trial. The control diet alone without the added energy from fructose was the comparator in all hypercaloric trials. Although comparisons in all isocaloric trials were matched for energy, 86% of the isocaloric trials provided fructose and the carbohydrate comparator under conditions of positive energy balance (that is, both arms provided excess energy), whereas 14% provided fructose and the carbohydrate comparator under conditions of neutral energy balance (that is, both arms provided energy to maintain weight). All isocaloric and hypercaloric trials administered fructose in fluid form at a median (IQR) dose of 182 g/day (115–204 g/day) for isocaloric trials and +193 g/day (+158–211 g/day) for hypercaloric trials. The median (IQR) excess energy provided by the hypercaloric trials was +25% (+23–33%). All isocaloric and hypercaloric trials featured high-carbohydrate and low-fat diets with similar macronutrient profiles: 50–55% energy carbohydrate, 30–35% energy fat and 13–15% energy protein. Metabolic feeding control was used in 14% of isocaloric trials and 17% of hypercaloric trials; partial-metabolic feeding control was used in 43 and 17% and the remainder provided fructose as a supplement. The median (IQR) dietary follow-up was 4 weeks (3–8 weeks) for isocaloric trials and 3 weeks (1.25–4 weeks) for hypercaloric trials.

The Heyland MQS was considered high (MQS ≥ 8) in 71% of isocaloric and 33% of hypercaloric trials. Lack of or poor description of randomization, nonconsecutive or poorly described patient selection and absence of double-blinding contributed to lower scores. Funding of all trials was from a combination of agency alone (69%) or agency–industry sources (31%). None were
funded by industry alone. None reported a potential conflict of interest.

Effect of fructose on IHCL

Figure 2a shows the effect of fructose on IHCL in isocaloric trials. Primary pooled analyses showed no effect of fructose on IHCL (standardized mean difference $\mu = -0.09$ (95% CI: $-0.36$ to $0.18$), $P = 0.51$), with no evidence of inter-study heterogeneity ($I^2 = 0\%$, $P = 0.95$). Sensitivity analyses did not alter the results. Meta-regression analyses showed no statistically significant subgroup effects (Supplementary Appendix Figure 1).

Figure 2b shows the effect of fructose on IHCL in hypercaloric trials. Primary pooled analyses showed that fructose raised IHCL (standardized mean difference $\mu = 0.45$ (95% CI: $0.18$ to $0.72$), $P = 0.001$), though there was significant inter-study heterogeneity ($I^2 = 55\%$, $P = 0.07$). Sensitivity analyses did not alter the results, but identified that the removal of Lê et al.,$^{23}$ a study conducted in offspring of individuals with T2DM, eliminated evidence of inter-study heterogeneity ($I^2 = 0\%$, $P = 0.87$). Meta-regression analyses showed no statistically significant subgroup effects (Supplementary Appendix Figure 2), and inter-study heterogeneity remained largely unexplained.

Effect of fructose on liver enzymes

Figure 3a shows the effect of fructose on ALT in isocaloric trials. To approximate paired analyses for crossover trials, we used a conservative correlation coefficient of 0.5. Primary pooled analyses showed no effect of fructose on ALT (MD = $0.15$ (95% CI: $-1.51$ to $1.82$), $P = 0.86$), with no significant evidence of inter-study heterogeneity ($I^2 = 0\%$, $P = 0.97$). Neither sensitivity analyses nor the use of more (0.75) or less (0.25) conservative correlation coefficients altered the results. Sensitivity analyses nor the use of more (0.75) or less (0.25) conservative correlation coefficients altered the results. Meta-regression analyses revealed no statistically significant subgroup effects (Supplementary Appendix Figure 3).

Figure 3b shows the effect of fructose on ALT in hypercaloric trials. To approximate paired analyses for crossover trials, we needed to use a conservative correlation coefficient of 0.5. Primary analyses showed a significant ALT-increasing effect (MD = $4.94$ (95% CI: $0.03$ to $9.85$), $P = 0.05$), with significant evidence of inter-study heterogeneity ($I^2 = 78\%$, $P = 0.003$). Sensitivity analyses revealed the removal of either Lê et al.,$^{23}$ Sobrecases et al.$^{24}$ or Johnston et al.$^{26}$ led to a loss of significance (MD = $2.97$ (95% CI: $1.40$, $7.35$); MD = $5.40$ (95% CI: $-1.97$, $12.78$); and MD = $4.84$ (95% CI: $-1.51$, $11.19$), respectively. Removal of Cox et al.,$^{25}$ significantly reduced evidence of inter-study heterogeneity ($I^2 = 56\%$, $P = 0.11$). Sensitivity analysis using higher (0.75) or lower (0.25) conservative correlation coefficients did not alter the results. Meta-regression analyses did not show any statistically significant subgroup effects (Supplementary Appendix Figure 4).

