ABSTRACT

High-glycemic index (high-GI) foods (so-called fast carbs) have been hypothesized to promote fat storage and increase risk of obesity. To clarify whether dietary GI impacts body weight, we searched PubMed and the Cochrane Database of Systematic Reviews for observational studies reporting associations between BMI and dietary GI, and for meta-analyses of randomized controlled trials (RCTs) comparing low-GI and high-GI diets for weight loss. Data on 43 cohorts from 34 publications, totaling 1,940,968 adults, revealed no consistent differences in BMI when comparing the highest with the lowest dietary GI groups. In the 27 cohort studies that reported results of statistical comparisons, 70% showed that BMI was either not different between the highest and lowest dietary GI groups (12 of 27 cohorts) or that BMI was lower in the highest dietary GI group (7 of 27 cohorts). Results of 30 meta-analyses of RCTs from 8 publications demonstrated that low-GI diets were generally no better than high-GI diets for reducing body weight or body fat. One notable exception is that low-GI diets with a dietary GI at least 20 units lower than the comparison diet resulted in greater weight loss in adults with normal glucose tolerance but not in adults with impaired glucose tolerance. While carbohydrate quality, including GI, impacts many health outcomes, GI as a measure of carbohydrate quality appears to be relatively unimportant as a determinant of BMI or diet-induced weight loss. Based on results from observational cohort studies and meta-analyses of RCTs, we conclude that there is scant scientific evidence that low-GI diets are superior to high-GI diets for weight loss and obesity prevention. Adv Nutr 2021;12:2076–2084.

Introduction

The glycemic index (GI) was introduced in 1981 as a means to classify foods according to their effects on postprandial blood glucose (1). Since then, >10,000 scientific articles have been published on GI (PubMed search May 2021), and several popular books have extolled the purported health benefits of low-GI diets (2–4), including better weight control and reduced obesity risk (5, 6). High-GI foods are frequently referred to as “fast carbs.” A May 2021 Google search for “fast carbs” produced >47,000 results, many of which featured websites that portrayed fast carbs as unhealthier and more fattening than low-GI “slow carbs.” Despite popular perception of the superiority of low-GI diets for weight loss and obesity prevention, published research on the topic has produced conflicting interpretations of results (5–14).

Several highly cited reviews have been published on the health implications of a high-GI diet (5, 6, 10, 11, 14), and a 2015 scientific consensus statement concluded diets low in GI were “probably” relevant to the prevention of obesity (8). In contrast, the 2010 US Dietary Guidelines Advisory Committee concluded that there was strong and consistent evidence showing that dietary GI was not associated with...
body weight (7). The conclusion that GI is not strongly or consistently associated with body weight is also supported from the results of 3 previous narrative reviews (9, 12, 13). The 2015 and 2020 US Dietary Guidelines Advisory Committees made no recommendations for using GI in dietary guidelines (15, 16).

More recently, a 2020 review on the importance of carbohydrate quality over quantity indicated that systematic reviews and meta-analyses of >50 randomized controlled trials (RCTs) have shown that low-GI dietary patterns lead to weight loss (17). However, some RCTs used control diets for which GI was not reported. Moreover, dietary fiber, which impacts body weight (18), was not always measured or reported as the dietary GI changed. Thus, inadequate reporting and residual confounding make attributing changes in body weight entirely to a low-GI diet problematic.

Central to the hypothesized link between high-GI diets and excess body weight is the carbohydrate-insulin model of obesity (19). This model proposes that high-GI foods are particularly fattening because they elevate postprandial insulin secretion, which has direct effects on accelerating storage of fat. However, the validity of the carbohydrate-insulin model of obesity has been questioned (17, 20–22). Although postprandial increases in insulin are greater after high-GI meals (23), most intervention studies show that low-GI diets are no better than high-GI diets for reducing fasting insulin concentrations (24–27).

