Review Article

Exercising Tactically for Taming Postmeal Glucose Surges

Elsamma Chacko

Connecticut Valley Hospital, 100 Silver Street, Middletown, CT 06457, USA

Correspondence should be addressed to Elsamma Chacko; elssmac@msn.com

Received 6 December 2015; Revised 28 January 2016; Accepted 2 March 2016

Academic Editor: Naohiko Ueno

Copyright © 2016 Elsamma Chacko. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

This review seeks to synthesize data on the timing, intensity, and duration of exercise found scattered over some 39 studies spanning 3+ decades into optimal exercise conditions for controlling postmeal glucose surges. The results show that a light aerobic exercise for 60 min or moderate activity for 20–30 min starting 30 min after meal can efficiently blunt the glucose surge, with minimal risk of hypoglycemia. Exercising at other times could lead to glucose elevation caused by counterregulation. Adding a short bout of resistance exercise of moderate intensity (60%–80% VO2max) to the aerobic activity, 2 or 3 times a week as recommended by the current guidelines, may also help with the lowering of glucose surges. On the other hand, high-intensity exercise (>80% VO2max) causes wide glucose fluctuations and its feasibility and efficacy for glucose regulation remain to be ascertained. Promoting the kind of physical activity that best counters postmeal hyperglycemia is crucial because hundreds of millions of diabetes patients living in developing countries and in the pockets of poverty in the West must do without medicines, supplies, and special diets. Physical activity is the one tool they may readily utilize to tame postmeal glucose surges. Exercising in this manner does not violate any of the current guidelines, which encourage exercise any time.

1. Introduction

The value of exercise in keeping body and mind in good health was an article of faith among physicians of the time of Hippocrates. More recently, before the insulin era began, doctors had been prescribing exercise for diabetes. Now, decades of clinical studies have established that blood glucose levels are sensitive to exercise timing, intensity, duration, and frequency [1, 2]. A 2013 review [1] concluded that postmeal exercise was better than premeal exercise for managing hyperglycemia.

In fact, it is no longer adequate to categorize exercise timing into just premeal and postmeal in diabetes studies. The postmeal period is an eventful stretch of time [3], unlike the usually quiet premeal period. In the healthy human body, insulin and the counterregulatory hormones work in tandem to keep blood glucose levels within normal bounds [4]. Glucose levels peak around the 1-hour mark after meal and decline to the premeal level in two to four hours [3]. Glucose levels in healthy people may not go above 140 mg/dL. In people with diabetes, a substantial peak could develop with the height, width, slopes, and peaking time showing wide variations based on several factors, including the state of diabetes, meal size, meal composition, level of activity, and medications.

This paper combs published data looking for ways to tame the postmeal glucose surge [5] through exercise without triggering hypoglycemia [1, 6, 7]. The assembled database consists of 39 articles [8–46] that report timing, intensity, and duration of the exercise activity. The feeding cycle is perceived as consisting of four time segments: before meal, early postprandial (0–29 min after meal), midpostprandial (30 min to 120 min after meal), and late postprandial (>120 min after meal). Exercise intensity in particular is measured using a large number of variables: %VO2max (maximal oxygen uptake), %VO2peak (peak oxygen uptake), %HRmax (heart rate max, calculated as 220 – age), %HRR (heart rate reserve), and %Wmax (maximal power output) are some of the commonly used units. The categorization of exercise intensity, a continuous variable, into light (<60% VO2max), moderate (60%–80% VO2max), and high (>80% VO2max), is dictated by the distribution of the values found in the assembled database. These categories serve the purpose extant here and may conform only approximately to others used elsewhere.
High-intensity premeal exercise resulted in significant postexercise glucose elevation in healthy men, men with type 2 diabetes, and people with type 1 diabetes [31–34]. These studies reported elevated levels of epinephrine, norepinephrine, glucagon, lactate, and pyruvate and depressed levels of glycerol, free fatty acids, and 3-hydroxybutyrate during the exercise activity [31, 33, 34, 54]. Some improvement in insulin sensitivity that lasted for up to 24 hours was also observed [15, 31, 47–51]. In people with type 1 diabetes nocturnal hypoglycemia was seen after high-intensity premeal exercise [6, 47, 51].

3.2. Early Postprandial Period. When light activity began 15 minutes after the start of meal, glucose levels kept rising during the first 15 minutes of the exercise at the end of which, starting at 30 min after meal, glucose started to go down [22–24]. The secondary peak that formed after the activity stopped was prominent, signaling suboptimal blunting of the glucose peak. The subjects in these studies were healthy men and women and people with type 1 diabetes. One study using moderate intensity (65% VO$_{2\text{max}}$) exercise at 15 min after meal reported that a one-third reduction in postmeal glucose elevation occurred when g-AUC was compared between exercising and nonexercising subjects [25].

3.3. Midpostprandial Period. Twenty-two midpostprandial studies using light to moderate exercise demonstrated glucose lowering with the activity [8–14, 17–19, 26–30, 44–46, 55–59]. These efforts spanned virtually all demographics of interest in diabetes studies: healthy women, healthy men, men with type 2 diabetes, women with type 2 diabetes, people with type 2 diabetes on insulin, men with metabolic syndrome, patients with type 1 diabetes, and healthy postmenopausal women. When the light to moderate exercise started 30 [12, 19, 26–28] or 45 min [17, 29, 44] after meal, there was good blunting of the postmeal glucose peak: for example, Larsen and colleagues estimated that the postprandial glucose surge was diminished by 50%, according to g-AUC calculations, when the subjects exercised for 45 min starting at 45 min after meal, at 53% VO$_{2\text{max}}$ [29].

High-intensity postmeal exercise during midpostprandial period did not result in significant postexercise glucose elevation in contrast to the glucose elevation found in the case of high-intensity premeal exercise. A delayed insulin sensitivity improvement was discernible [35–38]. The subjects in these studies were adults with type 2 diabetes, healthy adults, and inactive obese men.

3.4. Late Postprandial Period. The glycemic effect of light exercise during the late postprandial period (4.5 h) showed postprandial glucose elevation in people with prediabetes, reminiscent of the premeal effect [13]. Sasaki and colleagues demonstrated that the glycemic response to premeal exercise was nearly identical to that of the 4 h postmeal exercise in young trained athletes [56].