Publication bias

We examined funnel plots for evidence of publication bias (Supplementary Appendix Figure 5). There was some evidence of slight asymmetry in the hypercaloric trials for ALT on visual

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Table 1. Characteristics of controlled feeding trials investigating the effect of fructose on markers of NAFLD

| Study | Subjects | Age (years) | BMI (kg/m²) | Setting | Baseline IHCL* | Baseline ALT (U/l) | Design | Feeding control | Rando-mization | Fructose doseb | Fructose form | Comparatora | Dietb | Energy balance | Follow-up | MQSc | Funding sourceb |
|-------|----------|-------------|-------------|---------|---------------|-------------------|--------|----------------|---------------|---------------|--------------|-------------|------------|-------|----------------|-----------|-------|----------------|
| **Isocaloric trials** | | | | | | | | | | | | | | | | | | |
| Johnston et al.20 | 32 OW/OB (32M/0F) | 33.9 ± 10.0 | 29.4 ± 1.6 | OP, UK | 7.61 ± 5.3% | 28.9 ± 1.26 | P | Met | sup | No | n/a | 204-g/d | Sub | Neutral | 8 wk | 10 | Agency |
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| Cox et al.25 | 31 OW/OB (16M/15F) | 53.7 ± 8.1 | 29.3 ± 2.85 | IP/OP, USA | — | — | P | Met | sup | No | n/a | 204-g/d | Sub | Neutral | 8 wk | 10 | Agency |
| Aebeler et al.21 (HD) | 29 N (29M/0F) | 26.3 ± 6.6 | 22.4 ± 1.9 | OP, Switzerland | — | 23 ± 1.7 | C | Supp | No | n/a | 204-g/d | Sub | Neutral | 8 wk | 10 | Agency |
| Aebeler et al.21 (LD) | 29 N (29M/0F) | 26.3 ± 6.6 | 22.4 ± 1.9 | OP, Switzerland | — | 23 ± 1.7 | C | Supp | No | n/a | 204-g/d | Sub | Neutral | 8 wk | 10 | Agency |
| Silbernagel et al.20 | 20 N (12M/8F) | 30.5 ± 8.94 | 25.9 ± 2.24 | OP, Germany | 1.45 ± 0.85 % | Signal | — | P | Supp | No | n/a | 204-g/d | Sub | Neutral | 8 wk | 10 | Agency |
| Ngo Sock et al.19 | 11 N (11M/0F) | 24.6 ± 1.99 | 22 (19–25) | OP, Switzerland | 2.42 ± 0.83 log mmol/kg | 26 ± 13.3 | P | Met | sup | No | n/a | 204-g/d | Sub | Neutral | 8 wk | 10 | Agency |
| **Hypercaloric trials** | | | | | | | | | | | | | | | | | | |
| Johnston et al.20 | 15 OW/OB (15M/0F) | 35.0 ± 11.0 | 30.0 ± 1.4 | OP, UK | 7.20 ± 5.6% | 31.0 ± 1.50 | C | Supp | No | n/a | 203-g/d | Sub | Neutral | 8 wk | 10 | Agency |
| Cox et al.25 | 16 OW/OB (9M/7F) | 52.5 ± 9.3 | 29.3 ± 2.6 | IP/OP, USA | — | — | P | Met | sup | No | n/a | 204-g/d | Sub | Neutral | 8 wk | 10 | Agency |
| Silbernagel et al.20 | 10 N (7M/3F) | 30.5 ± 6.32 | 25.9 ± 1.58 | OP, Germany | 1.32 ± 0.92 % | Signal | — | C | Supp | No | n/a | 204-g/d | Sub | Neutral | 8 wk | 10 | Agency |
| Sobrecasses et al.24 | 12 N (12M/0F) | 23.9 ± 2.2 | 22.6 ± 1.1 | OP, Germany | 12.83 ± 2.38 | mmol/kg | 20.58 ± 8.7 | C | Supp | No | n/a | 204-g/d | Sub | Neutral | 8 wk | 10 | Agency |
| Lè et al.22 | 7 N (7M/0F) | 24.70 ± 3.44 | 22 (19–25) | OP, Switzerland | 6.21 ± 2.09 | mmol/kg | 16.4 ± 4 | C | Met | sup | No | n/a | 204-g/d | Sub | Neutral | 8 wk | 10 | Agency |
| Lè et al.23 | 16 OFF-DMM (16M/0F) | 24.7 ± 5.2 | 22 (19–25) | OP, Switzerland | 6.21 ± 2.09 | mmol/kg | 16.4 ± 4 | C | Met | sup | No | n/a | 204-g/d | Sub | Neutral | 8 wk | 10 | Agency |