To clarify whether dietary GI is important for weight control and obesity prevention, we searched PubMed and the Cochrane Database of Systematic Reviews for meta-analyses of observational studies that compared BMI of individuals across dietary GI strata and RCTs that compared low-GI and high-GI diets for weight loss. For observational studies, the search strategy included “glycemic index” OR “glycaemic index” AND “body mass index.” For the RCTs, the search strategy included “glycemic index” OR “glycaemic index” AND “weight” OR “body fat” OR “obesity.” Search results were filtered for “meta-analysis” and were limited to adult populations. No restrictions were placed on date of publication. Of the 56 results for the RCT search, 8 publications were identified that presented 1 or more meta-analyses that compared low-GI and high-GI diets for changes in body weight and/or body fat indices. However, we found no meta-analyses of observational studies, so we removed the “meta-analysis” filter. This produced 892 results, of which 35 publications provided required data on GI and BMI. Reference lists and electronic citation records of all identified meta-analyses and observational studies were also reviewed for additional publications not found in the initial searches.

**BMI in Relation to Dietary GI: Observational Cohort Studies**

A total of 35 observational studies, including data from 43 cohorts, provided information on baseline BMI across tertiles, quartiles, or quintiles of dietary GI (28–62). These included 18 cohorts of women, 8 cohorts of men, and 17 cohorts of combined women and men, with a total of 1,940,968 adults. In 27 cohorts, statistical analyses were performed to determine whether BMI differed significantly in the highest compared with lowest GI groups (Figure 1). In 16 cohorts, no statistical analyses were reported to assess whether BMI differed between the highest and lowest GI groups (Figure 2).

In the 27 cohorts for which statistical analyses were performed to compare BMI between GI groups, 12 showed no difference in BMI between the highest and lowest GI groups and 7 indicated that BMI was lower in the highest GI groups. Thus, in 70% of the 27 cohorts, dietary GI had either no association with BMI or high GI was associated with lower BMI. In only 8 of the 27 cohorts was BMI significantly lower in the group ingesting the lowest dietary GI.

BMI was not the primary outcome in any of these epidemiological studies. Thus, even when statistical differences were reported, the results must be viewed guardedly as BMI can be affected by many variables, such as total energy intake, fiber intake, and physical activity. Few, if any, adjustments were made for these potential confounders. In studies that did report statistical comparisons across GI groups regarding these potential confounders, no consistent pattern was evident. In studies that found BMI to be significantly lower in the highest GI groups, total energy intake in the highest GI group was significantly higher than in the lowest GI group in 3 cohorts (39, 46, 53) and lower in 1 cohort (28). Five cohorts that reported lower BMI in the highest GI group had significantly lower intakes of dietary fiber (28, 30, 39, 46, 53), whereas only 1 cohort with lower BMI in the highest GI group had significantly higher dietary fiber intake (61). Similar interactions were observed between physical activity and GI. For example, in 3 cohorts that showed lower BMI in the highest dietary GI groups, physical activity levels were also lower (39, 46, 61). In contrast, in only 1 study that showed lower BMI in the highest dietary GI group was physical activity level significantly higher (53).

Collectively, these results suggest that the significantly lower BMIs observed in the highest dietary GI group were not attributable to lower energy intake, lower fiber intake, or higher levels of physical activity. In contrast, in 4 of the 7 studies that reported higher BMI in the 3rd quartile of dietary GI, the results could be due in part to significantly higher total energy intake (31), lower intake of fiber (28, 31, 33, 38), and less physical activity (28, 33).
In the 16 cohorts for which no statistical analysis was performed, the mean BMI for the lowest and highest GI groups was virtually the same. This was observed across male, female, and mixed-sex cohorts (Figure 2). The reported BMI in the highest GI group was lower than that in the lowest dietary GI group in 8 of 16 cohorts and was the same in 3 of the cohorts. These observations are consistent with results of studies that reported statistical comparisons. Additional data reported in these studies suggest that these observations were not materially influenced by differences in total energy intake, fiber intake, or physical activity. For example, in the Nurses’ Health Study (NHS) I and NHS II, BMI was either exactly the same in the highest and lowest quintiles of dietary GI (NHS I) or 0.3 kg/m² lower in the highest quintile of dietary GI (NHS II), yet total energy intake was essentially the same across quintiles and physical activity levels were inversely related to GI (29). Among men in the Health Professionals Follow-Up Study, BMI was 0.6 kg/m² lower in the highest compared with the lowest quintiles of GI despite similar total energy intake and lower levels of physical activity (29). Similarly, for men and women in the European Prospective Investigation into Cancer and Nutrition cohort, BMI was 0.2 kg/m² lower in the highest compared with the lowest quintile of dietary GI, despite similar total energy and fiber intake and lower levels of physical activity (41).