High-intensity exercise at three hours after the meal in healthy men showed a significant postexercise glucose peak.

### 2. Search and Selection

The articles at the heart of this review came from a related project concluded in 2014 [2]. Iterative searches of MEDLINE using the search terms “exercise timing,” “post meal exercise,” “pre meal exercise,” “postprandial exercise,” “post-absorptive exercise,” and “HIT exercise.” Modifiers used in the database search were “high-intensity,” “intense,” “moderate,” “light,” “glucose,” “metabolic syndrome,” “obesity,” and “diabetes.” Other relevant studies came from the reference lists of landmark articles and from a hand search of appropriate journals and reviews. Studies of blood glucose response to a single bout of light to moderate exercise (<80% VO$_{2\text{max}}$) during premeal or postmeal period found their way into the database if intensity and duration were also among the study variables. The health benefits of high-intensity interval training (HIT) were well established through recent reviews [47–53]. A few articles from this group that showed glucose surges were included. Excluded were studies focused on glucose response to different meals or on medications along with exercise. This search did not use any cut-off dates.

Out of the 39 articles [8–46] found in Table 1, 23 studied the glucose response to light to moderate (<80% VO$_{2\text{max}}$) intensity exercise, before meal [8–21] or after meal [22–30]. Eight studied high-intensity (>80% VO$_{2\text{max}}$) exercise, before meal or after meal [31–38]. One study used premeal and postmeal resistance exercise [39]. The remaining seven articles compared effects such as training in fasted versus fed state [40–43], different intensities [44, 45], and different durations [46].

### 3. Results

3.1. Premeal Period. In general, glucose levels were steady during light to moderate premeal exercise and there was little risk of hypoglycemia in a wide range of populations: men with type 2 diabetes, individuals with insulin resistance, and healthy men and healthy women [8–10, 17, 19]. When glucose did go down, the extent of the decline depended on its preexercise level [8]. Light to moderate premeal exercise, however, led to postprandial glucose elevations immediately following the exertion. This effect was consistently seen in men with metabolic syndrome, men and women with type 2 diabetes, men and women with prediabetes, individuals with type 1 diabetes, and people with insulin resistance [11–15]. Melton and colleagues observed that moderate premeal exercise in people with prediabetes did not decrease oxidative stress in people with prediabetes; oxidative stress is closely linked to glucose surges [16]. Borer and colleagues found improved fasting glucose levels in healthy postmenopausal women after they completed two-two hour bouts of light premeal exercise [18]. The authors inferred liver glycogen depletion from markedly elevated levels of 3-hydroxybutyrate, free fatty acids, cortisol, and glucagon during the premeal exercise. A direct measurement showed that hepatic glucose production remained elevated during moderate premeal exercise [19]. At relatively high intensities (75% VO$_{2\text{max}}$) premeal exercise also showed a delayed, modest insulin sensitivity improvement according to glucose area-under-the-curve (g-AUC) calculations [20, 21].
Table 1: Glucose response to different exercise conditions: moderate exercise, before meal [8–21] and after meal [22–30]; high-intensity exercise, before meal [31–34] and after meal [35–39]; comparisons of training, fast versus fed [40–43], timings [44, 45], and durations [46].

| Study | Subjects | Exercise protocol | Results |
|-------|----------|------------------|---------|
| Gaudet-Savard et al. [8] | 43 men with T2D | Light, before meal for 1 h versus after meal tested at 6 time intervals 0-1, 1-2, 2-3, 3-4, 4-5, and 5–8, 60% VO2peak | Exercise in fasted state is safe, no hypoglycemia; decrease in blood glucose depends on preexercise glucose level |
| Poirier et al. [9] | 10 men with T2D | Light exercise, before meal versus 2 h after meal, both at 60% VO2peak for 1 h | Moderate exercise in fasted state has minimal impact on blood glucose; exercise 2 h after meal decreases plasma glucose |
| Poirier et al. [10] | 19 men with T2D | Light exercise, before meal for 1 h versus after meal tested at 6 time intervals 0-1, 1-2, 2-3, 3-4, 4-5, and 5–8, 60% VO2peak | Exercise in fasted state does not decrease blood glucose; blood glucose decreases after postprandial exercise, no clinical hypoglycemia is observed, and in postprandial state low blood sugar is seen |
| Derave et al. [11] | 7 men with metabolic syndrome | Light exercise before meal, versus 1 h after meal, 60% VO2peak for 45 min versus no exercise | Blunted glucose response with postmeal exercise, excessive glucose response with premeal exercise, and later meals unaffected |
| Colberg et al. [12] | 12 men and women with T2D | Brisk walk, before meal versus 30 min after meal, for 20 min versus no exercise | Postdinner walking better for blunting postprandial glucose excursion and the postdinner glucose peak bigger with predinner exercise |
| DiPietro et al. [13] | 10 prediabetes men and women | Light walks, 1 h after meal for 15 min each x 3 and 2.5 h and 4.5 h after meal (before dinner) for 45 min | Postmeal walks improve 24 h glycemia, there is no 24 h glucose improvement with predinner walk, and 3 bouts of 15 min postmeal walk are more effective than 45 min of morning or afternoon walk |
| Yamanouchi et al. [14] | 6 patients with T1D | Premeal walk versus postmeal walk at 60 min after meal, 50% VO2max for 30 min versus no walk | Glucose levels and glucose-AUC significantly lower only in the postmeal walking (premeal walk 17.8, postmeal walk 3.8, and no walk 11.8 mM) |
| Francois et al. [15] | 9 individuals with insulin resistance | Premeal HIT (three 10 min bouts) versus moderate premeal exercise, 60% VO2max for 30 min | Premeal HIT exercise results in improved insulin sensitivity; moderate exercise leads to postprandial glucose elevation |
| Melton et al. [16] | 16 prediabetes women | Moderate premeal exercise, 65% HRmax for 45 min versus no exercise | No effect on glucose, triglyceride, or oxidative stress |
| Kirwan et al. [17] | 6 healthy women | Exercise before meal versus after meal (45 min), 60% VO2peak to exhaustion | As insulin and glucose go up, FFA and glycerol are suppressed for 120 min of postmeal exercise, glucose is steady with premeal exercise for 120 minutes, and duration is not altered |
| Borer et al. [18] | 9 healthy postmenopausal women | Premeal exercise, versus postmeal exercise (1 h) at 43% max effort, 2 bouts of 2 h each | Only prolonged light premeal exercise improves fasting glucose; FFA and D-3 hydroxybutyrate go up more during premeal exercise indicating liver glycogen depletion |
| Marmy-Conus et al. [19] | 6 healthy men | Moderate premeal exercise versus moderate postmeal exercise starting 30 min after meal, 71% VO2max for 60 min | Muscle glucose uptake increased, liver glucose output decreased by 62% with the postmeal exercise, and glucose level goes up 20 min into the exercise |
| Short et al. [20] | Study 2: 11 young adults | Moderate aerobic exercise 17 h before meal versus 1 h before meal, 75% VO2peak for 45 min versus no exercise | Glucose-AUC 6% lower with the 1 h premeal trial within 3 h after the exercise, the effect not seen at 17 h after exercise |
Table 1: Continued.

