Effects of Diet, Lifestyle, Chrononutrition and Alternative Dietary Interventions on Postprandial Glycemia and Insulin Resistance

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Abstract: As years progress, we are found more often in a postprandial than a postabsorptive state. Chrononutrition is an integral part of metabolism, pancreatic function, and hormone secretion. Eating most calories and carbohydrates at lunch time and early afternoon, avoiding late evening dinner, and keeping consistent number of daily meals and relative times of eating occasions seem to play a pivotal role for postprandial glycemia and insulin sensitivity. Sequence of meals and nutrients also play a significant role, as foods of low density such as vegetables, salads, or soups consumed first, followed by protein and then by starchy foods lead to ameliorated glycemic and insulin responses. There are several dietary schemes available, such as intermittent fasting regimes, which may improve glycemic and insulin responses. Weight loss is important for the treatment of insulin resistance, and it can be achieved by many approaches, such as low-fat, low-carbohydrate, Mediterranean-style diets, etc. Lifestyle interventions with small weight loss (7–10%), 150 min of weekly moderate intensity exercise and behavioral therapy approach can be highly effective in preventing and treating type 2 diabetes. Similarly, decreasing carbohydrates in meals also improves significantly glycemic and insulin responses, but the extent of this reduction should be individualized, patient-centered, and monitored. Alternative foods or ingredients, such as vinegar, yogurt, whey protein, peanuts and tree nuts should also be considered in ameliorating postprandial hyperglycemia and insulin resistance. This review aims to describe the available evidence about the effects of diet, chrononutrition, alternative dietary interventions and exercise on postprandial glycemia and insulin resistance.

Keywords: postprandial hyperglycemia; insulin resistance; insulin secretion; chrononutrition; dietary interventions; exercise

1. Introduction

“Let food be thy medicine, and let medicine be thy food”, the famous quote by Greek physician Hippocrates stands well the test of time. Diet is considered the cornerstone of prevention and treatment of glucose dysmetabolism and insulin resistance (IR), and food has major effects on postprandial glycemia and overall physical health [1]. Our modern lives are characterized by being more often in a postprandial state, decreased energy expenditure, exposure to a “toxic” food environment, sedentary lifestyle with prolonged sitting time, high consumption of energy dense foods, irregular eating occasions and times of eating, skipping meals, chronic psychological stress, emotional eating, and food consumption late at night [1]. This way of life triggers mechanisms, such as development of insulin resistance (IR), proposed as a defense system against metabolic stress, especially for the heart [2].
Factors associated with IR include obesity, particularly abdominal obesity, increased waist circumference, familial history of type 2 diabetes (T2DM), sedentary lifestyle, hypertension, and fatty liver [3–6] (Figure 1). Obesity pathogenesis involves two related but distinct processes: a sustained positive energy balance and resetting of the body’s weight “set point” at an increased level [7]. Obesity is a heterogeneous phenotype and not every obese person is at risk for developing metabolic abnormalities; therefore, one needs to focus on the individual [8,9]. It has been reported that although there is a causal role of hyperinsulinemia in the progression of obesity with a well-established connection, its contribution varies [9]. The amount of visceral adipose tissue, estimated by increased waist circumference, may reflect the severity of IR [10]. IR is also characterized by hypertriglyceridemia, lower concentrations of high-density lipoprotein (HDL) cholesterol in blood, and increased inflammation [11]. As a result, glucose metabolism deteriorates, resulting in hyperglycemia, impaired glucose tolerance (IGT) or overt T2DM. Weight loss and regular moderate intensity physical activity/exercise are significant factors for IR prevention and/or treatment [12]. From a dietary perspective, dietary fiber, cereal fiber, fruit fiber, whole grains, full-fat dairy products [13–16], magnesium, and calcium lower IR, whereas high glycemic index (GI) and glycemic load (GL) foods, saturated fat, salt (deficiency or excess), and alcohol (>30 g/day) increase IR [17].

T2DM development stems from various genetic and/or environmental factors and is characterized by deficient pancreatic β-cell insulin secretion and decreased sensitivity/responsiveness of insulin-sensitive tissues to insulin [18]. The prevailing view in the physical history of T2DM is that IR precedes causing a progressive increase in insulin secretion to compensate for IR and maintain glucose tolerance (IGT-prediabetes). In the long term β-cell function declines, and hyperglycemia follows (overt T2DM) [19]. However, in some individuals, hyperinsulinemia may not be a compensatory response to insulin resistance but rather a primary defect due to hypersensitivity of β-cells explained by genetic/environmental factors, gastrointestinal/neural signals, or various substrates such as lactate, non-esterified fatty acids (NEFA)/triglycerides, or amino acids [20–23].

Uncontrolled T2DM may lead to serious consequences on health and health care cost [24,25] and its prevalence is high with over 427 million adults being affected worldwide expected to reach 629 million people by 2045 [25–27]. T2DM is related to micro- and macrovascular chronic complications, such as retinopathy, neuropathy, chronic kidney disease, cardiovascular disease (CVD), and non-alcoholic fatty liver disease (NAFLD) [25,27–29].

The current COVID-19 pandemic has led to dramatic societal changes leading to changes in consumers’ food practices with fewer cooking-related practices, bulk buying, increased fruit, vegetables and saturated fat intake, and increased body weight [30], all of which may have a significant effect on postprandial glycemia and IR.
Mechanisms for the Regulation of Postprandial Hyperglycemia

During meal ingestion, several mechanisms operate in concert (gastric emptying/intestinal glucose absorption, secretion/action of gastrointestinal hormones, changes in insulin/glucagon secretion/action, hyperglycemia mass action) to ensure optimal regulation of postprandial glucose fluctuations in the bloodstream via coordination by the central nervous system (CNS) (reviewed in detail in reference [31]): (a) After a meal, insulin plays a primary role in the regulation of glucose homeostasis through its effect on insulin-sensitive tissues (liver/skeletal muscle/adipose tissue) [32]. In the beginning of meal ingestion, there is a rapid increase in the secretion of insulin and a parallel decrease in that of glucagon from pancreatic β- and α-cells, respectively. These hormonal changes, rapidly inhibit hepatic glucose production (HGP) to allow incorporation of more than 30% of the ingested glucose into liver glycogen (direct pathway of glycogen synthesis) [33,34]. The preferential use of the liver to dispose a significant amount of the ingested glucose during first pass, along with a 60–70% hepatic clearance of insulin, protect the peripheral circulation from excessive hyperglycemia/hyperinsulinemia in the postprandial period [35–43]. The entrance of the remaining insulin into the peripheral circulation rapidly suppresses lipolysis in the adipose tissue and reduces plasma levels of NEFA; this facilitates the decrease of HGP/increase of glucose storage by insulin and permits the increase of insulin-stimulated glucose uptake by skeletal muscle for oxidation and storage as glycogen [32,44]. However, if the liver is overpowered by increased amounts of carbohydrate in the meal, the escape of glucose to the peripheral circulation will be much higher engaging skeletal muscle for its removal; this will require a substantial increase in insulin secretion and plasma insulin levels which, in the long term, may lead to the development/aggravation of IR [45–47]. If circulating glucose exceeds the capacity of skeletal muscle for oxidation/storage, muscle cells convert glucose to lactate/alanine/glutamine which are then delivered back to the liver and incorporate their carbons to glycogen (indirect pathway of glycogen synthesis) [34,48,49]. Considering both the direct (first pass) and indirect pathways (through muscle) of glycogen synthesis, overall, the liver disposes more than 50% of the ingested glucose and hence allows a much smaller amount to remain in the peripheral circulation [50]. Therefore, as suggested by Kowalski et al. [51] the liver may represent an “evolutionary conserved mechanism” for the regulation of postprandial hyperglycemia with a mission to reduce the secretory burden on β-cells, thus preventing plasma glucose and insulin levels from rising too high and induce serious consequences [37,38,40,41,43]. Insulin also affects the vascular endothelium and increases the rates of blood flow in skeletal muscle and adipose tissue. After meals, the increase of blood flow in muscle facilitates the delivery of substrates/hormones for metabolism, whereas in adipose tissue it ensures the clearance of NEFA from the circulation [52,53]. In conditions such as IGT, T2DM or obesity, insulin-stimulated rates of blood flow in adipose tissue and muscle are severely impaired [54–56].

(b) The role of insulin sensitivity in postprandial glucose regulation is important to restrain hyperinsulinemia after meals. The amount of insulin release from the β-cells under all circumstances depends on the metabolic requirements of the insulin-sensitive tissues. Therefore, if the sensitivity of these tissues to insulin is increased, the management of an increased entry of glucose into the bloodstream after meals could be achieved in the absence of marked increases in the plasma levels of insulin [57]. Indeed, Kahn et al. [58] provided experimental evidence showing that the pancreatic β-cells and insulin-sensitive tissues interact in a tightly regulated manner so that when insulin sensitivity is high, insulin secretion is low and vice versa. Therefore, lifestyle interventions, such as those described in the present review, are important to improve tissue sensitivity to insulin and hence avoid marked hyperglycemia and hyperinsulinemia after meals. (c) Glucose can regulate its own metabolism in tissues independently of insulin [59]. In the liver, hyperglycemia per se increases the rates of glucose uptake and suppresses HGP, thus facilitating glucose storage [60], whereas in muscle it increases the rates of glucose uptake and phosphorylation [61]. Therefore, as suggested by Kowalski et al. [51], the subtle increase in plasma glucose levels seen after meals in physiological conditions, may serve to collaborate with
insulin to minimize the amount of insulin required for its effects on insulin-sensitive tissues and, hence, reduce the secretory burden on \( \beta \)-cells. (d) The gastrointestinal tract plays a significant role in the regulation of hyperglycemia during food ingestion via feedback mechanisms coordinated by the CNS [62]. The rate of gastric emptying is a major determinant of the glycemic/insulin responses following a meal and has been shown to correlate positively with postprandial hyperglycemia [63,64]. Furthermore, in response to a meal the intestinal tract secretes several peptides (incretins) that regulate its motility/absorption capacity, and insulin/glucagon secretion and action [65,66]. Glucagon-like peptide-1 (GLP-1) and glucose-dependent insulino-tropic polypeptide (GIP) have been heavily investigated due to their clinical application in T2DM; they collaborate to increase glucose-dependent insulin secretion and insulin sensitivity [67,68]. The coordinated interplay between the gastrointestinal tract and pancreatic \( \beta \) - and \( \alpha \)-cells ensures a finely tuned adjustment of glucose supply to the metabolic needs and, therefore, helps to avoid high glucose/insulin fluctuations after meals. (e) Meal sequence within the day ensures that preceding meals sensitize the metabolic/incretin system to the following ones, thereby improving glucose tolerance during the day mostly by increasing glucose storage in the liver but also by facilitating glucose uptake in skeletal muscle [69–73] (discussed in detail in Section 2.3). Finally, meal composition, and ingestion of low GI/GL foods, particularly when consumed along with protein and/or fat, reduce postprandial glucose and insulin responses, thus improving insulin sensitivity [45,46,74–79] (discussed further in Section 4.2.1).