In the DiOGenes (Diet, Obesity, and Genes) study, there was a trivial difference in BMI (in kg/m²) between the lowest (25.7) and highest (25.9) quintiles, yet the highest GI quintile consumed nearly 400 kcal/d more than the lowest GI quintile (32). Thus, the slightly higher BMI in the highest GI quintile could be simply due to increased energy intake.

Only 1 of the cohort studies reported data on body fat, assessed by DXA (31). For 5830 adults in the Health Worker Cohort Study, BMI was significantly higher in the highest dietary GI quartile (26.9) compared with the lowest dietary GI quartile (26.6), but percentage body fat did not differ across quartiles (30.9% in quartile 1 vs. 31.4% in quartile 4) (31).

Several epidemiological studies have reported that the prevalence of overweight and obesity differs by very little across dietary GI categories. In the DiOGenes cohort, the percentage of individuals with obesity was essentially the
same in quintile 5 (12%) and quintile 1 (13%) of dietary GI, although the percentage of overweight participants was slightly higher in quintile 5 compared with quintile 1 (43% vs. 38%) (32). In the Black Women’s Health Study, the percentage of women with a BMI >30 (range = 28.2% to 29.5%) was similar across quintiles of dietary GI (63). In the NIH-AARP Diet and Health Study, the percentage of men with a BMI >25 was similar for quintile 1 (68.8%) and quintile 5 (69.1%) of dietary GI, but the percentage of women with a BMI >25 was higher in quintile 5 (59.9%) than in quintile 1 (48.8%) (64). The higher prevalence of women with a BMI >25 in quintile 5 compared with quintile 1 of dietary GI could be attributed in part to higher daily energy intake (1651 vs. 1499 kcal/d), lower fiber intake (13.8 vs. 22.1 g/d), and a lower percentage who regularly engaged in physical activity (11.0% vs. 23.6%) (64). Also, in the Framingham Offspring Cohort, waist circumference was not different across quintiles of dietary GI (65). Among older adults in the National Diet and Nutritional Survey, dietary GI was not correlated with BMI, body weight, or waist-to-hip ratio (66). In sum, data from these cohort studies do not support the hypothesis that high dietary GI is associated with a higher BMI or greater prevalence of obesity.

Results from RCTs

Eight publications presented a total of 30 meta-analyses of RCTs comparing low-GI and high-GI diets (24–27, 67–70). The RCTs in these meta-analyses were behavioral interventions in which participants received dietary advice. In some instances, participants were supplied with key foods that were aligned with the intervention GI. These studies examined the effects of dietary GI on several anthropometric outcomes, including body weight, BMI, body fat, and waist circumference. Results of these meta-analyses are presented in Table 1, which includes information on the populations examined and the number of studies in each meta-analysis.

Body weight

Eight publications, including a total of 15 meta-analyses, reported on body weight as an outcome measure. With only 2 exceptions (26, 70), low-GI diets provided no benefit over high-GI diets for weight loss. One small meta-analysis with only 4 RCTs reported that a low-GI diet produced significantly greater weight loss (≈1 kg) compared with a high-GI diet (26). In a meta-analysis of 28 RCTs by Zafar et al. (70) that included adults with normal glucose tolerance (NGT), impaired glucose tolerance (IGT), and type 2 diabetes (T2D), significantly greater weight loss was observed for a low-GI diet compared with a high-GI diet (standardized mean difference = 0.14, corresponding to ≈1.8 kg), but only if the low-GI diet was ≥20 units lower than the high-GI diet. However, when analyzed by glucose tolerance, the greater weight loss with the low-GI diet was observed only in NGT, with no differences between diets in
IGT and T2D. In the combined meta-analysis that included all 51 RCTs (28 RCTs with ≥20 units lower GI plus 23 RCTs with a GI difference <20 units), low-GI diets did not produce greater weight loss than high-GI diets (Table 1). In another meta-analysis by Zafar et al. (27), no significant differences in weight loss were observed among subjects with IGT, type 1 diabetes, or T2D, regardless of whether the GI of the low-GI diet was ≥20 units lower than the high-GI diets. Collectively, these meta-analyses suggest that weight loss with low-GI diets may be more effective than high-GI diets only if subjects have NGT and the GI difference between the diets is ≥20 units.