| Study | Subjects | Exercise protocol | Results |
|-------|----------|-------------------|---------|
| Oberlin et al. [21] | 9 sedentary patients with T2D | Moderate premeal exercise 60–75% HR_{max} for 1 h versus no exercise | Glucose-AUC improved 15% after the 2nd meal |
| Høstmark et al. [22] | 9 young and 10 middle-aged sedentary women, 10 young and 10 middle-aged trained women | Light bicycling starting 15 min after meal for 30 min versus no exercise | Light postmeal physical activity reduces blood glucose by a magnitude similar to that obtained by using drugs |
| Aadland and Høstmark [23] | 9 healthy people | Very light intensity (VLI) and light intensity (LI) walk starting 15 min after meal for 30 min versus no walking | Both VLI and LI exercise blunted and delayed the rise in blood glucose |
| Nygaard et al. [24] | 14 healthy women | Slow walking starting 15 min after meal for 15 min versus 40 min versus no walking | Even slow postmeal walking reduces postprandial glucose response to meal; this response is dose dependent on duration |
| Rasmussen et al. [25] | 7 people with T1D | Bicycling starting 15 min after meal, 65% VO_{2max} for 30 min versus no exercise | Moderate postmeal exercise starting 15 min after meal for 30 min reduces blood glucose response by one-third |
| Nelson et al. [26] | 9 people with T1D, 7 healthy people | Exercise starting 30 min after meal, 55% VO_{2max} for 45 min | Glycemic response to breakfast entirely normalized, symptomatic hypoglycemia seen after 35 min into exercise |
| Caron et al. [27] | 8 people with T1D | Exercise starting 30 min after meal, 50% VO_{2max} for 45 min | Glucose regulation improves with strategically timed postmeal exercise |
| Shin et al. [28] | 8 young healthy men | Exercise starting 30 min after meal, 50% VO_{2max} for 60 min | Higher insulin action decreases glucose, free fatty acid levels, and fat oxidation and increases growth hormone levels during exercise |
| Larsen et al. [29] | 9 sedentary men with T2D | Exercise starting 45 min after meal, 53% VO_{2max} for 45 min versus no exercise versus diet | Moderate postmeal exercise decreases glycemia, the effect does not persist to affect subsequent meal, and the effect is similar to what follows decreased calorie intake |
| Van Dijk et al. [30] | 60 T2D men (23 insulin-treated) | Endurance exercise, 2 h after meal, 35%–50% W_{max} versus 45–60 min versus no exercise | With exercise glycemic control per continuous glucose monitoring improves throughout the subsequent day. HbA{\textsubscript{1c}} is related to the magnitude of response to exercise |
| Kjaer et al. [31] | 7 men with T2D and 7 healthy men | Single bout premeal HIT exercise at 100–110% VO_{2max} | 60 min of postexercise hyperglycemia in T2D followed by increased insulin effect on glucose disposal that is present 24 h after exercise. This has less therapeutic value in T2D |
| Kreisman et al. [32] | 10 healthy men for premeal exercise and 8 healthy men for postmeal exercise | High-intensity exercise before meal versus 3 h after meal | Ra response to high-intensity exercise is preserved in postprandial exercise. Postexercise hyperglycemia is relatively reduced in postprandial exercise |
| Mitchell et al. [33] | 8 T1D and 8 healthy | High-intensity premeal exercise at 80% VO_{2max} | Postexercise hyperglycemia for 2 h, diabetes control deteriorates with intense premeal exercise |
| Yale et al. [34] | 8 lean and 12 obese people | High-intensity premeal exercise to exhaustion | Obese people had greater postexercise insulin resistance |
| Larsen et al. [35] | 8 sedentary men with T2D | High-intensity postmeal exercise, 45 min after meal, 98% VO_{2max} versus no exercise | Intense postmeal exercise does not deteriorate glucose homeostasis, effect related to energy expenditure, and the effect does not help lunch |
| Gillen et al. [36] | 7 adults with T2D | HIT 90 min after meal versus no exercise | HIT exercise at 90 min after meal reduces postprandial hyperglycemia up to 24 h |
### Table 1: Continued.