Abbreviations: BMI, body mass index; C, crossover; E, energy; F, female; HD, high dose; IHCL, intrahepatocellular lipids; IP, inpatient; LD, low dose; M, male; Met, metabolic; MQS, methodological quality score; n, normal; NAFLD, non-alcoholic fatty liver disease; Off-T2DM, offspring of persons with type 2 diabetes mellitus; OP, outpatient; P, parallel; Supp, supplement. *IHCL was measured by 1H-magnetic resonance spectroscopy in each trial. **Doses preceded by a " - " represent average doses calculated on the basis of the average reported energy intake or weight of participants. If these data were not available, then the average dose was based on an 2000-kcal intake. Plus signs indicate excess energy provided by fructose. †Fructose was provided as beverages or crystalline fructose to be added to beverages. ‡Comparators were the reference carbohydrate in the isocaloric trials and the control diet (weight-maintaining, background diet) alone without the added energy from fructose in the hypercaloric trials. Fructose was exchanged for the reference carbohydrate, providing an energy-matched comparison in the isocaloric trials, while it supplemented the control diet to provide excess energy in the hypercaloric trials. §Energy from carbohydrate:fat:protein. ‡Trials with a score > 8 were considered to be of higher quality according to the Heyland MQS. 17 †Agency funding is that from government, university or not-for-profit health agency sources. 18 The high dose of Aebeler et al.,21 was obtained by averaging the high fructose vs high sucrose and high fructose vs high starch comparisons, and the low dose was obtained from averaging the medium fructose vs medium glucose and the medium fructose vs starch (low fructose) comparisons.
inspection ($P = 0.056$ by Egger test; $P = 0.089$ by Begg test), but no small study effects were detected among the isocaloric and hypercaloric trials for either IHCL or ALT by Egger and Begg tests ($P > 0.05$).

**DISCUSSION**

The present aggregate analyses of 13 trials in 260 predominantly young, male participants, who were overweight/obese or otherwise healthy, investigated the effect of fructose on markers of

**a Isocaloric Trials**

| Trial                      | Year | Participants | % Weight | Standardized Mean Differences (95% CI) in IHCL |
|----------------------------|------|--------------|----------|-----------------------------------------------|
| Johnston et al.26 (E Neutral) | 2013 | 32           | 11.0%    | -1.10 [-6.11, 3.91]                           |
| Johnston et al.26 (E Positive) | 2013 | 32           | 13.5%    | 1.70 [-2.84, 6.24]                            |
| Cox et al.25                 | 2012 | 31           | 30.5%    | 0.00 [-3.02, 3.02]                            |
| Aeberli et al.21 (LD)        | 2011 | 29           | 20.2%    | 0.50 [-3.21, 4.21]                            |
| Aeberli et al.21 (HD)        | 2011 | 29           | 19.1%    | 0.00 [-3.81, 3.81]                            |
| Ngo Sock et al.19            | 2010 | 11           | 5.8%     | -1.00 [-7.93, 5.93]                           |
| **Total**                   |      | **164**      | **100.0%**| **0.15 [-1.51, 1.82]**                        |

Heterogeneity: $\text{ Tau}^2 = 0.00; \text{ Chi}^2 = 0.84, df = 5 (P = 0.97); I^2 = 0%$