2. Effects of Macronutrients in Foods and Meals on Postprandial Hyperglycemia and IR

The glycemic response depends on many factors including the amount of total consumed carbohydrates, carbohydrate type, starch type (i.e., resistant starch), food preparation (i.e., gelatinization, bran size), other macronutrients in foods (i.e., fat, protein, fiber), and the physiological organic functions (stomach, pancreatic and enteral hydrolysis, gastric emptying, intestinal absorption rate of nutrients, etc.) [78] (Figure 2). Regarding dietary components, a key dietary strategy for treating postprandial hyperglycemia, hyperinsulinemia, and IR is the consumption of foods and meals that diminish the glucose fluctuations known to induce oxidative stress and \( \beta \)-cell damage [80]. Indeed, increased glucose variability from peaks to nadirs has been recognized as a major metabolic defect leading to endothelial damage and CVD in T2DM [81]. The eating pace of consuming a meal has also been implicated in modulating postprandial hyperglycemia; for example, eating fast has been associated with higher glycemic excursions in healthy women [82]. Moreover, another important strategy to consider is the consumption of foods and meals that induce a lower GL and delay gastric emptying, thus leading to decreased insulin requirements and postprandial glucose excursions and may also reduce hunger and desire to eat [31]. Such foods and meals typically contain high fiber, particularly soluble fiber, low amounts of easily absorbable carbohydrates, low amounts of total carbohydrates, and are high in proteins [31,46,75].

Carbohydrates are well known to be the major macronutrients affecting postprandial glycemia. A study using continuous glucose monitoring in subjects with T2DM showed that for achieving lower 24-h glucose peaks it was better to consume at least half of the amount of carbohydrates at lunch time and avoid consuming significant amount of carbohydrates in the morning (at breakfast) or in the evening dinner [83]. A cross-sectional study examining the effects of different proportions of carbohydrates at breakfast on postprandial glucose fluctuations in IGT (\( N = 55 \)) and normal glucose tolerance (NGT) individuals (\( N = 78 \)), using continuous glucose monitoring, recorded breakfast meals according to the proportion of carbohydrates into low (<45%), medium (45% to 65%) and high carbohydrates (>65%), and reported a gradual increase in postprandial glucose fluctuations with increasing proportions of carbohydrates in breakfast, higher postprandial glucose excursions, higher postprandial glucose spikes, and longer time period in which glucose levels decreased to baseline in subjects with IGT compared to NGT individuals; this study concluded that in IGT a high-
carbohydrate meal at breakfast should be avoided and a low carbohydrate meal should be recommended instead [84]. Carbohydrate-rich foods that may be more effective in ameliorating postprandial hyperglycemia and IR include legumes/pulses [85–87], whole grains [88–91], and pasta [92]. In 2017, the American Diabetes Association (ADA) proposed consuming the following foods for preventing T2DM along with weight loss: whole grain cereal products, nuts, yogurt, coffee and tea and limit consumption of red meat and sodas containing sugar [93].

In conclusion, macronutrient content of foods and meals affects significantly postprandial hyperglycemia and IR. Carbohydrates are the main macronutrients that affect glycemic responses. However, the type, total amount, other macronutrients consumed in parallel, physiological organic function, the GI/GL of carbohydrate containing foods, the extent to glucose excursions and fluctuations, and the time of day that the majority of carbohydrates is consumed, will be determinant factors of postprandial glucose and insulin increases and amelioration or not of IR.

2.1. Effects of Diet on Postprandial Hyperglycemia and IR

There are several dietary patterns that have been suggested as choices for prevention and/or treatment of T2DM individuals, independently of body weight status. All rigorously investigated healthy dietary patterns, i.e., Mediterranean-style diet [94–96], vegetarian diets [97–100], Nordic diet [101], and the Dietary Approaches to Stop Hypertension (DASH) [94,102] have been associated with a lower risk of developing T2DM. The PREDIMED trial that compared a Mediterranean-style to low-fat eating pattern for prevention of T2DM, reported a 30% lower relative risk in people of high cardiometabolic risk [103,104], whereas another trial examining a healthy Nordic diet, reported a 25% lower risk in women and 38% in men [101]. Results from the Dietary Intervention Randomized Controlled Trial (DIRECT) assigning obese adults with T2DM to a calorie-restricted Mediterranean-style, a calorie-restricted lower-fat, or a low carbohydrate eating pattern (28% of calories from carbohydrate), showed that HbA1c was lowest in the low carbohydrate group after 2 years, whereas fasting plasma glucose was lower in the Mediterranean-style group than in the lower-fat group [105]. Results from another RCT showed that despite only a 2-kg difference in weight loss, the group following a low carbohydrate Mediterranean-style eating pattern experienced greater rates of at least partial diabetes remission, with rates of 15% at year 1 and 5% at year 6 compared with 4.7% and 0%, respectively, in the group following a low-fat eating plan [106].

Results from a network meta-analysis of RCTs in T2DM adults examining 9 different dietary approaches (vegetarian, Mediterranean-style diet, high protein, moderate-
carbohydrate, low carbohydrate, low-GI/GL, paleolithic, low-fat and control diet) with a duration of at least 12 weeks, showed that all dietary approaches significantly reduced Hb1Ac, and fasting blood glucose levels, although the low carbohydrate and the Mediterranean-style diets were the most effective for HbA1c, and the Mediterranean-style and vegetarian diets were the most effective for fasting glucose reduction [107]. Although this network meta-analysis reported some benefits of the paleolithic diet, there was only one available study included in their analysis [107]. Moreover, this network meta-analysis reported that low carbohydrate diets were more effective in reducing HbA1c in T2DM aged ≥60 years, whereas the Mediterranean-style style, the moderate-carbohydrate, the low-GI/GL, the high protein, and the low-fat diets were more effective in HbA1c reduction in T2DM aged <60 years [107]. Compared to the 9 dietary approaches examined, the Mediterranean-style diet was a more effective dietary approach for improving postprandial hyperglycemia and IR compared to the other 8 analyzed dietary schemes [107].

2.2. Effects of Weight Loss on Postprandial Glycemia and IR

Weight loss has been proposed as a key strategy for the treatment of postprandial hyperglycemia, hyperinsulinemia, and IR [108]. The proposed treatment for overweight and obesity in people with prediabetes and/or overt T2DM includes firstly diet, physical activity, and behavioral therapy, then pharmacotherapy in those with body mass index (BMI) >27 kg/m² and finally bariatric surgery in those with BMI >35 kg/m² [109]. In the obese, insulin resistant, and T2DM individuals, the beneficial effects of weight loss on postprandial glycemia and IR are mostly due to the decreased blood levels of NEFA [110,111] and to an improvement in the insulin-mediated suppression of fat oxidation [112]. Postprandial switch from fat to carbohydrate oxidation was also observed in prediabetic subjects who had 14 kg weight loss [113]. This suggests that impairments in the regulation of substrate utilization by skeletal muscle can be reversible and contribute to improvements in metabolic health and glycemia. These changes are accompanied by improvements in insulin sensitivity and have been observed in most weight loss interventions. A recent consensus report of the ADA recommended a 7–10% of initial body weight reduction, if required, and maintenance to prevent progression from prediabetes to T2DM, and noted the potential for diabetes remission [114].

Eating plans that create an energy deficit should be customized to fit the person’s preferences, metabolic goals, and resources, to achieve long-term sustainment [114]. Regular physical activity, which can contribute to both weight loss and prevention of weight regain, and behavioral strategies are also important components of lifestyle therapy for weight management [115–120]. Structured weight loss programs with regular visits have been shown to enhance weight loss in T2DM [121–123]. The threshold of weight loss for maximal clinical benefits in T2DM is unknown, but it has been shown that the greater the weight loss, the greater the benefits [114]. The UK Prospective Diabetes Study (UKPDS) demonstrated that decreases in fasting blood glucose levels were correlated with the degree of weight loss [124]. A meta-analysis found that lifestyle interventions producing <5% weight loss had less effect on HbA1c, lipid profile, or blood pressure (BP) compared to studies achieving weight loss >5% [119]. Table 1 describes the effects of dietary schemes/patterns and lifestyle interventions in postprandial hyperglycemia and cardiovascular disease risk factors.

There is still a great debate as to which is the best dietary macronutrient composition for weight loss, amelioration of postprandial hyperglycemia and IR. Individualization of the macronutrient composition of the diet should depend on the health status of the individuals, their metabolic goals (glycemia, lipid profile, BP, hepatic status, renal status, etc.), physical activity/exercise levels, food preferences or aversions, and food availability [114].

2.2.1. Low Calorie Diets for Weight Loss

The Look AHEAD Trial, the Diabetes Remission Clinical Trial (DiRECT) and the Diabetes Intervention Accentuating Diet and Enhancing Metabolism (DIADEM-I) highlighted the potential for T2DM remission, defined as the maintenance of euglycemia (complete
remission) or prediabetes level of glycemia (partial remission) with no diabetes medication for at least 1 year, with a weight loss of 15 kg or more within one year following a low or a very low-calorie diet, and also using meal replacements [125–128] in people undergoing weight loss treatment. In the Look AHEAD trial, when compared with the control group, the intensive lifestyle arm (more visits, 175 min/week unsupervised physical activity, dietary modification counseling, and a weight loss goal of 10%) resulted in at least partial diabetes remission in 12% of participants as compared with 2% in the control group [125]. The DiRECT trial showed that at 1 year, weight loss associated with the lifestyle intervention resulted in diabetes remission in 46% of participants [121]. Remission rates were related to the magnitude of weight loss, rising progressively from 7% to 86% as weight loss for 1 year increased from <5% to >15% [121]. However, there is a disagreement on whether a low-calorie diet (800 kcal/day) is sustainable or really wanted by all IGT and T2DM patients. Moreover, an early rapid weight loss is typically followed by weight plateau and progressive regain [129], linked to weight loss induced cell stress, altered adipokine secretion, reduced lipolysis and induced inflammatory response in adipose tissue [130]. Therefore, a low-calorie diet may be used to reach adequate weight loss rapidly; however, for maintenance of the lost weight a less restrictive diet is used, typically a more balanced, lower fat diet.

2.2.2. Low(er)-Fat Diets

The typical dietary scheme used in almost all weight loss studies, either as the primary diet or as the control diet is a balanced, low-fat diet, and its beneficial effects on weight loss (of about 3.2 kg greater weight loss as a result of consuming a low-fat ad libitum diet) and maintenance have been well documented: compliance is easier in the long-term, and its reported effects are more pronounced in subjects with a higher body weight [131]. The strongest evidence for T2DM prevention using a low-fat diet approach comes from several studies, including the U.S. Diabetes Prevention Program (DPP) [115,132] and the Finnish Diabetes Prevention Study [133], demonstrating that a lifestyle intervention (combination of a low-fat diet combined with at least 150 min of moderate intensity weekly exercise) led to weight loss and decreased T2DM incidence for adults with overweight/obesity and IGT by 58% over 3 years [115]. Moreover, follow-up of 3 large studies of lifestyle intervention for diabetes prevention has shown sustained reduction in the rate of conversion to T2DM: 43% reduction at 20 years in the Da Qing Diabetes Prevention Study [134]; 43% reduction at 7 years in the Finnish Diabetes Prevention Study [135] and 34% reduction at 10 years [132], and 27% reduction at 15 years extended follow-up in the DPP [136] in the U.S. Diabetes Prevention Program Outcomes Study (DPPOS). The follow-up of the Da Qing study also demonstrated a reduction in CVD and all-cause mortality [137].