| Study                      | Subjects                  | Exercise protocol                                                                 | Results                                                                 |
|----------------------------|---------------------------|------------------------------------------------------------------------------------|------------------------------------------------------------------------|
| Little et al. [37]         | 10 inactive obese men     | HIT (high-intensity interval training) 2 h after breakfast versus continuous moderate intensity exercise, 65% VO_{2max} for 30 min versus no exercise | No effect on lunch. PPG-AUC for dinner and for next breakfast better for HIT, absolute AUC and absolute spikes not different |
| Szewieczek et al. [38]     | 14 T2D and 14 healthy     | HIT 2 h after meal versus no exercise                                              | Hyperglycemia reduced during recovery period with HIT                  |
| Heden et al. [39]          | 13 obese T2D patients     | Resistance exercise (RE) 45 min after dinner, before dinner versus no exercise      | Predinner RE improves postprandial glucose concentration; postdinner exercise improves both glucose and TAG concentrations |
| Gillen et al. [40]         | 16 women                  | HIT, fasted versus fed, starting at 60 min, 3/week for 6 weeks                      | HIT is time efficient, fed versus fasted: both improve body composition and muscle oxidative capacity |
| De Bock et al. [41]        | 20 healthy men            | Endurance training, 10 fasted versus 10 fed starting at 90 min \times 3/week for 6 weeks versus 75% VO_{peak} | Fat oxidation similar, glycogen breakdown less in fasted training        |
| van Proeyen et al. [42]    | 27 healthy men            | Endurance training, 10 fasted versus 10 fed starting at 90 min \times 4/week for 6 weeks versus 7 no training | Fasted training is (slightly) more potent in muscle adaptations         |
| Nybo et al. [43]           | 15 healthy men            | Endurance training, 7 fasted versus 8 fed starting at 3 h postprandial \times 4/week for 8 weeks at 70–85% VO_{peak} | Muscular adaptations similar in fast versus fed training except GLUT4 and glycogen content more in fasted training |
| Achten and Jeukendrup [44] | 8 healthy men             | 45 min after meal 40%, 65%, 80% VO_{2max} for 20 min                                 | Insulin peaks at 30 min after meal; insulin and glucose levels decrease in 10 min similarly (then glucose level goes up for 80%) |
| Manders et al. [45]        | 9 sedentary men with T2D  | Starting 60 min after meal light (35% W_{max}) for 60 min versus moderate intensity (70% W_{max}) exercise for 30 min | Light exercise as opposed to moderate exercise reduces hyperglycemia throughout the subsequent 24 h, prevalence of hyperglycemia 50% versus 19% |
| Van Dijk et al. [46]       | 30 patients with T2D      | 90 min after meal, 50% W_{max} 30 min every day versus 60 min every other day versus no exercise | Hyperglycemia lower in both exercise regimens                             |

that was slightly smaller than the peak that resulted from high-intensity premeal exercise, but much bigger than the postprandial glucose peak itself [32].

#### 3.5. Comparison Studies

A training study that compared the effect of high-intensity exercise in fasted versus fed state (60 min after meal) in women found comparable improvements in body composition and muscle oxidative capacity. This study did not detect any significant change in insulin sensitivity in either group [40].

Three endurance training studies compared health effects, before meal versus after meal (two at 90 min after meal and one at 3 h after meal) all in healthy men [41–43]. Fat oxidation [41] and muscle adaptation did not change but GLUT4 [43] and glycogen content went up in the fasted training.

Glucose response to exercise at different intensities was compared in two studies on healthy men and men with type 2 diabetes [44, 45]. The lower intensity groups had lower glucose levels during the exercise [44] and throughout the subsequent 24 h period [45].

One study compared duration, 30 min every day versus 60 min every other day, at 90 min after meal in people with type 2 diabetes [46]. Both experiments showed comparable decreases in glycemia.

#### 4. Discussion

Many of the 39 studies presented in Table 1 have straightforward graphical depictions of what happens to the typical glucose profiles of people with diabetes when exercise...

Scientifica
intervenes. These time versus glucose plots, viewed holistically, reveal the profound impact that timely exercise can have on blood glucose. First of all, the impact that exercise has on the glucose profile is highly dependent on the time elapsed between the start of the previous meal and the start of the exercise. Exercising at 30–45 min after meal at light to moderate intensity can reduce the postmeal glucose surge whereas exercising at a different time can increase the surge. Secondly, the effect appears to be direct and without any significant delay, as befits the source-sink relationship between exogenous glucose and physical activity. Thirdly, enough studies have accumulated to permit promising conjectures about what could be going on at the gross-physiological level, aside from the molecular processes involved. Clearly, energy balance has to be conserved: glucose molecules from the gut enter the blood stream and enrich the fuel supply and the excess thus generated is available to be used up in underwriting the physical activity rather than contributing to the surge. Which way the glycemic balance tilts depends upon the rate of arrival of meal-derived glucose in the blood and the rate at which exercise draws away the fuel. Combine this process with the proclivity of hepatic glucose to enter the picture and fill any supply gaps, and what emerges is a simple, internally consistent view of the exercise-blood glucose dynamics.

4.1. Liver Glucose. During the prebreakfast hours counterregulation is up and active and elective physical activity uses liver glucose (along with fat and muscle glycogen) [8–21, 31–34]. Hepatic glucose involvement in fueling physical activity during early [22–25] and late [13, 32] postprandial periods depends upon the intensity of the exercise and also on how much glucose is present in the blood. Rising blood glucose during exercise indicates glucose arrival in the bloodstream in amounts greater than what the body expends. Premeal exercise resulted in postprandial glucose elevation [11–15] or, at high intensities, postexertion glucose elevation [31–34]. For light exercise during the early postprandial period [22–24] the overall blunting of the peak was suboptimal. Exercise during late postprandial period resulted in glycemic response similar to what was seen in the premeal studies [13, 32, 56]. Further, in four of the midpostprandial studies a transient rise of blood glucose was seen with increased intensity or duration of the activity signifying the arrival of endogenous glucose in the blood [17–19, 44]. In short, hepatic glucose output can be minimized by tactically adjusting the timing, intensity, and duration of the elective physical activity.

4.2. Meal-Derived Glucose. Exercising during the midpostprandial period led to decreased blood glucose levels [8–14, 17–19, 26–30, 44–46, 55–59]. Light to moderate aerobic exercise at 30 min after meal gave optimal blunting of the postmeal glucose peak [12, 26–28]. When exercise started at 30 min after meal, insulin-to-glucagon ratio went up and hepatic glucose output and fat oxidation remained suppressed during the activity [26, 28]. Glucose transport to muscles is increased via insulin-mediated or contraction-mediated transport systems or both. The body utilizes the excess glucose molecules appearing in the blood before they can accumulate and form the peak.