Test for overall effect: $Z = 0.18 (P = 0.86)$

**b Hypercaloric Trials**

| Trial                      | Year | Participants | % Weight | Standardized Mean Differences (95% CI) in IHCL |
|----------------------------|------|--------------|----------|-----------------------------------------------|
| Johnston et al.26          | 2013 | 15           | 26.1%    | 5.80 [1.40, 10.20]                            |
| Cox et al.25               | 2012 | 16           | 28.7%    | -0.80 [-4.15, 2.55]                           |
| Sobrecases et al.24        | 2010 | 12           | 25.4%    | 4.66 [0.36, 8.96]                             |
| Lê et al.23 (Off-T2DM)     | 2009 | 16           | 18.8%    | 12.90 [5.45, 20.35]                           |
| **Total**                  |      | **59**       | **100.0%**| **4.94 [0.03, 9.85]**                         |

Heterogeneity: $\text{ Tau}^2 = 18.96; \text{ Chi}^2 = 13.76, df = 3 (P = 0.000); I^2 = 78%$

Test for overall effect: $Z = 1.97 (P = 0.05)$

**Figure 2.** Forest plots of the effect of fructose on intrahepatocellular lipid (IHCL) in healthy participants in (a) isocaloric and (b) hypercaloric feeding trials. Pooled effect estimates shown as diamonds. Data are expressed as weighted MD with 95% CI using generic inverse variance random effects models. Inter-study heterogeneity was tested by Cochrane's $Q$ statistic ($\chi^2$-test) at a significance level of $P < 0.10$ and quantified by $I^2$, where $I^2 > 50\%$ is considered to be evidence of substantial heterogeneity and $> 75\%$, considerable heterogeneity. Any CHO denotes any carbohydrate comparator; E neutral, neutral energy balance; E positive, positive energy balance; and Off-T2DM, offspring of T2DM.

**Figure 3.** Forest plots of the effect of fructose on ALT in healthy participants in (a) isocaloric and (b) hypercaloric feeding trials. Pooled effect estimates shown as diamonds. Data are expressed as weighted MD with 95% CI using generic inverse variance random effects models. Inter-study heterogeneity was tested by Cochrane's $Q$ statistic ($\chi^2$-test) at a significance level of $P < 0.10$ and quantified by $I^2$, where $I^2 > 50\%$ is considered to be evidence of substantial heterogeneity and $> 75\%$, considerable heterogeneity. Any CHO denotes any carbohydrate comparator; E neutral, neutral energy balance; E positive, positive energy balance; HD, high dose; LD, low dose; and Off-T2DM, offspring of T2DM.
NAFLD under two different types of trial conditions: one where fructose in beverage form was isocalorically exchanged for other carbohydrates and the other where fructose in beverage form supplemented control diets with excess energy (± 21–35% energy) at extreme doses (104–220 g/day) relative to the same control diets without the excess energy. These two types of trial conditions produced different results. Although there was no effect of fructose in isocaloric trials, fructose increased both IHCL and ALT in hypercaloric trials.

Relation of findings to other lines of evidence
Our finding of a lack of effect of fructose on NAFLD markers in isocaloric trials contradicts evidence from animal models and observational studies. The ability of fructose to induce a metabolic syndrome phenotype and NAFLD is thought to lie in its ability to act as an unregulated substrate for de novo lipogenesis, bypassing the major rate-limiting step of glycolysis at phosphofructokinase.7,27 This mechanism contributes significantly to de novo lipogenesis in rodent models, in which fructose fed at supraphysiological doses under isocaloric (~60% energy) or hypercaloric (~30% excess energy) conditions induces steatosis and steatohepatitis.8-11 Small cross-sectional and retrospective case–control studies have also shown an association between fructose-consuming sugar intake and NAFLD.12-14 Clinical translation of these data, however, has several limitations. Rodent models are complicated by supraphysiological doses and excess energy,28 and marked differences exist in the metabolic fate of fructose between animals and humans. Although de novo lipogenesis from fructose accounts for 60–70% of fatty acids in rodents,28 its contribution in humans is quantitatively insignificant.29,30 Two carefully conducted reviews of the available isocaloric tracer studies showed that de novo lipogenesis from fructose contributes <1% of fatty acids, whereas glucose (~50%), lactate (~25%) and glycogenosis (~15%) synthesis remain the major pathways of hepatic fructose disposal in humans.29,30 Cross-sectional and retrospective case–control studies do not provide evidence of causation and have found positive associations with many other factors that might be equal or better predictors of NAFLD, such as increased intake of energy, total fat, total carbohydrate, animal protein, cholesterol and the n-6:n-3 ratio of polyunsaturated fatty acids and decreased intake of dietary fiber.6 No large prospective observational studies have evaluated the relationship between fructose and NAFLD.