Results from one meta-analysis showed a greater, but not clinically significant, body fat change (by 16 g/day) favoring lower fat diets compared to lower carbohydrate diets [138]. A systematic review in people with T2DM [139], several studies [140–143], and a meta-analysis [144] suggested that lowering total fat intake did not consistently improve glycemia or CVD risk factors in T2DM, but the benefit from a low-fat eating pattern appeared to be mostly related to weight loss and not to the eating pattern itself [145,146].

In conclusion, weight loss can be achieved with low(er)-fat diets, but this seems to be inferior to low carbohydrate diets, and remission is expected to be lower than that achieved with low carbohydrate diets.

2.2.3. Low(er)-Carbohydrate Diets

The beneficial effects of lower carbohydrate diets on IR and other cardiometabolic risk factors in obese individuals are independent of weight loss [77]. There are many types of low carbohydrate diets. Each diet has varying restrictions on the types and amounts of carbohydrates consumed. However, severe carbohydrate restriction should be carefully monitored. It has been reported that short-term adoption of a ketogenic diet induces more severe hepatic IR than an obesogenic high-fat diet [147].
Table 1. Effect of dietary schemes/patterns and lifestyle interventions on metabolic outcomes (glycemia, lipidemia, cardiovascular disease factors) in individuals at high risk for developing or with diagnosed type 2 diabetes.

| Study | Health Status | Age (Years) | BMI (kg/m²) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|-------|---------------|-------------|-------------|---------------------------------------------|-------------|-----------------------|----------------------|--------------------------|
| Look AHEAD Research Group, 2010 [123] | T2DM 58.6 ± 6.8 | 35.9 ± 6.0 | 11 Years Randomized, controlled trial | 5145 | Intensive lifestyle intervention (ILI) Diabetes support and education (DSE, control group) | ILI: usual medical care combined with an intensive 4-year program designed to increase physical activity and reduce initial weight by 7% or more. DSE: usual medical care, provided by their own primary care physicians, plus three group educational sessions per year for the first 4 years | After 4 years: BW (kg) − 6.15% in ILI vs − 0.88% in DSE HbA1c (mmol/L) − 0.36% in ILI vs 0.09% in DSE systolic BP (mmHg) − 5.33 in ILI vs − 2.97 in DSE diastolic BP (mmHg) − 2.92 in ILI vs − 2.48 in DSE HDL (mg/dL) 3.67 in ILI vs 1.97 in DSE TG (mg/dL) − 25.56 in ILI vs − 19.75 in DSE |
| Johansen, et al., 2017 [148] | T2DM 54.6 | 25–40 | 12 months Randomized, assessor-blinded, single-center study | 98 | Lifestyle group (LG) Standard care (SC) | LG: 5–6 weekly aerobic sessions of 30–60 min, with 2–3 sessions of resistance training and an individual dietary plan with 45–60% CHO, 15–20% PRO, and 20–35% FAT (<7% saturated fat) SC: medical counseling, lifestyle advice | HbA1c (mmol/L) 6.65–6.34% in LG 6.74–6.66% in SC Reduction in Glu-lowering medications − 73.5% in LG − 26.4% in SC |
| Linmans, et al., 2011 [149] | IGT or T2DM 62.9 ± 11.8 | 30.4 ± 4.9 | 1 year Randomized trial | 2818 | Intervention group (IG) Control group (CG) | IG: with lifestyle coaches supervising the program, >30 min exercise for >5 days/wk CG: usual care according to a diabetes management program | HbA1c (mmol/L) − 0.12% Fasting Glu (mmol/L) − 0.17 NS changes in the CG |
### Table 1. Cont.

| Study                        | Health Status Age (Years) BMI (kg/m²) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups                                      | Dietary Intervention                                                                 | Selected Clinical Outcomes                                                   |
|------------------------------|--------------------------------------|---------------------------------------------|--------------|-----------------------------------------------------------|--------------------------------------------------------------------------------------|--------------------------------------------------------------------------------|
| Chee, et al., 2017 [150]     | Overweight/obese, T2DM, and HbA1c 7%–11% 30–65 >23 | 6 months Randomized controlled clinical trial | 230          | Usual care (UC) Trans-cultural diabetes nutrition algorithm-conventional counseling (tDNA-CC) Trans-cultural diabetes nutrition algorithm-motivational interviewing (tDNA-MI) | UC: clinical care according to Malaysian Clinical Practice Guidelines for T2DM (2009) and low-calorie diet (1200 or 1500 kcal/day) tDNA-CC low-calorie meal plan (1200 or 1500 kcal/day) and a physical activity prescription for >150 min/wk with conventional counseling tDNA-MI low-calorie meal plan (1200 or 1500 kcal/day) and a physical activity prescription for >150 min/wk with motivational interviewing | tDNA-MI BW (kg) –6.9 ± 1.3 in tDNA-MI –5.3 ± 1.2 in tDNA-CC –0.8 ± 0.5 NS in UC HbA1c (mmol/L) –1.1 ± 0.1% in tDNA-MI –0.5 ± 0.1% in tDNA-CC –0.2 ± 0.1%, NS in UC Fasting plasma Glu (mmol/L) –1.1 ± 0.3 in tDNA-MI –0.6 ± 0.3, NS in tDNA-CC 0.1 ± 0.3, NS in UC Systolic BP (mm Hg) –9 ± 2 in tDNA-MI –9 ± 2 in tDNA-CC –1 ± 2, NS in UC |
| Lindström, et al., 2006 [135] | Overweight with IGT 55 31.1            | 7 years Randomized controlled trial          | 522          | Intervention group (IG) Control group (CG)                | IG: <30% FAT, <10% saturated FAT, >15 g per 1000 kcal Fibers, and moderately intense physical activity 30 min per day or more CG: general health information at baseline without specific individualized advice | Incidence of T2DM 4.3 per 100 person-years (IG) vs 7.4 per100 person-years (CG) ↓43% relative risk in IG |
| Diabetes Prevention Program Research Group 2015 [136] | At high risk for T2DM 50.6 ± 10.7 34.0 ± 6.7 | 15 years Randomized controlled clinical trial | 3234         | Intensive lifestyle intervention (ILS) Metformin (MET) Placebo (PLBO) | ILS: low calorie and low lipid diet, plus 150 min physical activity per week MET: 850 mg x2/day PLBO: x2/day | ↓18% diabetes incidence rate MET: ↓27% diabetes incidence rate ILS: ↓8.7% aggregate microvascular prevalence in women |
Table 1. Cont.

| Study                | Health Status Age (Years) BMI (kg/m²) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups                                                                 | Dietary Intervention                                                                 | Selected Clinical Outcomes |
|----------------------|--------------------------------------|---------------------------------------------|-------------|---------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------|---------------------------|
| Ujvari, et al., 2014 [151] | 18–40 PCOS and BMI >27 kg/m² Healthy overweight/obese women (OB-C) PCOS and normal weight BMI 18.5–25 Healthy women and BMI 18.5–25 | 3 months Randomized trial                    | 49          | Overweight/obese women with PCOS (OB-PCOS) Overweight/obese controls (OB-C) Normal-weight PCOS (NW-PCOS) Healthy normal-weight controls (NW-C) | OB: PCOS-dietary restriction diet high in PRO and low in CHO (40% CHO, 30% FAT, and 30% PRO), and activity for 45 min 2–3 times/wk | BW (kg) –4.7 Ins (uU/mL) –11.9 Relative mRNA levels IRS1 +0.28 GLUT1 +0.06 |
| O’ Brien, et al., 2017 [152] | IGT 45.1 ± 12.5 33.3 ± 6.5 | 12 months Randomized, pilot study           | 96          | Intensive lifestyle intervention (ILI) Metformin (MET) Standard care (SC)                | ILI: weight loss (5–7% of initial body weight) by improving dietary patterns (decreasing fat and calories) and promoting moderate physical activity (≥150 min per week) MET: 850 mg of metformin x2/day SC: medical care and educational materials on diabetes prevention from the National Diabetes Education Program | BW (kg) –4.0 in ILI –0.9 in MET +0.8 in SC Waist circumference (cm) –4 in ILI –1.8 in MET –0.2 in SC |
| Study                               | Health Status Age (Years) BMI (kg/m²) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups                                                                 | Dietary Intervention                                                                 | Selected Clinical Outcomes                                                                 |
|-------------------------------------|--------------------------------------|---------------------------------------------|-------------|---------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------|
| Slentz, et al., 2016 [153]          | Overweight/obese) 45–75 25–35         | 6 months Randomized, parallel clinical trial | 150         | High amount/moderate intensity physical activity (1)                                   | (1) High amount—(67 KKW)/moderate intensity: equivalent of expending 67 KKW (~22.3 km (13.8 miles) per week) with moderate-intensity exercise | BW (kg)                                                                                   |
|                                     |                                      |                                             |             | High amount/vigorous intensity physical activity (2)                                   | (2) High amount (67 KKW)/vigorous intensity—equivalent to group 2, but with vigorous-intensity exercise (75% peak VO$_2$reserve) | −1.94 in (1)                                                                               |
|                                     |                                      |                                             |             | Low amount/moderate intensity physical activity (3)                                    | (3) Low amount—(42 kJ kg body weight$^{-1}$ week$^{-1}$ (KKW)/moderate intensity: equivalent of expending 42 KKW (e.g., walking ~16 km (8.6 miles) per week) with moderate-intensity (50% peak VO$_2$reserve) | −1.67 in (2)                                                                               |
|                                     |                                      |                                             |             | Lifestyle intervention with low amount/moderate intensity physical activity + diet (4)  | (4) diet + 42 KKW moderate intensity same as group 1 but with diet and weight loss (7%) to mimic the first 6 months of the DPP. | NS in (3)                                                                                 |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | −6.44 in (4)                                                                               |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | Fat mass (kg)                                                                              |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | −2.2 in (1)                                                                                |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | −2.3 in (2)                                                                                |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | NS in (3)                                                                                 |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | −6.0 in (4)                                                                                |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | AUC$_{Glu}$ (mmol/L × 120 min)                                                           |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | −73 in (1)                                                                                |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | −22 in (2)                                                                                |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | NS in (3)                                                                                 |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | −96 in (4)                                                                                |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | AUC$_{Ins}$ (pmol/L × 2 h)                                                                |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | −264 in (1)                                                                               |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | −246 in (2)                                                                               |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | −166 in (3)                                                                               |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | −348 in (4)                                                                               |
| Mensink, et al., 2003 [154]         | IGT 55.6 ± 0.9 29.8 ± 0.5             | 2 years Randomized trial                    | 114         | lifestyle intervention group (INT) Control group (CON)                                |                                                                                       | INT:                                                                                      |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | BW (kg) −2.4 ± 0.7                                                                         |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | Body fat (%) −1.0 ± 0.3                                                                   |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | Waist (cm) −1.9 ± 0.7                                                                    |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | Fast Glu (Mm) + 0.2 ± 0.1                                                                 |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | 2 h Glu (Mm) −0.6 ± 0.3                                                                   |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | Fast Ins (Mm/L) −1.8 ± 1.7                                                                |
|                                     |                                      |                                             |             |                                                                                       |                                                                                       | HOMA −0.5 ± 0.5                                                                           |
Table 1. Cont.