### Fat mass and body fat percentage
Change in fat mass was reported in 4 of the 8 meta-analyses (25, 26, 69, 70), and 1 study also reported results for body fat mass and % body fat (70). In a meta-analysis of 4 RCTs with overweight and obese subjects, low-GI diets resulted in 1.13-kg greater body fat loss (26). Among individuals with NGT (70), consumption of a low-GI diet that differed by ≥20 units from the comparison high-GI diet resulted in more fat loss than the high-GI diet (Table 1). In contrast, among individuals with IGT, and in the combined group of individuals with IGT, NGT, and T2D, a low-GI diet that was ≥20 units lower than the high-GI diet did not result in greater fat loss (70). When the GI difference between diets was unstipulated, the percentage of fat loss did not differ between high- and low-GI diets (25, 69, 70). In the only meta-analysis that reported on changes in body fat percentage, no differences were observed between low-GI and high-GI diets among individuals with NGT, IGT, or T2D (70).

### Fat-free mass
Two relatively small meta-analyses, that included only 2 or 3 RCTs, reported data on changes in fat-free mass in overweight and obese subjects. One reported no difference

| Table 1 | Effects of low-GI diets compared with high-GI diets on body weight, body fat, fat-free mass, and waist circumference: results from meta-analyses of RCTs |
|---------|---------------------------------------------------------------|
| **Meta-analysis** | **Subject characteristics** | **Number of RCTs included** | **Difference between low-GI and high-GI diets** |
| **Body weight (kg)** | | | |
| Kelly et al., 2004 (24) | At risk for CVD | 13 | 0.14 (−0.68, 0.95) kg |
| Thomas et al., 2007 (26) | Overweight, obese | 4 | −1.09 (−1.99, −0.18) kg |
| Ajala et al., 2013 (67) | T2D | 3 | 1.39 (−1.58, 4.36) kg |
| Schwingshackl et al., 2013 (69) | Overweight, obese, T2D | 14 | −0.62 (−1.28, 0.03) kg |
| Schwingshackl et al., 2013 (69) | Obese, T2D | 9 | −1.26 (−2.17, −0.34) kg |
| Schwingshackl et al., 2013 (69) | Overweight, T2D | 6 | 0.04 (−0.90, 0.98) kg |
| Clar et al., 2017 (68) | At risk for CVD | 20 | −0.16 (−0.54, 0.21) kg |
| Reynolds et al., 2019 (25) | Overweight, obese, IGT, T2D | 8 | −0.29 (−0.62, 0.03) kg |
| Zafar et al., 2019 (70) | NGT, IGT, T2D | 51 | −0.06 (−0.15, 0.03) SMD |
| Zafar et al., 2019 (70) | NGT, IGT, T2D; GI difference ≥20 units | 28 | −0.14 (−0.25, −0.03) SMD |
| Zafar et al., 2019 (70) | NGT; GI difference ≥20 units | 13 | −0.26 (−0.43, −0.09) SMD |
| Zafar et al., 2019 (70) | IGT; GI difference ≥20 units | 8 | −0.07 (−0.28, 0.14) SMD |
| Zafar et al., 2019 (70) | T2D; GI difference ≥20 units | 7 | −0.02 (−0.22, 0.18) SMD |
| Zafar et al., 2019 (27) | IGT, T1D, T2D | 22 | 0.00 (−0.92, 1.92) kg |
| Zafar et al., 2019 (27) | IGT, T1D, T2D; GI difference ≥20 units | 12 | −0.64 (−3.33, 2.05) kg |
| **Body fat (kg)** | | | |
| Thomas et al., 2007 (26) | Overweight, obese | 4 | −1.13 (−1.89, −0.38) kg |
| Schwingshackl et al., 2013 (69) | Overweight, obese | 5 | −0.56 (−1.24, 0.12) kg |
| Reynolds et al., 2019 (25) | Overweight, obese, IGT, T2D | 5 | −0.27 (−0.79, 0.26) kg |
| Zafar et al., 2019 (70) | NGT, IGT, T2D | 24 | −0.09 (−0.19, 0.02) SMD |
| Zafar et al., 2019 (70) | NGT | 17 | −0.10 (−0.21, 0.01) SMD |
| Zafar et al., 2019 (70) | IGT | 5 | −0.06 (−0.38, 0.26) SMD |
| Zafar et al., 2019 (70) | NGT, IGT, T2D; GI difference ≥20 units | 12 | −0.15 (−0.35, 0.04) SMD |
| Zafar et al., 2019 (70) | NGT; GI difference ≥20 units | 7 | −0.28 (−0.52, −0.04) SMD |
| Zafar et al., 2019 (70) | IGT; GI difference ≥20 units | 4 | 0.17 (−0.24, 0.58) SMD |
| **Body fat percentage** | | | |
| Zafar et al., 2019 (70) | NGT, IGT, T2D | 21 | 0.00 (−0.14, 0.13) SMD |
| Zafar et al., 2019 (70) | NGT | 17 | −0.10 (−0.21, 0.01) SMD |
| Zafar et al., 2019 (70) | IGT | 5 | −0.06 (−0.38, 0.26) SMD |
| **Fat-free mass (kg)** | | | |
| Thomas et al., 2007 (26) | Overweight, obese | 2 | −0.13 (−0.03, 0.56) kg |
| Schwingshackl et al., 2013 (69) | Overweight, obese | 3 | −1.04 (−1.73, −0.35) kg |
| **Waist circumference (cm)** | | | |
| Schwingshackl et al., 2013 (69) | Overweight, obese, IGT, T2D | 10 | 0.06 (−0.83, 0.96) kg |