Energy represents an important confounding factor in the effect of fructose. Overfeeding of a ‘fast food’ diet has been shown to relate to an increase in ALT in healthy participants.31 Randomized trials of energy-restricted diets focusing on total energy reduction and exercise to promote weight loss have also shown reversal of NAFLD markers in people with NAFLD.32 In the present analyses, we observed increases in IHCL and ALT only in hypercaloric trials. The lack of effect in the isocaloric trials was seen even under conditions of positive energy balance. Six of the isocaloric trials (three of four trials assessing IHCL19,20,26 and five of six trials assessing ALT19,25,26) used excess energy diets in both the fructose and comparator arms, so permitting the effect of fructose to be isolated from that of energy under matched, yet excess energy-feeding conditions. Restricting our analyses to these trials did not show an effect of fructose on NAFLD markers. We made similar observations for the lack of effect of fructose on both body weight33 and uric acid34 in two earlier systematic reviews and meta-analyses. These data suggest that the effect of fructose on NAFLD markers may not be different from that of other carbohydrates as long as energy remains matched.

Previous meta-analyses have identified subgroup effects on related metabolic end points. A dose threshold was observed for a triglyceride-raising effect of fructose: ≥100 g/day for fasting and ≥50 g/day for postprandial triglycerides across different participant groups35 and ≥60 g/day for fasting triglycerides in type 2 diabetes.36 A fasting triglyceride-raising effect of fructose was also seen where starch was the comparator and follow-up was ≤4 weeks in type 2 diabetes,36 whereas a weight-loss effect of fructose was seen in overweight/obese individuals and where fructose was in fruit form.37 None of these subgroup analyses were significant in the present analysis. Although the number of trials was small, the lack of effect modification across a priori subgroup analyses was consistent with that seen in our earlier meta-analyses for blood pressure37 and uric acid.34

Implications
Although our results bear on the question of whether fructose-containing sugar-sweetened beverages have a unique role in the development of NAFLD, their translation to ‘real-world’ intake patterns is complicated. The median level of fructose exposure was >95th percentile U.S. intake (87 g/day) across all trials: 2.5-fold greater than this threshold (~ 215 g/day providing ~35% excess energy) in the hypercaloric trials, in which there was an effect, and 1.4-fold greater than this threshold (115 g/day) in the isocaloric trials, in which there was no effect. Also, no trials used non-beverage grain or fruit sources of fructose, which together account for >30% of fructose in the U.S. diet38 and have been linked (as whole grains and fruits) to weight loss and improved metabolic outcomes in large prospective cohort studies39,41 and randomized trials.43,44 Dietary trials of more representative sources of fructose at more representative levels of exposures remain a research priority.

CONCLUSIONS
In conclusion, our preliminary systematic review and meta-analysis does not support a NAFLD-inducing effect of fructose in isocaloric exchange for other carbohydrates at levels of exposure that are
well above that found in Western diets. The evidence does support an IHCL- and ALT-increasing effect of diets supplemented with fructose providing excess energy (+ 21–35% energy) at extreme doses (104–220 g/day). Confounding from excess energy, however, cannot be excluded in the hypercaloric trials, such that the observed NAFLD-inducing effect is more attributable to the excess energy than the fructose itself. Other sources of uncertainty in our analyses include the small number of available trials (four for IHCL and six for ALT), as well as the relatively small sample sizes (<30 participants/trial) and narrow participant demographics (most participants were young and relatively healthy). The short follow-up (all trials were <12-weeks) using IHCL and ALT as markers of NAFLD may also not be relevant to the natural history of NAFLD over the longer term, especially in people who may be at low risk. It is unclear whether a larger number of trials which address these many issues will show the same findings when analyzed collectively. To understand the role of fructose in the epidemic of NAFLD, there remains a need for larger, longer, high-quality trials of the effect of ‘real-world’ intake patterns of fructose on histopathological changes of NAFLD in at risk populations.