| Study                      | Health Status Age (Years) BMI (kg/m²) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention                                                                 | Selected Clinical Outcomes |
|----------------------------|---------------------------------------|---------------------------------------------|--------------|------------------------|---------------------------------------------------------------------------------------|----------------------------|
| Roumen, et al., 2008 [155] | IGT 54.2 ± 5.8 29.6 ± 3.8            | 3 years Randomized controlled lifestyle intervention | 106          | Intervention group (INT) Control group (CON) | INT: Dietary recommendations as per Dutch guidelines for a healthy diet, and physical activity of at least 30 min a day for at least 5 days a week
CON: usual care according to a diabetes management program | After 3 years in INT:
- BW (kg) −1.08 ± 4.30 vs +0.16 ± 4.91
- BMI (kg/m²) −0.36 ± 1.47
- BF (Kg) −1.16 ± 3.80
- Fasting Glu (Mm) +0.2 ± 0.1
- 2 h Glu (Mm) −0.05 ± 2.02
- Fasting Ins (Mu/L) −1.17
- HOMA −0.19 |

Abbreviations: IGT: impaired glucose tolerance; T2DM: type 2 diabetes; Glu: Glucose; Ins: Insulin; AUC: Area Under the Curve; CHO: carbohydrates; PRO: proteins; FAT: fats; NS: no statistically significant difference; BW: body weight; HbA1c: glycated haemoglobin A1c; BP: blood pressure; HDL: high density lipoprotein cholesterol; TG: triglycerides; BMI: body mass. index. An arrow pointing downwards indicates a decrease.
One study reported that 93% of subjects with prediabetes attained a normal HbA1c with a lower carbohydrate diet [156]. Results from a meta-analysis of 38 studies assessing a total of 6499 adults aiming to address the debate between low-fat and low carbohydrate diets and compare their effects on adiposity and lipid profiles, reported that at 6–12 months, the low carbohydrate diets were effective in improving weight loss, HDL-cholesterol and triglycerides, whereas low-fat diets were effective in improving LDL-cholesterol and total cholesterol, raising concerns about long-term adoption of low carbohydrate on potentially increasing CVD risk [157]. Another meta-analysis of RCTs [158] compared a low carbohydrate eating pattern (defined as <40% of calories from carbohydrate) to high carbohydrate eating pattern (defined as <30% of calories from fat). In trials up to 6 months long, the low carbohydrate eating pattern improved HbA1c more, and in trials of varying lengths, lowered triglycerides, raised HDL cholesterol, lowered BP, and resulted in greater reductions in diabetes medications [158]. Another meta-analysis of RCTs comparing low carbohydrate eating patterns (defined as <45% of calories from carbohydrate) to high carbohydrate eating patterns (defined as >45% of calories from carbohydrate) found that HbA1c benefits were more pronounced in the very low carbohydrate interventions (where <26% of calories came from carbohydrate) at 3 and 6 months, but not at 12 and 24 months [159], suggesting that severe carbohydrate restriction may not be health effective in the long-term, possibly due to diet compliance difficulties after 6 months of adopting a very low carbohydrate diet. Studies investigating severe carbohydrate restricted diets in T2DM, such as the paleolithic diet, are small and few, ranging from 13–29 participants, lasting no longer than 3 months, and have reported mixed effects on HbA1c, body weight and blood lipids [160–162]. A recent meta-analysis of 23 trials with 1357 participants examining the efficacy and safety of low carbohydrate diets and very low carbohydrate diets in T2DM, reported that at 6 months, compared with control diets, low carbohydrates diets achieved higher rates of diabetes remission (57% vs 31%; defined as HbA1c <6.5%), but data on remission at 12 months were sparse, ranging from a small effect to a trivial increased risk of T2DM [163]. The authors of this meta-analysis reported large clinically important improvements for weight loss, triglycerides, and insulin sensitivity at 6 months, which was diminished at 12 months [163]. Very low carbohydrate diets were found to be less effective than less restrictive low carbohydrate diets for weight loss at 6 months, explained by low dietary adherence [163]. Moreover, participants were found to have deteriorated quality of life and LDL-cholesterol at 12 months [163]. Finally, in another meta-analysis comparing low carbohydrate to high carbohydrate eating patterns, the larger the carbohydrate restriction, the greater the reduction in HbA1c, though HbA1c was similar at duration of 1 year and longer for both eating patterns [164]. However, it is questionable whether a low carbohydrate diet is sustainable long-term. In addition, low carbohydrate diets need to be higher in plant protein sources than animal protein sources to avoid unwanted elevations of plasma lipids. Both low carbohydrate diets and low-calorie diets require supplemental micronutrients (as do regular vegetarian and vegan diets), although it is unclear to what extent micronutrients can be provisioned by animal protein sources without unwanted increases in plasma lipid levels.

Overall findings tend to support evidence from existing RCTs and observational studies showing that people with markers indicating higher risk for diabetes, prediabetes or IR have lower risk when they reduce calorie, carbohydrate, or saturated fat intake and/or increase fiber or protein intake (lean animal protein or plant protein) compared with their peers [114]. For purposes of weight loss, the ability to sustain and maintain an eating plan that results in an energy deficit, irrespective of macronutrient composition or eating pattern, is very important for success [165–168]. Studies investigating specific weight loss eating plans using a broad range of macronutrient composition in people with T2DM have produced mixed results regarding efficiency and efficacy on body weight, HbA1c, lipid profiles, and BP [140,141,144,169–176]. As a result, the evidence does not identify one eating plan or a certain macronutrient composition that is clearly superior to other [177] and can be generally recommended for weight loss for people with T2DM [178]. Thus,
an individualized plan is warranted, taking into consideration dietary preferences along with food preferences, metabolic goals, and ability to comply to and maintain the eating plan [179]. In each case (low-calorie or low fat or low carbohydrate diets), a Mediterranean-style diet is likely one of the best diet options after reaching weight loss goals, with intermittent re-establishment of optimal weight by reverting to either the low-calorie diet or to the low carbohydrate diet [95]. A partial reason for suggesting the Mediterranean-style diet as the maintenance diet after reaching weight loss goals is that it is a low GL diet, which supports a lower body weight.

In conclusion, weight loss is the best remedy for ameliorating postprandial hyperglycemia and IR. However, it is unclear whether weight gain and loss per se are the primary or intermediate cause relevant to T2DM or its remission. Weight loss should be rapid or steady, and the choice of dietary pattern and/or dietary macronutrient composition, should be individualized, patient-centered, and depend entirely on the person’s lifestyle, habits, eating practices, preferences, and metabolic goals.

2.3. Effects of Nutrient and Meal Sequence on Postprandial Glycemia and IR

Diurnal regulation of glucose metabolism in the postprandial and postabsorptive state have brought into light the importance of food and meal sequence. The sequence of meals plays an important role in postprandial glycemic responses; preceding meals sensitize the metabolic and incretin system to the following ones, thereby improving glucose tolerance during the day (Figure 3) [31].

During sleep at night, the gradual development of IR, due to growth hormone and cortisol surges, ensures that blood glucose levels will be maintained within normal levels until awakening, by switching from glucose to NEFA oxidation in skeletal muscle [31]. The increase in lipolysis and supply of NEFA to the liver and kidneys will also ensure stimulation of gluconeogenesis and HGP [31]. Thus, in addition to meal composition and size, the timing of macronutrient consumption during a meal seems to be a key regulator of postprandial hyperglycemia [31,74]. There is some evidence suggesting that premeal consumption of nutrients such as water, fat, protein, or fiber as “preloads” delay the rate of postprandial glucose absorption from the small intestine and attenuate insulin secretion and glucose excursions [74,180–185]. Moreover, it has been shown that the timing of carbohydrate ingestion (i.e., carbohydrate-last meal patterns) can markedly reduce postprandial glycemia by delaying gastric emptying, enhancing glucose-stimulated insulin release, and affecting insulin clearance [31,74]. One study showed that ingestion of olive oil (fat) half-hour before the consumption of a carbohydrate meal (potato) slowed the gastric emptying rate, mitigated the postprandial rises in the levels of glucose and insulin, and GLP-1 secretion was stimulated after the meal [181]. Similarly, protein preload (in many studies whey protein) has been examined in multiple clinical trials, and favorable effects on postprandial glycemia have been reported [182,186,187]. Consumption of whey protein half-hour before a carbohydrate meal, resulted in reduced postprandial hyperglycemia and increased plasma insulin levels [182]. Reduced postprandial glucose levels was also a key finding in other trials as well when protein preloads were ingested 15 min or in one study 2 h before a meal by subjects with IGT [188–190]. In those studies, protein preloads before meals were also associated with increased satiety, reduced hunger, and decreases in HbA1c [189,190]. The magnitude of food and meal sequence is greater in T2DM than in healthy people, being comparable and additive to current antidiabetic medications and has been shown to sustain over time, offering a simple, effective, safe, and inexpensive tool for treating postprandial hyperglycemia and IR [31,74].
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Figure 3. Overall metabolic effects of diet, nutrients, timing of eating occasions, speed of eating, physical activity, and alternative dietary interventions on postprandial hyperglycemia and insulin resistance. Abbreviations: Glu: glucose; GLP-1: glucagon-like peptide 1; CHO: carbohydrates; IL: interleukin; PYY: peptide YY; GLUT: glucose transporter; ppd: postprandial; BW: body weight; Ins: insulin; HOMA: homeostatic model assessment for insulin resistance; HbA1c: glycated hemoglobin A1c; IRS: insulin receptor substrate; TG: triglycerides; HDL: high density lipoprotein cholesterol; chol: cholesterol; LDL: low-density lipoprotein cholesterol. An arrow pointing upwards or downwards indicates an increase or decrease.

A recently proposed nutrient/meal sequence for reduced postprandial glycemic responses and improved insulin sensitivity as has been illustrated in a food pyramid paradigm from bottom (highest consumption) to top (minimum consumption) may be the following: (a) low-energy dense foods containing water, such as soups, vegetables (5 colors: fresh, cooked, sprouted, fermented) and fruits should be consumed first; followed by (b) light dairy foods (milk, yogurt, cheese), fish/seafood (lightly cooked and better cooked and consumed with vegetables), poultry and eggs (boiled); followed by (c) lean red meat or soy protein (tofu, tempeh, isolate); followed by (d) whole (legumes/pulses (sprouted, cooked), whole grains (cooked, pasta, bread, porridge, cereals), and starchy vegetables [1]. In contrast, the following foods were proposed to be consumed scarcely if at all and include processed/refined grain/flour products (breads, cakes, breakfast cereals, baked goods) fried/fatty/processed carbohydrate foods (chips, other snacks), processed/refined grain/flour foods (breads, cakes, breakfast cereals, baked goods), fried/fatty foods (chips, other snacks), sweets and sugary drinks [1]. These meal sequence options have been sug-
gested to ameliorate postprandial glucose/insulin excursions and oxidative responses by enhancing incretin (i.e., GLP-1) secretion and delaying gastric emptying [1,191]. Moreover, carbohydrate sources containing slower absorbable carbohydrates, low in GI/GL, high in fiber and protein (i.e., whole grain cereals, beans, lentils, pasta, etc.) seem to be optimal for increased satiety, and reduced energy intake [1]. Therefore, it seems that meal and food sequence may play an important role in ameliorating postprandial hyperglycemia and IR and may be included in the dietary recommendations.