4.3. Exercise Intensity. The body at rest relies on free fatty acids for its major fuel needs. With rising intensity blood glucose and muscle glycogen enter the mix. Glucose (muscle glycogen and blood glucose) becomes the main fuel at high intensities (>80% VO\textsubscript{2max}). Fat oxidation peaks around the intensity range of 50%–72% VO\textsubscript{2max} [60, 61]. At high-intensity exercise fat oxidation drops precipitously, catecholamine levels rise as much as 15-fold, and hepatic glucose output rises up to 8-fold [54, 62]. The extent of hepatic glucose production is determined by insulin-to-glucagon ratio during light to moderate exercise. Catecholamines play a comparable role when the activity is very intense. Improved insulin sensitivity observed with high-intensity exercise is brought about by the glycogen repletion that follows the glycogen depletion [15, 31, 35–38, 47–51, 54, 62, 63]. High-intensity exercise has mixed effects for diabetes patients: improved body composition, physical fitness, GLUT4 protein expression, mitochondrial function, oxygen capacity, and, to some extent, insulin sensitivity [15, 31, 47–52, 63]. Although the magnitude is modest the improvement in insulin sensitivity following high-intensity exercise is consistently observed [15, 31, 47–51, 53, 63]. Once the exercise training is stopped, the insulin sensitivity benefit disappears in about 60 hours [64]. In short, the insulin sensitivity improvement following high-intensity exercise is of slow onset, short-lived, and not nearly enough to adequately tame the glucose surges to come.

Markedly increased hepatic glucose output, on the other hand, resulted in a postexertion glucose elevation usually seen with intense premeal exercise [31–34]. Improved insulin sensitivity following muscle glycogen depletion can also lead to nocturnal hypoglycemia in people with type 1 diabetes [6, 47, 51]. The net effect of high-intensity premeal exercise on glycemia can be major glucose swings.

4.4. High-Intensity Interval Training. High-intensity interval training efficiently improves body composition, physical fitness [40, 47–52], and insulin sensitivity [47–51]. But it is not clear whether high-intensity interval training is appropriate for diabetes patients. Apart from the basic question of physical feasibility on the part of diabetes patients, exercise with intensity >80% VO\textsubscript{2max} comes with some negative effects: negligible fat oxidation, elevated catecholamines, and increased hepatic glucose output [54, 62, 63]. None of these is diabetes friendly, although fat oxidation is observed to go up following intense activity [53]. The high levels of catecholamines are not salutary to a population that can ill afford high blood pressure too. The worst effect of all is wide glucose swings: from postexertion glucose elevation [31–34] to nocturnal hypoglycemia [6, 47, 51].

High-intensity interval training has received much attention lately as an efficient way to deal with glucose surges [15, 47, 49]. According to the review by Kessler and colleagues high-intensity interval training for 8–12 months showed improvement in some cardiometabolic risk factors, but the reported effects on fasting glucose, HbA\textsubscript{1C}, and HDL cholesterol were inconsistent [48]. Lipids in general did not
improve. Moreover, two studies found no improvement in HbA1C in patients with type 2 and type 1 diabetes with high-intensity interval training [65, 66].

4.5. Energy Expenditure. Of the 5 studies at 30 min after meal that showed efficient blunting of the glucose peak, one was a brisk walk for 20 minutes [12]; two were at 55% VO2max for 45 min [26, 27]. The other two studies had durations of 60 min each and intensities of 50% VO2max and 71% VO2max, respectively [19, 28]. It looked like the effectiveness of the exercise depended upon intensity and duration which together determine energy expenditure. Two studies, one at 45 min after meal [44] and the other at 60 min after meal [45], compared glucose responses to exercise at different intensities. In both cases the result showed that the lower intensity produced better glucose control. At relatively high intensities, glucose from endogenous sources rushing to meet the increased demand can lead to elevated glucose levels. For example, when the intensity was 50% VO2max, glucose level did not rise during the 60 min duration of the exercise [28]. On the other hand, when the intensity rose to 71% VO2max glucose level went up 20 min into the activity [19].

The best tactic for taming glucose surges is to go with a light aerobic activity (<60% VO2max) at 30 min after meal for up to 60 min [28]. At higher intensities duration may be reduced to 20–30 min [19]. Patients can fine-tune the intensity and duration guided by glucose monitoring. Those who do not have a glucometer can be trained with relative intensity and duration guided by glucose monitoring. Those who do not have a glucometer can be trained with relative intensity and duration guided by glucose monitoring.

4.6. Resistance Plus Aerobic Exercise. Three studies demonstrated the role of postmeal resistance exercise in lowering glucose surges [7, 39, 67]. When Heden and colleagues used moderate intensity resistance exercise for 45 min starting at 45 min after meal, glucose levels came down during the exercise partially blunting the surge in obese patients with type 2 diabetes [39]. Another study showed that an acute bout of moderate resistance exercise 2.5 hours after meal prevalence of hyperglycemia was reduced by 35% over the next 24 hours in people with impaired glucose tolerance and in those with type 2 diabetes with or without insulin [67]. Yet another study used combined resistance and endurance exercise of moderate intensity during the postmeal surge to get hyperglycemia reduced by 39% during the 24 hours following the exercise bout [7]. Although combined training is better for glucose control (HbA1C) than resistance training or endurance training alone the data on these when the training was conducted at postmeal period are not available [68–73]. Resistance training 3 times a week [74] at midpostprandial period may have added benefits in controlling glucose surges.

4.7. Hypoglycemia. Transient, inconsequential hypoglycemia could strike when exercising in the late postprandial period [1, 7] and delayed nocturnal hypoglycemia [6, 47, 51] is a distinct possibility after intense premeal exercise in people with type 1 diabetes. Nelson and colleagues observed no hypoglycemia during the first 35 minutes of light exercise in both healthy people and people with type 1 diabetes when the activity started at 30 min after meal [26]. Praet and colleagues noted mild hypoglycemia, glucose concentration below 3.9 mmol/L, in 6 out of 11 subjects when the exercise was done during the late postprandial period [7]. Most diabetes patients can blunt the glucose peak and simultaneously minimize the risk of hypoglycemia by starting a bout of light to moderate physical activity at 30 minutes after meal. In the midpostprandial period the glucose still coming in from the gut will guard against hypoglycemia. Patients with diabetes can further fine-tune the intensity and duration with the help of glucometer readings.

5. Strengths and Limitations

Out of the 39 studies included in Table 1, 17 had sample sizes numbering in the single digits. Another limitation is the nonuniform categorizations used for exercise intensities in different studies included in this review.