CONFLICT OF INTEREST

JLS has received research support from the Canadian Institutes of Health Research (CIHR), Calorie Control Council, The Coca-Cola Company (investigator initiated, unrestricted grant), Pulse Canada and The International Tree Nut Council Nutrition Research & Education Foundation. He has received travel funding, speaker fees and/or honoraria from the American Heart Association (AHA), American College of Physicians (ACP), American Society for Nutrition (ASN), National Institute of Diabetes and Digestive and Kidney Diseases (NIDDK) of the National Institutes of Health (NIH), Canadian Diabetes Association (CDA), Canadian Nutrition Society (CNS), Calorie Control Council, Diabetes and Nutrition Study Group (DNSG) of the European Association for the Study of Diabetes (EASD), International Life Sciences Institute (ILSI) North America, International Life Sciences Institute (ILSI) Brazil, University of South Carolina, Calorie Control Council, Canadian Sugar Institute, Abbott Laboratories, Pulse Canada, Dr Pepper Snapple Group, and The Coca-Cola Company. He is on the Clinical Practice Guidelines Expert Committee for Nutrition Therapy of both the Canadian Diabetes Association (CDA) and European Association for the study of Diabetes (EASD), as well as being on the American Society for Nutrition (ASN) writing panel for a scientific statement on the metabolic and nutritional effects of fructose, sucrose and high fructose corn sirup. He is a member of the Transcultural Diabetes Algorithm (tDNA) Collaborative Group and the International Carbohydrate Quality Consortium (ICQC). He is an unpaid scientific advisor for the International Life Science Institute (ILSI) North America, Food, Nutrition, and Safety Program (FNSP). His wife is an employee of Unilever Canada. RJDS has received research support from the CIHR, Calorie Control Council, the Canadian Foundation for Dietetic Research (CFDR), and The Coca-Cola Company (investigator initiated, unrestricted grant). He has served as an external resource person to the World Health Organization’s (WHO) Nutrition Guidelines Consultative Revision (NUGAG) and was a lead author of a systematic review and meta-analysis commissioned by the WHO of trans-fatty acids and health outcomes. The WHO paid for his travel and accommodation to attend NUGAG Meetings in Hangzhou, China and Copenhagen, Denmark. AIC and VH received a travel award to attend the ‘Journey Through Science Day’ hosted by PepsiCo and the New York Academy of Sciences (NYAS). VH has also received research support from the CIHR and WHO for work on a systematic review and meta-analysis commissioned by the WHO of the relation of saturated fatty acids with health outcomes. AM and AIC have received research support from the CIHR. ALJ is a part owner, Vice-President, and Director of Research of Glycemic Index Laboratories, Toronto, Canada. She has received research support from the CDA. TMSW is a part owner and the President of Glycemic Index Laboratories, Toronto, Canada. She has received research support from the CDA. TAMSG is a part owner and the President of Glycemic Index Laboratories, Toronto, Canada and has authored several popular diet books on the glycemic index for which he has received royalties from Phillipa Sandall Publishing Services and CABI Publishers. He has received consultant fees, honoraria, travel funding, or research support from or served on the scientific advisory board for CIHR, CDA Dairy Farmers of Canada, McCain Foods, Temasek Polytechnic, Northwestern University, Royal Society of London, Glycemic Index Symbol program, CreaNutrition AG, McMaster University, Canadian Society for Nutritional Sciences, National Sports and Conditioning Association, Faculty of Public Health and Nutrition—Autonomous University of Nuevo Leon Diabetes and Nutrition Study Group of the European Association for the Study of Diabetes. 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DJAJ has received consultant fees, honoraria, travel funding or research support from or served on the scientific advisory board for the CIHR, Canadian Foundation for Innovation (CFI), Ontario Research Fund (ORF), and Advanced Foods and Material Network (AFINet) Calorie Control Council, The Coca-Cola Company (investigator initiated, unrestricted), Barilla, Solae, Unilever, Hain Celestial, LoblawS Supermarkets, Inc., Sanitarium Company, Herbalife International, Pacific Health Laboratories, Inc., Metabolon and Metaproteomics, Bayer Consumer Care, Oldways Preservation Trust, The International Tree Nut Council Nutrition Research & Education, The Peanut Institute, Procter and Gamble Technical Center Limited, Griffin Hospital for the development of the NuVal System, Soy Advisory Board of Dean Foods, Alpro Soy Foundation, Nutritional Fundamentals for Health, Pacific Health Laboratories, Kellogg’s, Quaker Oats, The Coca-Cola Sugar Advisory Board, Pepsi Company, Agrifoods and Agriculture Canada (AAFC), Canadian Agriculture Policy Institute (CAPI), The Almond Board of California, The California Strawberry Commission, Orafti, the Canola and Flax Councils of Canada, Pulse Canada, the Saskatchewan Pulse Growers, and Abbott Laboratories. The remaining authors declare no conflict of interest.

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