3. Effects of Intensive Lifestyle (Diet + Exercise + Behavior Modification) Interventions on Postprandial Hyperglycemia and IR: A Focus on Exercise

Intensive lifestyle interventions include: (a) An energy reduced (500–750 kcal/day deficit), typically a low-fat diet (≤30% total calories, 1200–1500 kcal/d for women, 1500–1800 kcal/d for men), although other dietary schemes have been also used, such as low carbohydrates, Mediterranean-style diet, etc., (b) Moderate to high intensity physical activity (≥150 min/week of aerobic physical activity, i.e., brisk walking). (c) ≥14 at least 45-min sessions (individually or in groups) supervised by specialists (doctors, dietitians, psychologists, trainers, etc.). (d) Daily monitoring of food intake and physical activity, facilitated by paper diaries or applications, weekly monitoring of weight. (e) Structured curriculum of behavior change (i.e., DPP, including goal setting, problem solving, and stimulus control), regular feedback and support from a trained interventionist, in 6 months, with (f) a goal to achieve 7–10% weight loss [151,152,192,193].

Individuals who achieve short-term weight loss should be advised to attend long-term (≥1 year) comprehensive weight management programs, with at least monthly contact and ongoing monitoring of body weight, follow a reduced calorie diet and engage in high levels of physical activity (200–300 min/week) [192,193]. To achieve long-term weight loss >5% there should be a short-term (3-month) high-intensity lifestyle interventions using very low-calorie diets (≤800 kcal/day) and long-term comprehensive weight management counseling to maintain weight loss [192,193].

Behavioral therapy (using cognitive behavior techniques, goal setting, nutrition education, monitoring, and feedback exchange) by trained dieticians and specialized health care professionals is an important component and may be achieved either by meeting in person or using technology in individual or group sessions [192,193].

Technological advances (phone applications, online appointments, phone calls, meeting online platforms and social media) can offer valid alternatives and may be equally effective [24,194,195]. One study demonstrated that adherence to self-monitoring via a technological application (mobile health, mHealth, supported by mobile devices) significantly improved weight loss [196]. Preventing negative emotions and distress can be beneficial for people with obesity, prediabetes or T2DM, and IR and can enhance adherence to the suggested dietary plan and better management of food choices [197].

Lifestyle modification programs typically prescribe 150–180 min per week of moderately vigorous aerobic activity, such as brisk walking or cycling [192]. People lacking time to exercise are encouraged to engage in multiple brief bouts (i.e., 10 min) of activity throughout the day and to increase their lifestyle activity, i.e., by using stairs vs taking the elevator [192]. The combination of diet and exercise is the most effective for reducing body weight and achieving better metabolic control compared to diet or exercise alone [148,153,193]. This combination has a dual effect both on mediating energy intake and on increasing energy expenditure to achieve a negative energy balance [148,153,193]. It has been shown that weight loss alone is effective in improving insulin sensitivity, but it is more likely to improve fasting fat oxidation and mitochondrial function if combined with exercise training [111]. It has been shown that for each kg of mean weight loss, there is a mean HbA1c reduction of 0.1% in obese and overweight subjects with T2DM; and that HbA1c lowering is greater in poor glycemic controlled vs well controlled populations with the same degree of weight loss [198]. It has also been reported that chronic exercise training improves the capacity of skeletal muscle to utilize fatty acids for fuel during exercise, and
may also improve fasting fat oxidation, indicating increased metabolic flexibility, thus improving fine-tuning of the balance between fatty acid uptake, oxidation and intramyocellular triacylglycerol turnover in skeletal muscle, reducing lipid intermediates, and improving insulin sensitivity [111]. The increased exercise-induced insulin sensitivity is accompanied by a reduction in the mRNA expression of acetyl-CoA carboxylase-2 and an increase in the protein levels of hydroxyacyl-CoA dehydrogenase in muscle [199]. It has been shown that the mRNA levels of both insulin receptor substrate-1 (IRS-1) and glucose transporter-1 (GLUT-1) were increased in overweight/obese women with polycystic ovary syndrome following a lifestyle intervention that included increased physical activity (prescribed aerobic activity for 45 min two or three times/week) [151], suggesting that the changes occurring at a molecular level after exercise directly affect the mechanisms involved in insulin sensitivity and glucose metabolism in muscle. One study showed that afternoon exercise (i.e., high intensity interval training/HITT; individuals cycled at 75 rpm with a load of forced rest after 1 min, repeated six times with 1 min rest in between, three sessions/week, using continuous glucose monitoring, and recorded daily food intake) was found superior to morning exercise in improving glucose levels in T2DM [200]. A recent systematic review of 20 studies with a total of 352 participants (15 studies with T2DM and 5 with healthy subjects) concluded that exercise (moderate-intensity aerobic exercise in 18 studies, HIIT in 3 studies, and resistance exercise in one) performed post-meal instead of pre-meal regardless of time of day, improved postprandial hyperglycemia [201]. Finally, the combination of weight loss and exercise improved the aerobic capacity of skeletal muscle, increased the number of mitochondria per muscle cell and improved electron chain transport activity in obese subjects [202], having a favorable impact on postprandial glucose excursions.

Results of a meta-analysis of 16 RCTs with ≥24 weeks of follow-up using lifestyle interventions including diet and exercise, showed a four-fold increased probability of metabolic syndrome remission compared to pharmaceutical interventions [203]. Another meta-analysis of 20 RCTs (9 with people with IGT, and 11 with T2DM) with diagnosis of 6–48 months duration showed that lifestyle interventions lowered T2DM risk by 20% in people with IGT after 10 years of follow-up but did not reduce all-cause mortality in T2DM [204]. A more recent meta-analysis of 28 RCTs showed that lifestyle interventions were more effective than the standard care regarding the glycemic control of subjects with T2DM, particularly when there was weight loss [29]. These results have also been reported in other systematic reviews and meta-analyses all supporting the beneficial effects of lifestyle interventions, regardless of study design, on weight loss and weight maintenance, delay of T2DM onset and progression and decreased risk for CVD [24,27,28,205–210], as well as reduced or eliminated need for anti-diabetic medications [27,127,128,148]. Both shorter-term (6–12 months) and longer-term (>12 months) studies showed that more intensive lifestyle interventions (i.e., medications alone vs medications plus diet plus 5–6 aerobic training sessions/week, duration 30–60 min, of which 2 to 3 sessions were combined aerobic and resistance training) led to greater HbA1c reduction (6.5% compared to 1% with less intensive interventions) [148–150] and significant reduction in anti-diabetic medications (up to 73% with lifestyle interventions compared to 23% in the control group) [148]. Studies have also shown that the beneficial metabolic changes (body weight loss of 6%, decreased HbA1c by ~0.4%, decreased insulin resistance/HOMA-IR index, ameliorated postprandial hyperglycemia, decreased BP, increased HDL cholesterol, and decreased plasma triglycerides) may last in the long-term [123,154,155]. One study showed that after lifestyle-induced weight loss, improvements in insulin secretion in older, obese, nondiabetic subjects seemed to be largely dependent on improved insulin sensitivity. However, in older, obese diabetic subjects, improved insulin secretion was a consequence of better β-cell function, demonstrating that changes in insulin secretion after lifestyle interventions (including fully supervised aerobic treadmill-walking exercise conducted for 1 h/day, 5 days/week, with increased intensity at week 4 maintaining 80–85% maximum heart rate) may be mediated via alterations in the secretion of incretins, such as GIP [211].
In conclusion, the combination of a reduced energy diet and regular moderate to high intensity physical activity aiming at moderate body weight loss and maintenance, and nutritional behavior modification may ameliorate postprandial hyperglycemia and IR.

3.1. Metabolic Effects of Different Types of Physical Activity

In general, exercise can be classified as aerobic (long duration exercise involving a large number of muscles, such as brisk walking or jogging, swimming, cycling) or anaerobic (shorter duration exercise such as sprinting or weightlifting using free weights/weight machines/elastic resistance bands), depending on the predominant mechanisms used by muscle to provide energy [212]. HIIT is also a popular form of aerobic training consisting of short periods of intense exercise interrupted by short periods of active recovery or rest (e.g., such as using a stationary cycle ergometer) [213]. For the working muscle, a constant supply of ATP is required; this is accomplished via either the anaerobic or the aerobic metabolic pathways, which do not function independently but rather synergistically by interactions between the exercising muscles and distant organs/tissues (liver/adipose tissue/cardiovascular system/brain) to provide energy and maintain blood glucose levels within the normal euglycemic range (reviewed in detail in reference [12]).

3.1.1. Aerobic Exercise

At rest (typically during sleeping at night), skeletal muscle derives most of its energy from the oxidation of NEFA than that of glucose [214]. James et al., first reported in rats that aerobic exercise increased the sensitivity of skeletal muscle and adipose tissue but not liver to insulin using euglycemic-hyperinsulinemic clamps with infusions of radiolabeled glucose; the rates of glucose oxidation in muscle were increased [215].

During exercise, the rapid increase in energy demands in the working muscles requires a substantial change in the mobilization and oxidation of carbohydrates and lipids depending on the intensity and duration of exercise. In a systematic study, Romijn et al. [216] described the effects of aerobic exercise intensity (low/25% VO\textsubscript{2max}, moderate/65% VO\textsubscript{2max}, or high/85% VO\textsubscript{2max}) and duration (30 or 120 min) on the utilization of glucose and NEFA in endurance-trained cyclists in the fasting state on 3 consecutive days using a stationary cycle ergometer; indirect calorimetry and infusions of stable isotopes were used to estimate energy turnovers and substrate mobilization; the mechanisms are as follows [216]: (a) At the onset of exercise, insulin secretion is rapidly decreased and that of glucagon increased. The sympathetic nervous system is activated to mediate increases in catecholamines (their levels in plasma increase from ~2-fold to ~4-fold and ~300-fold at low, moderate, or high intensity exercise, respectively) to maintain BP, increase blood flow rates in muscle for the effective delivery of hormones and substrates, and facilitate increases in endogenous glucose production and lipolysis in adipose tissue depots; plasma levels of growth hormone and cortisol also increase with exercise intensity to supplement catecholamine effects [216–219]. The decrease in plasma insulin levels and the increase in anti-insulin hormones stimulate hepatic glucose production (~1.5-fold, ~4-fold, or ~6-fold during low, moderate, or high intensity exercise, respectively) via glycogenolysis/gluconeogenesis to maintain plasma glucose levels within the normal range and increase the rates of lipolysis in the adipose tissue to release NEFA and glycerol [216,217,219]. The rates of glucose uptake in the contracting muscle increase independently of insulin due to the translocation of GLUT4 transporters from intracellular pools to the surface membrane; hexokinase activity and the rates of glucose phosphorylation also increase [217,220–222]. (b) At low intensity exercise, muscle derives energy mainly from the utilization of circulating NEFA produced by lipolysis in the peripheral adipose tissue (~90%) rather than glucose (~10%); the subtle increase in glucose oxidation is met solely by glucose uptake from the circulation and therefore, muscle glycogen is not utilized. Lipolysis from intramuscular triglycerides is not increased [216]. (c) When the intensity of exercise increases from low to moderate, there is a progressive shift from the utilization of NEFA (~50%; derived from the circulation or from muscle triglycerides with equal contribution) to that of glucose (~50%, mostly from muscle
glycogen stores, for anaerobic metabolism and production of lactate) [216,223,224].