In spite of the small sample sizes, a wide spectrum of metabolic conditions could be seen among the 615 subjects in the 39 studies. The actual numbers were 230 (type 2), 50 (type 1), 70 (obesity, prediabetes, metabolic syndrome, or insulin resistance), and 265 (healthy.) Twenty-one studies used control groups, whereas the others used a crossover design. All 39 studies used supervised exercise and all but two reported using standardized diets. The outcome measures in general could be seen across multiple studies, yielding consistent and reproducible results. Also on the positive side, the results of this review are the exposition of a few systematic problems bedeviling diabetes research. These are identified immediately below.

6. Looking Ahead

A valuable next step would be a lifestyle study to measure the effect on metabolic parameters of a timely postmeal exercise of light to moderate intensity starting 30 min after major meals. Also, it is useful to see the effect of adding a 10 min resistance exercise to the above lifestyle before the endurance exercise [68–72], 2-3 times a week [73].

Exercise timing is not specified in many studies and this makes it quite difficult to make sense of the results. Timing, intensity, frequency, and duration of the physical activity are needed to assess the full glucose response. Also, many long-term studies involving high-intensity interval training discuss improvements in insulin sensitivity without reporting the values of HbA1C [47–51].

AUC calculations are not properly standardized. Incremental AUC, total AUC, and positive AUC as well as other arbitrarily defined AUC calculations have been reported, and it is difficult to compare the results. The abundance of misguided creativity seen here appears to be born of the notion that calculations can be used in such a manner that "less pronounced differences" between observed outcomes "would be easier to detect" [75]. Calculations are not microscopes, and calculation-dependent effects are not scientifically trustworthy. Sticking with the total AUC is the only legitimate option here.
7. Summary and Conclusions

Light to moderate exercise during the midpostprandial period preferentially uses meal-derived glucose and lowers blood glucose levels because insulin and glucose levels are high and, therefore, hepatic glucose output and fat oxidation are inhibited. To achieve the critical goal of blunting the postmeal glucose peak, it is best to start the aerobic exercise [<80% VO₂max] at 30 min after meal and continue for up to 60 min. Hypoglycemia risk is minimal with this approach. Exercising at any other time, before meal, early after meal, and late after meal, can trigger endogenous glucose production, resulting in elevated blood glucose levels. Although high-intensity (>80% VO₂max) exercise is effective in improving body composition, physical fitness, and insulin sensitivity it's feasibility and efficacy in improving cardiometabolic markers for diabetes patients need to be established. This is because high-intensity exercise can offer wide fluctuations in glucose excursion: postexercise glucose elevation and delayed nocturnal hypoglycemia. There are indications that appending a bout of resistance training at moderate intensity (60%–80% VO₂max) to the daily endurance exercise 2-3 times a week is appropriate, if the patient is fit enough to undertake it.

We have come a long way with diabetes management, but the progress has not touched the majority of patients who live in developing countries and in the pockets of poverty in the West. Medications, medical care, glucometers, and choice diets are simply not accessible to the vast majority of insulin-treated diabetic patients who live in these parts of the world. They must fight the disease with physical activity and little else. Promoting a light to moderate, timely postmeal exercise after major meals would be of tremendous benefit universally to all with insulin resistance. Such a lifestyle practice would not violate any current guidelines: guidelines encourage exercise any time.

Competing Interests

The author declares that there is no duality of interests associated with this paper.

References

[1] J. Haxhi, A. Scotto Di Palumbo, and M. Sacchetti, “Exercising for metabolic control: is timing important?” Annals of Nutrition and Metabolism, vol. 62, no. 1, pp. 14–25, 2013.

[2] E. Chacko, “Timing and intensity of exercise for glucose control,” Diabetologia, vol. 57, no. 11, pp. 2425–2426, 2014.

[3] S. Daenen, A. Sola-Gazagnes, J. M’Bembé et al., “Peak-time determination of post-meal glucose excursions in insulin-treated diabetic patients,” Diabetes and Metabolism, vol. 36, no. 2, pp. 165–169, 2010.

[4] P. E. Cryer, S. N. Davis, and H. Shamoon, “Hypoglycemia in diabetes,” Diabetes Care, vol. 26, no. 6, pp. 1902–1912, 2003.

[5] F. Cavallet, A. Petrelli, M. Traversa et al., “Postprandial blood glucose is a stronger predictor of cardiovascular events than fasting blood glucose in type 2 diabetes Mellitus, particularly in women: lessons from the San Luigi Gonzaga Diabetes Study,” Journal of Clinical Endocrinology and Metabolism, vol. 91, no. 3, pp. 813–819, 2005.

[6] A. Maran, P. Pavan, B. Bonsembiante et al., “Continuous glucose monitoring reveals delayed nocturnal hypoglycemia after intermittent high-intensity exercise in nontrained patients with type 1 diabetes,” Diabetes Technology and Therapeutics, vol. 12, no. 10, pp. 763–768, 2010.

[7] S. F. Praet, R. J. Manders, A. G. Lieverse et al., “Influence of acute exercise on hyperglycemia in insulin-treated type 2 diabetes,” Medicine and Science in Sports and Exercise, vol. 38, no. 12, pp. 2037–2044, 2006.

[8] T. Gaudet-Savard, A. Ferland, T. L. Broderick et al., “Safety and magnitude of changes in blood glucose levels following exercise performed in the fasted state and the postprandial state in men with type 2 diabetes,” European Journal of Cardiovascular Prevention and Rehabilitation, vol. 14, no. 6, pp. 831–836, 2007.

[9] P. Poirier, S. Mawhinney, L. Grondin et al., “Prior meal enhances the plasma glucose lowering effect of exercise in type 2 diabetes,” Medicine and Science in Sports and Exercise, vol. 33, no. 8, pp. 1259–1264, 2001.

[10] P. Poirier, A. Tremblay, C. Catellier, G. Tancrède, C. Garneau, and A. Nadeau, “Impact of time interval from the last meal on glucose response to exercise in subjects with Type 2 diabetes,” Journal of Clinical Endocrinology and Metabolism, vol. 85, no. 8, pp. 2860–2864, 2000.

[11] W. Derave, A. Mertens, E. Muls, K. Pardaens, and P. Hespel, “Effects of post-absorptive and postprandial exercise on glucosecoregulation in metabolic syndrome,” Obesity, vol. 15, no. 3, pp. 704–711, 2007.