1. CVD, cardiovascular disease; GI, glycemic index; IGT, impaired glucose tolerance; NGT, normal glucose tolerance; RCT, randomized controlled trial; SMD, standardized mean difference; T1D, type 1 diabetes; T2D, type 2 diabetes.
between diets (26), whereas one reported that low-GI diets resulted in significantly greater loss (1.04 kg) of fat-free mass (69).

Waist circumference

Only 1 meta-analysis has been published comparing the effects of low-GI and high-GI diets on waist circumference (69). This meta-analysis of 10 RCTs demonstrated that low-GI and high-GI diets did not differ with regard to changes in waist circumference.

Expected Compared with Observed Results: Problems with Interpreting Research Findings

High-GI meals consistently result in greater glucose and insulin secretion (23) and lower postprandial fat oxidation than low-GI meals (71). These findings are consistent with the carbohydrate-insulin model of obesity and the hypothesis that low-GI diets are associated with greater weight loss and reduced risk of obesity. However, data from observational studies and meta-analyses of RCTs do not substantiate the superiority of low-GI diets for weight loss or obesity prevention. Interestingly, body weight and fat loss were observed following low-GI diets in individuals with NGT but not in those with IGT (70), where insulin responses to a glycemic challenge are almost always greater. If the carbohydrate-insulin model of obesity were to hold true, it would be expected that individuals with the greatest insulin responses to meals would show the greatest reductions in body weight in response to a low-GI diet. It must be noted, however, that most intervention studies showed that low-GI diets are no better than high-GI diets for reducing fasting insulin (24–27). Even in controlled-feeding studies, fasting insulin concentration is not affected by the GI of the diet (72).

Several physiological and methodological limitations of GI research may help explain our null findings. High-GI meals rarely predict short-term energy intake (73). Also, GI values found in tables are measured under rigidly controlled laboratory conditions where foods are eaten singly by generally healthy young subjects and typically involve small sample sizes. This may not represent real-world eating situations because most foods are rarely ingested singly and in prescribed amounts. The glycemic response to a meal with carbohydrate-containing foods can change depending upon the macronutrient composition and dietary fiber content of the meal, preparation of the food, and the time of day that the food is consumed (74–78). Using the published or measured GI value of individual foods to determine the GI of a meal has been reported to overestimate the directly measured GI of a meal by 12 to 19 GI units (75). For perspective, in the observational studies presented in Figures 1 and 2, the difference in median GI between the highest and lowest dietary GI groups rarely exceeded 10 GI units.