(d) When the intensity of exercise increases further from moderate to high, energy consumption relies almost exclusively to glucose (derived ~10% from the circulation, and ~60% from muscle glycogen) than to NEFA (~30%; from the circulation or from muscle triglyceride stores) [216], (e) Increase in the duration of exercise from 30 to 120 min does not modify substrate contribution at low intensity; however, at moderate/high intensity there is a progressive increase in the reliance on circulating NEFA and glucose, leading to a decrease in muscle glycogen stores. (f) In healthy subjects (age ~64 years, BMI ~34 Kg/m\(^2\)) the combination of diet intervention with aerobic exercise (treadmill walking or cycle ergometer at ~80% maximum heart rate, 5 days/week, 60 min/day for 12 weeks) improved insulin sensitivity in muscle (euglycemic-hyperinsulinemic clamps), and increased the expression of genes regulating enzyme activities and oxidative capacity in the mitochondria (muscle biopsies) improving NEFA transport and oxidation in this tissue; interestingly, these effects were independent of weight loss (8–10%) and the glycemic index of the diets [225].

3.1.2. Anaerobic Exercise

Exercising against resistance (strength training) has gained popularity since it can improve body composition by increasing lean body mass, an effect aerobic exercise practically does not have. Although the mechanisms increasing glucose uptake into muscle cells are generally the same as those described earlier in this section, in contrast to aerobic exercise that requires a complex interplay between glucose and lipids, anaerobic exercise (such as weightlifting or sprinting) requires only glucose derived from muscle glycogen for ATP production [12]. The anaerobic metabolism of glucose in the glycolytic pathway will produce lactate which will be turned back into glucose in the liver. The recycling of glucose between muscle and liver constructs a “substrate cycle” (Cori cycle). The role of substrate cycles in metabolic pathways is important since they can improve the sensitivity of the pathway to external signals, such as hormones (e.g., catecholamines through increases in the sympathetic nervous system activity during exercise) [57]; they also produce heat, an aspect of substrate cycling that is involved in weight control and, therefore, obesity [226]. In anaerobic exercise the “Cori cycle” is of major importance since it is responsible for providing most of the ATP during muscle contractions [12].

An important effect of resistance exercise is that in can increase muscle mass quantity but also quality through the IGF-1/phosphatidylinositol-3 kinase/protein kinase-B pathways [227]. Resistance exercise but not aerobic exercise in rats has been shown to increase IGF-1 expression and subsequent GLUT-4 translocation and increase of glucose uptake in skeletal muscle preparations in-vitro [228]; increases in plasma IGF-1 concentrations have also been reported in humans during high-intensity resistance training [229]. Earlier studies in rats showed that IGF-1, with its insulin-like effects, increases the apparent sensitivity of skeletal muscle to insulin either in-vitro [230] or after acute or chronic (10 days) administration in-vivo [231].

Muscle hypertrophy with resistance training is popular and has attracted attention regarding its impact on glucose homeostasis and the mechanisms involved. It has been suggested that the increase in insulin-stimulated glucose disposal with resistance exercise may be due, at least in part, to the increase in muscle mass, whereas aerobic training may enhance insulin sensitivity by changes in the intrinsic metabolic pathways in muscle cells [232]. In a comprehensive recent narrative review, Paquin et al, summarized all the available literature on this subject aiming to provide a mechanistic explanation for this type of exercise in improving insulin sensitivity and health outcomes [233]. As these authors [233] conclude, regular resistance exercise improves insulin sensitivity by a number of mechanisms, including greater muscle vascularization and increases in blood flow rates. However, the question if the beneficial effects of resistance exercise are due mostly to muscle hypertrophy or to a better quality of muscle cells needs further investigation [233].

The mechanisms described in this section regarding the crosstalk between tissues for the mobilization of substrates during exercise depend on how much glycogen is stored
in the liver or muscle at the beginning of exercise, which in turn depends on the amount of carbohydrates in the previous meal or on any performance of physical activity prior to exercise, as well as on the intensity/duration of exercise, and the physical fitness of the subjects. These factors should be considered since they may explain, at least in part, the discrepancies between studies in the literature.

In subjects with diabetes or obesity, exercise is the cornerstone of therapeutic interventions. It increases insulin sensitivity independently of changes in body fat with nutritional interventions, improves metabolic control, contributes to the decrease in body weight, decreases the risk for cardiovascular/kidney complications and hypertension, and improves physical fitness and well-being [234–242].

4. Chrononutrition

Most living organisms exhibit circadian (diurnal) rhythms, which essentially control rhythmicity in physiological activities, such as rest/active, and feeding/fasting cycles [243,244]. In mammals, the functions of nearly all organs and systems, such as the pancreas, the gastrointestinal system (including the intestinal microbiome), the adipose tissue, the immune system, the endocrine system, the cardiovascular system, thermoregulation, brain activity, etc. are regulated by circadian rhythms, and may exhibit daily oscillations [243,244]. Endogenous molecular circadian clocks display 24-h oscillations and govern the circadian rhythms [243,244]. The center of these clocks is in the hypothalamus, within the suprachiasmatic nucleus (SCN), which contains neurons oscillating periodically approximately every 24 h, and acts as a “master clock” for the peripheral clock systems present in all other tissues and cells in the body [243,244].

The molecular clocks regulate the transcription of a myriad of clock-controlled genes either directly by the two master heterodimeric transcription factors CLOCK and BMAL1 and other clock regulated transcription factors, or indirectly through other clock output proteins [245,246], to drive rhythmic gene expression and regulate biological functions under circadian control [247]. Also, the heterodimer CLOCK and BMAL1 rhythmically activates the expression of their transcriptional repressors, Period (Per1 and Per2) and Cryptochrome (Cry1 and Cry2) [247]. Several factors involved in glucose metabolism, such as insulin and cortisol are expressed and secreted following circadian stimuli similarly to the main organs involved in glucose uptake and metabolism do (i.e., liver, pancreas, adipose tissue, muscles) [248,249]. Glucose tolerance peaks during daylight and is lower during the night/dark cycle. Insulin follows a temporal control of production, and its release is controlled by both feeding-fasting patterns and circadian rhythms [250]. The levels of nutrients in the blood are a physical signal for insulin production, and the circadian and SCN systems modulate both insulin and glucagon by controlling their synthesis at the cellular level [249,251–253]. Animal and in vitro studies with BMAL1−/− and Per1−/− show a disruption on glucose homeostasis and insulin release despite displaying normal activity and feeding-fasting rhythms [251,254]. This demonstrates how the molecular clock at the cellular and tissue levels can have significant effects on physiology [249]. Furthermore, the pancreatic β-cells receive parasympathetic input, which is under circadian control by gamma-aminobutyric acid/GABA-ergic projections from the SCN [252]. Similarly, the SCN has both glutamatergic and GABAergic projections to modulate the liver and influence glucose production [252]. The time of day that the function of the pancreatic β-cells peaks is still unknown, but some data suggest that it may peak during lunch hours; indicating that carbohydrate intake is better metabolized during these hours resulting in ameliorated postprandial hyperglycemia [83]. Finally, cortisol, a steroid hormone involved in metabolism and stress responses, also follows a daily rhythmicity. Adrenal gland cells have cellular clocks to temporally influence production and release of cortisol [249,255]. Peak levels of cortisol synchronize with the beginning of the active phase to aid in arousal (early morning in diurnal animals and early night in nocturnal animals) [255].

The circadian clocks can respond to environmental variables which may act as circadian time cues also known as zeitgebers or time givers or time cues [243]. While light
(light/dark cycles in the day) is the most potent time signal and it caters to the SCN clock system, food availability (feeding/fasting cycles) and activity-rest pattern are other important zeitgebers that entertain the peripheral clocks, which dominate local physiological processes such as glucose and lipid homeostasis, hormonal secretion, the immune responses, and the digestive system [243]. Circadian rhythms may also influence energy balance [243]. Synchronization of peripheral clocks is essential to ensure temporally coordinated physiology [243]. Significant variations from this circadian rhythmicity have been shown to disrupt metabolism and homeostasis in many organs and can manifest in different ways ranging from irritability and fatigue to several chronic diseases, including obesity, T2DM, CVD, and inflammation [243]. Unlike the changes in daylight, which are fixed depending on the geographical location, changes in food intake and feeding time in a day can affect nutrient sensing pathways that act to maintain homeostasis [243]. Synchronization of food consumption, food quality, and metabolic rhythms in a day may provide optimal metabolism and positively impact health. In the past years we have valuable information regarding our organism synchronization including the following approximations: melatonin drops at 7:00, cortisol rises at 8:00, at 8:30 we have bowel movements, at 10:00 insulin sensitivity peaks, at 11:30 we have high alertness, at 14:30 muscle performance peaks, at 20:00 melatonin rises, at 22:00 body cools down, at 1:00 sleep deepens, at 3:00 body temperature rises, and so on [256].

The term “chrononutrition” describes the direct relationship between time of day (eating during light time vs evening vs night), eating occasion (consuming or skipping meals, macronutrients’ amount and type, and time of day consumed, main course eaten at early lunch hour before 14:00–15:00 vs late lunch hour after 15:00 vs evening dinner, etc.), the body’s daily circadian rhythms and their effects in metabolic health. This concept reflects the basic idea that, in addition to the amount and content of food ingested, the time of ingestion itself is also critical for the well-being of an organism. As such, one can envision an “optimal” feeding schedule synchronous to the body’s metabolism that may provide benefits for the overall health.

4.1. Feeding and Circadian Solidarity

A study tracking the participants’ eating behaviors via a smartphone app, revealed erratic diurnal feeding patterns [257]. This may be expected considering the modern way of life where late eating (in many cases with over 30% of the daily calories being consumed after 18:00), lack of stable eating patterns, and skipping meals are indeed very common, particularly in western societies (Figure 3). This type of lifestyle can lead to circadian misalignment and negatively impact metabolism and glucose control. Circadian misalignment has been observed in chronic jet lag and night shift workers, who exhibit a lowered glucose and lipid tolerance, increased liver lipogenesis, higher body mass index (BMI), and are prone to various metabolic diseases such as obesity, CVD, gastrointestinal disorders, and T2DM [258–260]. In nocturnal rodents, feeding during the daytime hours for a week completely altered the phase of the circadian expression of clock and clock-controlled genes in the peripheral tissues [261,262]. In humans, a circadian misalignment protocol with a 12-h shift of all meals resulted in elevated blood glucose and reduced insulin levels after the evening dinner [263]. Finally, individuals with an inherent later chronotype also tend to have a higher BMI presumably because they consume their meals later in the day [264]. All the above, underline the importance of timing of food consumption.