[12] S. R. Colberg, L. Zarrabi, L. Bennington et al., “Postprandial walking is better for lowering the glycemic effect of dinner than pre-dinner exercise in type 2 diabetic individuals,” Journal of the American Medical Directors Association, vol. 10, no. 6, pp. 394–397, 2009.

[13] L. DiPietro, A. Gribok, M. S. Stevens, L. F. Hamm, and W. Rumpler, “Three 15-min bouts of moderate postmeal walking significantly improves 24-h glycemic control in older people at risk for impaired glucose tolerance,” Diabetes Care, vol. 36, no. 10, pp. 3262–3268, 2013.

[14] K. Yamanouchi, R. Abe, A. Takeda, Y. Atsumi, M. Shichiri, and Y. Sato, “The effect of walking before and after breakfast on blood glucose levels in patients with type 1 diabetes treated with intensive insulin therapy,” Diabetes Research and Clinical Practice, vol. 58, no. 1, pp. 11–18, 2002.

[15] M. E. Francois, J. C. Baldi, P. J. Manning et al., “‘Exercise snacks’ before meals: a novel strategy to improve glycaemic control in individuals with insulin resistance,” Diabetologia, vol. 57, no. 7, pp. 1437–1445, 2014.

[16] C. E. Melton, P. S. Tucker, K. H. Fisher-Wellman, B. K. Schilling, and R. J. Bloomer, “Acute exercise does not attenuate postprandial oxidative stress in prediabetic women,” Physician and Sportsmedicine, vol. 37, no. 1, pp. 27–36, 2009.

[17] J. P. Kirwan, D. J. O’Gorman, D. Cyr-Campbell, W. W. Campbell, K. E. Yarasheski, and W. J. Evans, “Effects of a moderate glycemic meal on exercise duration and substrate utilization,” Medicine and Science in Sports and Exercise, vol. 33, no. 9, pp. 1517–1523, 2001.

[18] K. T. Borer, E. C. Wuorinen, J. R. Lukos, J. W. Denver, S. W. Porges, and C. F. Burant, “Two bouts of exercise before meals, but not after meals, lower fasting blood glucose,” Medicine and Science in Sports and Exercise, vol. 41, no. 8, pp. 1606–1614, 2009.

[19] N. Marmy-Conus, S. Fabris, J. Froitto, and M. Hargreaves, “Preexercise glucose ingestion and glucose kinetics during...
exercise," *Journal of Applied Physiology*, vol. 81, no. 2, pp. 853–857, 1996.

[20] K. R. Short, L. V. Pratt, and A. M. Teague, "The acute and residual effect of a single exercise session on meal glucose tolerance in sedentary young adults," *Journal of Nutrition and Metabolism*, vol. 2012, Article ID 278678, 9 pages, 2012.

[21] D. J. Oberlin, C. R. Mikus, M. L. Kearney et al., "One bout of exercise alters free-living postprandial glycaemia in type 2 diabetes," *Medicine and Science in Sports and Exercise*, vol. 46, no. 2, pp. 232–238, 2014.

[22] A. T. Høstmark, G. S. Ekeland, A. C. Beckstrøm, and H. D. Meen, "Postprandial light physical activity blunts the blood glucose increase," *Preventive Medicine*, vol. 42, no. 5, pp. 369–371, 2006.

[23] E. Aadland and A. T. Høstmark, "Very light physical activity after a meal blunts the rise in blood glucose and insulin," *The Open Nutrition Journal*, vol. 2, pp. 94–99, 2008.

[24] H. Nygaard, S. E. Tomten, and A. T. Høstmark, "Slow post-meal walking reduces postprandial glycaemia in middle-aged women," *Applied Physiology, Nutrition and Metabolism*, vol. 34, no. 6, pp. 1087–1092, 2009.

[25] O. W. Rasmussen, F. F. Lauszus, and K. Hermansen, "Effects of moderate exercise on postprandial glucose homeostasis in Type II diabetic patients," *Diabetologia*, vol. 42, no. 11, pp. 1282–1292, 1999.

[26] J. B. Gillen, J. P. Little, Z. Punthakee, M. A. Tarnopolsky, M. C. Riddell, and M. J. Gibala, "Acute high-intensity interval exercise reduces the postprandial glucose response and prevalence of hyperglycaemia in patients with type 2 diabetes," *Diabetes, Obesity and Metabolism*, vol. 14, no. 6, pp. 575–577, 2012.

[27] J. P. Little, M. E. Jung, A. E. Wright, W. Wright, and R. J. F. Manders, "Effects of high-intensity interval exercise versus continuous moderate-intensity exercise on postprandial glycemic control assessed by continuous glucose monitoring in obese adults," *Applied Physiology, Nutrition and Metabolism*, vol. 39, no. 7, pp. 835–841, 2014.

[28] J. Szewieczek, J. Dulawa, D. Strzałkowska, B. Hornik, and G. Kawecki, "Impact of the short-term, intense exercise on postprandial glycaemia in type 2 diabetic patients treated with gliclazide," *Journal of Diabetes and its Complications*, vol. 21, no. 1, pp. 101–107, 2007.

[29] T. D. Heden, N. C. Winn, A. Mari et al., "Postdinner resistance exercise improves postprandial risk factors more effectively than predinner resistance exercise in patients with type 2 diabetes," *Journal of Applied Physiology*, vol. 118, no. 5, pp. 624–634, 2015.

[30] J. B. Gillen, M. E. Percival, A. Ludzki, M. A. Tarnopolsky, and M. J. Gibala, "Interval training in the fed or fasted state improves body composition and muscle oxidative capacity in overweight women," *Obesity*, vol. 21, no. 11, pp. 2249–2255, 2013.

[31] K. De Bock, W. Derave, B. O. Eijnde et al., "Effect of training in the fasted state on metabolic responses during exercise with carbohydrate intake," *Journal of Applied Physiology*, vol. 104, no. 4, pp. 1045–1055, 2008.

[32] K. van Proeyen, K. Szlufcik, H. Nielsens et al., "Training in the fasted state improves glucose tolerance during fat-rich diet," *Journal of Physiology*, vol. 588, no. 21, pp. 4289–4302, 2010.