Accurate assignment of GI values to foods from a diet record or FFQ is difficult (79). Factors that strongly affect a food’s GI, such as variety and cooking/processing methods, are rarely specified in sufficient detail to ensure that the actual GI of the food eaten accurately reflects the assigned GI. For instance, the University of Sydney’s GI database lists 27 values for brown rice ranging from 48 to 87 and 66 GI values for white rice ranging from 17 to 94 (80). Moreover, significant interindividual variability and intrapersonal reproducibility in repeat GI testing under controlled conditions further complicate interpretation of GI data (81, 82). In a cohort of nearly 800 adults, the glycemic response to white bread varied by >5-fold when comparing the bottom 10% and top 10% of the individual postprandial glycemic responses (82). Finally, glycemic responses across individuals are highly heterogeneous and subject to variation due to age, genetics, physical activity, insulin sensitivity, and BMI (82).

Interpretation of results from observational studies is also limited by the inherent inaccuracies of self-reported diet and physical activity data. Energy intake data from self-report invariably underestimates energy intake determined from doubly-labeled water (DLW) (83, 84), and self-reported physical activity data may not accurately correspond to objectively measured physical activity from accelerometry or physical activity energy expenditure from DLW (85).

Conclusions

Data from observational cohort studies show no consistent association between BMI and dietary GI, and results of meta-analyses of RCTs provide little support for the notion that low-GI diets are superior for weight loss. In the 27 cohort studies that performed statistical comparisons of BMI by dietary GI, the majority (70%) reported either no differences in BMI between extremes of dietary GI or a significantly lower BMI in the highest dietary GI group. We acknowledge that our results may not include all published observational studies that show data on BMI and dietary GI. However, we doubt that additional cohorts missed in our search would change the interpretation regarding BMI–GI relations of the 43 cohorts presented herein. Moreover, previous reviews of the association between BMI and dietary GI (9, 12, 13) included <10 cohorts.

Similarly, results of RCTs generally do not support a case for greater weight loss with low-GI diets. The 1 notable exception is that low-GI diets with a dietary GI at least 20 units lower than the comparison diet resulted in greater weight loss, but this was only observed in individuals with NGT. Because low-GI diets may result in greater loss of fat-free mass (69), this may explain why none of the meta-analyses showed a benefit of low-GI diets for reducing body fat percentage. Also, it must be noted that the RCTs included in the meta-analyses were behavioral interventions in which participants received dietary advice. Although controlled-feeding trials are necessary to determine efficacy of dietary GI for weight loss, the RCTs used in the meta-analyses in Table 1 are more relevant to the effectiveness of dietary GI in real-world conditions.

In view of all the contrary evidence from observational cohort studies and meta-analyses of RCTs, it is surprising that the hypothesis that low-GI (“slow-carb”) diets are superior...
for weight loss and obesity prevention persists. Carbohydrate quality, including GI, clearly impacts many health outcomes (17, 18, 25), and several meta-analyses have reported higher risks of T2D, cardiovascular disease, and stroke associated with high-GI diets (25, 86–89). However, GI as a measure of carbohydrate quality appears to be unimportant as a determinant of BMI or diet-induced weight loss. We contend that GI is an imprecise measure of the glycemic response of a food when applied to foods in a meal, and that the GI assigned to foods from an FFQ may differ significantly from actual GI. Nutrient density, dietary fiber and whole-grain content of carbohydrates, and percentage of added sugar, are more important qualities (17, 18, 78). Further, a focus on staple carbohydrate foods and the positive nutrients they contribute to diet quality as compared with the detractor nutrients associated with indulgent foods is important in characterizing the quality of carbohydrates. As for body weight and obesity, there is scant scientific evidence that low-GI diets are superior to high-GI diets for weight loss and obesity prevention.

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