Timing of food intake according to the body’s circadian rhythms can align the zeitgeber with the metabolic responses from different organs resulting in a healthier postprandial state. Lipogenesis in the liver can also be reset by changes in nutrition modulating fat usage by muscle via the circulating lipids [265,266]. The liver clock is affected by the balance between food intake and starvation intervals [267–269]. In this context, because breakfast is consumed after the longest fasting period of the day, it is usually the most important meal to modulate the phase of the liver clock [270]. On the other hand, late evening dinners or midnight snacks change the length of the fasting period and thus alter the phase of
peripheral clocks. Regarding the effect of nutrients on the liver clock, a combination of carbohydrate and protein is essential to reset it and cause a phase-shift, whereas protein, carbohydrates, or lipids, when consumed alone, are insufficient [270].

In conclusion, the feeding time (day vs night), the hours of daily fasting until the first meal is consumed and irregular meal patterns may play a significant role in postprandial hyperglycemia and IR, and personalized advice to patients should be provided, taking these factors into consideration.

4.2. Effects of Meal Timing on Postprandial Glycemia and IR

There is intense scientific interest in examining the validity or not of the famous quote “Eat breakfast like a king, lunch like a prince, and dinner like a pauper” (Figure 3). In modern times a tendency for late-eating patterns has been observed and in many cases, it is accompanied by skipping breakfast [257]. One study showed that the unfavorable effects of evening eating on metabolic risk in healthy volunteers were due to lower concentrations of epinephrine/norepinephrine and higher concentrations of acylated ghrelin after the evening dinner (at 20:00), with the opposite behavior exhibited after the morning meal (at 8:00) [271]. Another study in healthy adults showed that eating in the evening (at 17:00) vs morning (at 9:00) resulted in higher postprandial glucose concentrations and GIP, and lower levels of five glycolysis/tricarboxylic acid cycle/nucleotide-related metabolites, and eighteen amino acid-related metabolites, all involved in an exacerbation of postprandial hyperglycemia [272]. Another study showed that transferring 100 kcal of fat intake from night (20:30–05:00) to earlier periods was associated with lower low-density lipoprotein (LDL) cholesterol levels, especially when transferred to lunch time (11:30–13:30) or evening (17:30–20:30) [273]. In a large cross-sectional study with 11,594 subjects with T2DM, evening chronotype was independently associated with T2DM, and people had lower odds of having HbA1c at the recommended levels of <7% [274].

Table 2 describes the effects of the first meal of the day on indices of glycemic control. Several studies have suggested that skipping breakfast may be associated with increased BMI (change in BMI from baseline over time), altered lipid and glucose homeostasis, increased LDL, metabolic syndrome and T2DM [260,275–278], although reverse causality cannot be ruled out, i.e., people may feel fuller when obese and so more likely to skip breakfast. Regular breakfast consumption has also been shown to increase satiety and thermogenesis and improve the quality of the diet by inclusion of fibers and nutrient-rich foods [279,280]. The effect of breakfast on weight loss is controversial with some short-term studies showing a modest effect with a slightly greater weight loss in individuals who consumed breakfast [281], longer term prospective studies showing no clinically significant effect [276], and others in healthy individuals showing minor differences in nutritional and perceived characteristics of breakfast regarding medium-term [282].

Similarly, the effects of breakfast consumption on postprandial glycemia and IR is also controversial. Results from few RCTs in lean and obese subjects suggest that skipping breakfast regardless of body weight may adversely affect insulin sensitivity [280,283]. A recent meta-analysis of prospective cohort studies reported that breakfast omission was associated with 55% greater risk of T2DM compared with breakfast consumption [277]. In a 16-year follow up cohort, men who skipped breakfast had higher risk for developing IR and T2DM [278]. Additionally, individuals with later chronotypes and a pre-existing history of T2DM were reported to have worse glycemic control when breakfast was skipped [284]. However, reverse causality cannot be ruled out, i.e., the increased risk for IR and T2DM may be due to the higher BMI and not to skipping breakfast.

A few clinical trials have been conducted in individuals with prediabetes or T2DM (summarized in Table 2). Breakfast omission, and thus continued fast until lunch in T2DM, overweight or obese individuals was shown to result in reduced insulin and GLP-1 after lunch (the first meal of the day in one study), and lower glycemic and insulinemic responses, leading to postprandial hyperglycemia after lunch consumption [72,283,285]. In contrast, healthy individuals skipping breakfast had better glycemic control after lunch, with blood
glucose levels slightly elevated or unchanged and insulin secretion and sensitivity slightly increased [286,287]; only glucose variability during the day was reported to be higher in the fasting group [280]. A Japanese study in healthy individuals showed that breakfast omission together with late-evening dinner consumption, but not breakfast skipping alone, resulted in hyperglycemia [288]. Another study examining the acute effects of breakfast consumption or omission in both healthy individuals and T2DM, reported that breakfast consumption vs breakfast skipping affected positively clock and clock-controlled gene expression leading to normal oscillation patterns, while skipping breakfast resulted in increased postprandial glycemic responses [285]. Few available RCTs suggest that the reduced insulin sensitivity observed later in breakfast skippers may be due to the anti-insulin hormones peaking in the morning hours (i.e., cortisol), the prolonged elevation of plasma NEFA levels due to the extension of overnight fast, and the low recruitment of GLUT-4 transporters and insulin-stimulated glucose uptake in muscle (post-fasting) [289].

4.2.1. Effects of Meal Macronutrient Composition on Postprandial Glucose and IR

The carbohydrate type and content of a meal are the main determinants of postprandial hyperglycemia and insulin response (Figure 3). Although the amount of carbohydrate intake required for optimal health is unknown, an intake of >130 g/day seems to be required for brain’s demand for glucose, and of body’s metabolic processes, including glucose metabolism [114]. Breakfast is the most studied meal (summarized in Table 2). Possibly breakfast hours may be the worst time of the day to consume high amounts of carbohydrates, while small amounts would initiate an increase in insulin sensitivity. Thus, limiting the readily available carbohydrates in breakfast and replacing them with other energy and nutrient sources may be beneficial for achieving better glucose regulation and lower glycemic excursions [84].

Results of a recent meta-analysis of prospective cohort studies with 4–26 years follow-up showed a role for GI and GL as causal factors contributing to incident T2DM and recommended considering the inclusion of GI and GL in food and nutrient-based recommendations [290]. Another recent meta-analysis reported that consuming low GI and GL at breakfast attenuated acute postprandial hyperglycemia [291]. Consumption of fibers at breakfast (i.e., beta glucans, whole grain cereals) may slow the rates of gastric emptying and intestinal glucose absorption, and contribute to increased satiety, thus reducing postprandial glucose responses in a dose-dependent manner [292]. We have shown in a series of studies that foods (such as Ceratonia siliqua, carob) and characteristics (such as large bran size or the sucrose to oligosaccharide ratio in honey varieties of similar botanical origin and characterization explaining 30% of the postprandial glucose differences) are able to reduce glucose excursions and overall postprandial glycemic responses [293–295], indicating the need for more clinical trials investigating the effects of functional foods on postprandial glycemia and IR.

Accordingly, protein and fat consumption at breakfast have been shown to increase satiety and better regulate post-meal glucose responses [275,276]. Long term substitution of available carbohydrates at breakfast with proteins also decreased blood triglycerides and regulated BP [275]. Short- and long-term clinical trials in subjects with T2DM [84,296–300] (Tables 2 and 3) suggested that consuming more lower fat containing proteins than carbohydrates at breakfast help to reduce postprandial hyperglycemia, insulin responses and blood lipids. These studies had varying protocols and meal compositions; nevertheless, it seems that including proteins and fats and not only carbohydrates at breakfast may be beneficial for glycemic regulation.
Table 2. Effect of First meal of the Day on Glycemia in healthy individuals and subjects at high risk for developing or with type 2 diabetes.

| Study                           | Health Status | Age (Years) | Duration & Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention                                                                 | Selected Clinical Outcomes                                                                 |
|---------------------------------|---------------|-------------|-------------------------------------------|-------------|-----------------------|--------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------|
| Morris, et al., 2015 [263]      | Healthy       | 28 ± 9      | 2 weeks Within-participant cross-over     | 14          | Circadian alignment   | Alignement protocol had B at 8:00 AM                                                | +8% and +14% ppd AUC_{Glu} and ppd AUC_{Ins} at Dinner time                               |
|                                 |               | 25.4 ± 2.6  |                                           |             | Circadian misalignment (12 h shift)       | Misalignment protocol had “B” at 8:00 PM                                            | +3% ISR at Dinner time                                                                  |
|                                 |               |             |                                           |             | 2 weeks               | Isocaloric diets of 15–20% PRO, 45–50% CHO, 30–35% FAT                            | +12% and −27% ppd AUC_{Glu} and ppd AUC_{Ins} in biological evening                    |
|                                 |               |             |                                           |             | 2 weeks               | +8% ISR in biological evening                                                      | −21% Fasting Ins in biological evening                                               |
|                                 |               |             |                                           |             | 2 weeks               | +14% and +9% late phase Ins/ISR and 24 h Ins at circadian misalignment             | +14% and +9% late phase Ins/ISR and 24 h Ins at circadian misalignment               |
|                                 | Healthy lean  | 36 ± 11     | 6 weeks Randomized controlled trial       | 33          | B group               | B: ≥700 kcal before 11:00, Fasting group: extend o/n fast until 12:00, ad libitum intake for the rest of the day | Glu (mg/dL) +1.3 (fast) vs +1.1 (B)                                                 |
|                                 |               | 22.4 ± 2.2  |                                           |             | Fasting group         |                                                                                     | Ins (µIU/mL) +0.32 (fast) vs +0.35 (B)                                               |
|                                 |               |             |                                           |             |                       |                                                                                     | HOMA-IR +0.10 (fast) vs +0.10 (B)                                                    |
|                                 |               |             |                                           |             |                       |                                                                                     | C-ISI Matsuda index +0.38 (fast) vs −0.97 (B)                                         |
|                                 |               |             |                                           |             |                       |                                                                                     | Index of adipose insulin sensitivity (%) +3.3 (fast) vs +9.9 (B)                      |
|                                 |               |             |                                           |             |                       |                                                                                     | Peak Glu until 12:00 + 1.1 mmol/L in B vs fasting                                  |
|                                 |               |             |                                           |             |                       |                                                                                     | Mean morning Glu +0.3 mmol/L in B vs fasting                                   |
|                                 |               |             |                                           |             |                       |                                                                                     | Greater Glu variability in fasting group                                            |
Table 2. Cont.