[33] L. Nybo, K. Pedersen, B. Christensen, P. Aagaard, N. Brandt, and B. Kientz, "Impact of carbohydrate supplementation during endurance training on glycogen storage and performance," *Acta Physiologica*, vol. 197, no. 2, pp. 117–127, 2009.

[34] J. Achten and A. E. Jeukendrup, "Effects of pre-exercise ingestion of carbohydrate on glycaemic and insulinaemic responses during subsequent exercise at differing intensities," *European Journal of Applied Physiology*, vol. 88, no. 4-5, pp. 466–471, 2003.

[35] R. J. F. Manders, J.-W. M. Van Dijk, and L. J. C. van Loon, "Low-intensity exercise reduces the prevalence of hyperglycaemia in type 2 diabetes," *Medicine and Science in Sports and Exercise*, vol. 42, no. 2, pp. 219–225, 2010.

[36] J.-W. Van Dijk, K. Timmers, C. D. A. Stehouwer, F. Hartgens, and L. J. C. van Loon, "Exercise therapy in type 2 diabetes: is daily exercise required to optimize glycemic control?" *Diabetes Care*, vol. 35, no. 5, pp. 948–954, 2012.

[37] O. P. Adams, "The impact of brief high-intensity exercise on blood glucose levels," *Diabetes, Metabolic Syndrome and Obesity: Targets and Therapy*, vol. 6, pp. 113–122, 2013.

[38] H. S. Kessler, S. B. Sisson, and K. R. Short, "The potential for high-intensity interval training to reduce cardiometabolic disease risk," *Sports Medicine*, vol. 42, no. 6, pp. 489–509, 2012.

[39] C. K. Roberts, J. P. Little, and J. P. Thiyault, "Modification of insulin sensitivity and glycemic control by activity and exercise," *Medicine and Science in Sports and Exercise*, vol. 45, no. 10, pp. 1868–1877, 2013.
diabetes: a meta-analysis of studies using continuous glucose monitoring,” *Diabetes/Metabolism Research and Reviews*, vol. 29, no. 8, pp. 593–603, 2013.

[51] J. Yardley, R. Mollard, A. MacIntosh et al., “Vigorous intensity exercise for glycemic control in patients with type 1 diabetes,” *Canadian Journal of Diabetes*, vol. 37, no. 6, pp. 427–432, 2013.

[52] M. J. Gibala and A. M. Jones, “Physiological and performance adaptations to high-intensity interval training,” *Nestle Nutrition Institute Workshop Series*, vol. 76, pp. 51–60, 2013.

[53] R. W. Asano, M. M. Sales, R. A. V. Browne et al., “Acute effects of physical exercise in type 2 diabetes: a review,” *World Journal of Diabetes*, vol. 5, no. 5, pp. 659–665, 2014.

[54] E. B. Marliss and M. Vranic, “Intense exercise has unique effects on both insulin release and its roles in glucoregulation: implications for diabetes,” *Diabetes*, vol. 51, supplement 1, pp. S271–S283, 2002.

[55] C. Manohar, J. A. Levine, D. K. Nandy et al., “The effect of walking on postprandial glycemic excursion in patients with type 1 diabetes and healthy people,” *Diabetes Care*, vol. 35, no. 12, pp. 2493–2499, 2012.

[56] S. Sasaki, S. Nakae, N. Ebine, W. Aoi, A. Higashi, and K. Ishii, “The effect of the timing of meal intake on energy metabolism during moderate exercise,” *Journal of Nutritional Science and Vitaminology*, vol. 60, no. 1, pp. 28–34, 2014.

[57] T. Terada, A. Friessen, B. S. Chahal, G. J. Bell, L. J. McCargar, and N. G. Boulé, “Exploring the variability in acute glycemic responses to exercise in type 2 diabetes,” *Journal of Diabetes Research*, vol. 2013, Article ID 591574, 6 pages, 2013.

[58] S. R. Colberg, M. J. Hernandez, and F. Shahzad, “Blood glucose responses to type, intensity, duration, and timing of exercise,” *Diabetes Care*, vol. 36, no. 10, article e177, 2013.

[59] L. W. L. Tobin, B. Kiens, and H. Galbo, “The effect of exercise on postprandial lipemia in type 2 diabetic patients,” *European Journal of Applied Physiology*, vol. 102, no. 3, pp. 361–370, 2008.

[60] J. Achten, M. Gleeson, and A. E. Jeukendrup, “Determination of exercise intensity that elicits maximal fat oxidation,” *Medicine & Science in Sports & Exercise*, vol. 34, no. 1, pp. 92–97, 2002.

[61] P. D. Feo, C. D. Loreto, P. Lucidi et al., “Metabolic response to exercise,” *Journal of Endocrinological Investigation*, vol. 26, no. 9, pp. 851–854, 2003.

[62] M. Kjaer, “Hepatic glucose production during exercise; in *Skeletal Muscle Metabolism in Exercise and Diabetes*, E. A. Richter, B. Kiens, H. Galbo, and B. Saltin, Eds., vol. 441 of *Advances in Experimental Medicine and Biology*, pp. 117–127, Springer, New York, 1998.

[63] J. Jensen, P. I. Rustad, A. J. Kolnes, and Y.-C. Lai, “The role of skeletal muscle glycogen breakdown for regulation of insulin sensitivity by exercise,” *Frontiers in Physiology*, vol. 2, article 112, 2011.

[64] R. Burstein, C. Polychronakos, C. J. Toews, J. D. MacDougall, H. J. Guyda, and B. I. Posner, “Acute reversal of the enhanced insulin action in trained athletes. Association with insulin receptor changes,” *Diabetes*, vol. 34, no. 8, pp. 756–760, 1985.

[65] T. Terada, A. Friessen, B. S. Chahal, G. J. Bell, L. J. McCargar, and N. G. Boulé, “Feasibility and preliminary efficacy of high intensity interval training in type 2 diabetes,” *Diabetes Research and Clinical Practice*, vol. 99, no. 2, pp. 120–129, 2013.

[66] A. R. Harmer, D. J. Chisholm, M. J. McKenna et al., “High-intensity training improves plasma glucose and acid-base regulation during intermittent maximal exercise in type 1 diabetes,” *Diabetes Care*, vol. 30, no. 5, pp. 1269–1271, 2007.