| Study                        | Health Status | Age           | Duration & Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention                                                                 | Selected Clinical Outcomes                                                                 |
|------------------------------|---------------|---------------|------------------------------------------|--------------|-----------------------|---------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------|
| Chowdhury, et al., 2016 [283]| Obese         | 44 ± 10       | 33.7 ± 4.9                               | 6 weeks      | B group               | Fasting group: extend o/n fast until 12:00, ad libitum intake for the rest of day       | Fasting Glu (mg/dL) +1.7 (fast) vs +1.4 (B)                                                  |
|                              |               |               |                                          | Randomized controlled trial | Fasting group |                                                                      | Fasting Ins (µIU/mL) −0.62 (fast) vs +0.39 (B)                                                |
|                              |               |               |                                          | 23           |                       |                                                                                       | HOMA-IR −0.13 (fast) vs +0.018 (B)                                                        |
|                              |               |               |                                          |              |                       |                                                                                       | C-ISI Matsuda index −0.05 (fast) vs +0.05 (B)                                               |
|                              |               |               |                                          |              |                       |                                                                                       | Ins AUC Glu, mg-120 min/dL +171 (fast) vs −231 (B)                                          |
| Jakubowicz, et al., 2017 [285]| Healthy:      | 44.3 ± 2.9    | 23.1 ± 0.4                               | 2 test days  | YesB, NoB            | Each test meal:                                                                          | +15–18% AUC<sub>Glu</sub> after lunch w/o B                                               |
|                              |               |               |                                          | Randomized open-label crossover-within-subject clinical trial |            |                       | 572 ± 8 kcal                                                                          | −25% AUC<sub>Ins</sub> after L for T2DM grp w/o B                                        |
|                              |               |               |                                          | 32           |                       | 32% PRO                                                                                | −35% AUC<sub>iGLP-1</sub> after L on NoB                                                   |
|                              |               |               |                                          |              |                       | 49% CHO                                                                                 |                                                                                              |
|                              |               |               |                                          |              |                       | 19.4% FAT                                                                               |                                                                                              |
|                              | T2D:          | 66.8 ± 1.9    | 30.7 ± 1.1                               |              |                       |                                                                                       |                                                                                              |
|                              |               |               |                                          |              |                       |                                                                                       |                                                                                              |
| Nas, et al., 2017 [286]      | Healthy adults| 24.6 ± 3.3    | 23.7 ± 4.6                               | 3 test days  | Control (C)           | Isocaloric diets 55% CHO, 30% FAT, 15% PRO                                              | HOMA-IR 1.96 ± 0.82 (C), 2.07 ± 0.91 (BSD), 1.96 ± 1.05 (DSD)                            |
|                              |               |               |                                          | Randomized crossover nutritional intervention | (3 meals)    | BSD-washout-C-DSD or DSD-washout-C-BSD                                                 | Glycemia<sub>AUC</sub> (mg/dLx24 h): 2360 ± 111 (C), 2425 ± 131 (BSD), 2374 ± 165 (DSD) |
|                              |               |               |                                          | 17           | BSD (B skipping)      |                                                                                        | MAGE 3.90 ± 1.32 (C), 3.65 ± 1.52 (BSD), 3.28 ± 1.75 (DSD)                              |
|                              |               |               |                                          |              | DSD (D skipping)      |                                                                                        | C-peptide (µg/day) 74 ± 38 (C), 86 ± 40 (BSD), 75 ± 42 (DSD)                            |
|                              |               |               |                                          |              |                       |                                                                                        | iAUC<sub>Ins</sub> (µU/mLx 2 h) after L: 211 ± 74 (BSD), 144 ± 74 (DSD)              |
|                              |               |               |                                          |              |                       |                                                                                        | iAUC<sub>Glu</sub> (mg/dLx 2 h) after L: 114 ± 41 (BSD), 62 ± 40 (DSD)              |
|                              |               |               |                                          |              |                       |                                                                                        | HOMA-IR pp after L: 59 ± 44 (BSD), 27 ± 23 (DSD)                                      |
Table 2. Cont.

| Study                        | Health Status | Age (Years) | BMI (kg/m²)       | Duration & Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes                                                                 |
|------------------------------|---------------|-------------|-------------------|-------------------------------------------|--------------|-----------------------|---------------------|------------------------------------------------------------------------------------------|
| Kobayashi, et al., 2014 [287] | Healthy       | 25.3 ± 1.2  | BW 74.5 ± 4.3 kg  | 2 test days Randomized crossover           | 8            | B (3 meals) noB (2 meals) | PRO 364 ± 16 kcal CHO 1310 ± 77 kcal FAT 462 ± 33 kcal Individually adjusted meals of 2190 ± 124 kcal/day | +9 mg/dL Blood Glu after L in noB vs B group (p < 0.05) +10 mg/dL sleep Blood Glu in noB vs B group (p < 0.05) AUCGlu after L 409 ± 99 mg/dL min in B group vs 811 ± 101 mg/dL min in noB group AUCGlu after D 1049 ± 144 mg/dL min in B group vs 1196 ± 204 mg/dL min in noB group |
| Jakubowicz, et al., 2015 [301] | T2DM          | 56.9 ± 1.0  | 28.2 ± 0.6        | 2 test days Randomized, open-label, crossover-within-subject clinical trial | 22           | YesB (B, L, D) NoB (L, D) | Each test meal: 701 ± 8 kcal; 26% PRO, 54% CHO, 20% FAT, 7% fiber | NoB vs YesB after B: Glu (mg/dL min) −43%, Ins (µIU/mL·min) −72.1%, Glucagon (pg/mL·min) −20.6% C-peptide (ng/mL·min) −63.3%, iGLP−1 (pmol/L·min) −60.5% NoB vs YesB after L: Glu (mg/dL min) +39.8%, Ins (µIU/mL·min) −24.7%, Glucagon (pg/mL·min) +9.7% C-peptide (ng/mL·min) −13.6%, iGLP−1 (pmol/L·min) −21.5% NoB vs YesB after D: Glu (mg/dL min) +24.9%, Ins (µIU/mL·min) −10.8%, Glucagon (pg/mL·min) +8.5% C-peptide (ng/mL·min) −14.5%, iGLP−1 (pmol/L·min) −14.5% |
Table 2. Cont.

| Study                      | Health Status Age (Years) BMI (kg/m²) | Duration & Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|----------------------------|---------------------------------------|------------------------------------------|-------------|-----------------------|----------------------|---------------------------|
| Kang, et al., 2013 [84]    | Subjects with prediabetes and normal (NGR) Glu regulation 46.4 ± 13.8 18.5–24.9 | 3 days Cross-sectional study | 133         | LC (low CHO) MC (medium CHO) HC (high CHO) | Diet of 30 kcal/kg/day calorie intake from three daily meals According to CHO in B: Low carbohydrate (LC) meal with <45%CHO Medium-carbohydrate (MC) meal with CHO between 45% and 65% High-carbohydrate (HC) meal with >65% CHO | In subjects with impaired Glu regulation: Significantly ↑ ppd Glu, Glu peak, Glu excursion, and iAUCGLU in subjects with impaired Glu regulation after B with >50% CHO |
| Rosi, et al., 2018 [282]   | Healthy 24 ± 2 23.4 ± 1.6 | 7 weeks Randomized, crossover, and controlled trial | 15          | F-CTRL BR-BREAD BR-MUESLI BR-RICE | Energy-free meal with a cup of decaf coffee (~fasting) 3 isoenergetic meals with similar PRO: with a cup of semi-skimmed milk, an apple, and cereal foods as indicated below: White bread with chocolate hazelnut spread, GI <55, GL~22 Muesli with dark chocolate chips and nuts, ↑fiber, GI <55, GL~23 Chocolate-flavored puffed rice, ↓FAT, ↑CHO, GI >55, GL~38 | The RICE group had significantly higher: -AUCIns 120 min after B -AUCGlu 120 min after B -Plasma Glu after B |
Table 2. Cont.

| Study | Health Status Age (Years) | BMI (kg/m²) | Duration & Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|-------|---------------------------|-------------|-------------------------------------------|-------------|-----------------------|----------------------|---------------------------|
| Jakubowicz, et al., 2017 [296] | T2DM 59.0 ± 0.7 | 32.11 ± 0.1 | 12 weeks Randomized, open-label, parallel-arm clinical trial | 48          | 42 g total PRO: WBdiet (whey, 28 g) PBdiet (42 g various PRO sources) CBdiet (high CHO B, 17 g PRO from various sources) | At B: WBdiet: 25% PRO (mainly whey), 50% CHO, 25% FAT PBdiet: 25% PRO (mainly from eggs, tuna, soy), 50% CHO, 25% FAT CBdiet: 11% PRO (soy), 64% CHO, 29% FAT Hypocaloric diets: B 660 ± 25 kcal L 560 ± 20 kcal (23% PRO, 48% CHO, 29% FAT) D 280 ± 15 kcal (31% PRO, 31% CHO, 38% FAT) B at 6:00–8:30 h, L at 12:30–14:30 h, D at 18:30–20:30 h | HbA1C (%) WB −0.89 ± 0.05 PB −0.6 ± 0.04 CB −0.36 ± 0.04 Fasting Glu (mmol/L) WB −0.73 ± 0.06 PB −0.43 ± 0.06 CB −0.12 ± 0.04 Overall glycemia was −12% in PB and −19% in WB Glu peak was −18% in PB and −31% in WB Rapid Glu levels decrease after B in PB and WB Overall AUC Ins was +37% in PB and 62% in WB (same for after L and D) AUC C-pept was +53% in PB and 96% in WB (same for after L and D) AUC iGLP_1 was +70% in WB and +33% in PB after B, L, and D HbA1C reduced in all grps |
Table 2. Cont.

| Study | Health Status | Age (Years) | BMI (kg/m²) | Duration & Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|-------|---------------|-------------|-------------|-------------------------------------------|-------------|------------------------|----------------------|-----------------------------|
| Jakubowicz, et al., 2012 [297] | 20–65 | 32.3 ± 1.8 | 32 weeks | Randomized, treatment controlled, open clinical trial | 144 | Lcb | Low kcal and low CHO diet (Lcb) with low kcal and low HCO B | BW (kg): 70.6 ± 8.7 (HCPb) vs 86.9 ± 9.7 (Lcb) | Fasting Glu (mg/dL): 84.2 ± 4.6 (HCPb) vs 95.5 ± 4.9 (Lcb) |
| | | | | | | | High CHO and high PRO diet (HCPb) with daily dessert for B | Fasting Ins (µU/mL): 8.9 ± 3.9 (HCPb) vs 23.69 ± 3.8 (Lcb) | HOMA-IR: 1.6 ± 0.4 (HCPb) vs 5.9 ± 0.9 (Lcb) |
| | | | | | | Similar L & D composition, differences for B | Total Chol (mg/dL): 179.2 ± 11.1 (HCPb) vs 190.8 ± 18.2 (Lcb) | TG (mg/dL): 122.6 ± 9.7 (HCPb) vs 174.5 ± 20.9 (Lcb) |
| Neumann, et al., 2016 [298] | Healthy | 24.1 ± 2 | n/a | 8 days | Randomized, controlled study | 24 | SKP (Skipping breakfast) | CHO group: 351 kcal; 59 g CHO, 10 g PRO, 8 g fat | No difference in fasting blood Glu CHO and PRO groups lead to greater ppd Glu vs SKP |
| | | | | | | | PRO group: 350 kcal; 39 g CHO, 30 g PRO, 8 g Fat | ↓10% Glu in PRO vs CHO at 30 min ppd after B |
| Pedersen, et al., 2016 [299] | Obese/T2DM | 63.9 ± 2.15 | 33 ± 1.25 | 4 exp. days | Randomized crossover study | 28 | CHO-B | Fast ≥ 8 h before the test diets | Peak blood Glu (mmol/L): 11.3 ± 0.5 after CHO-B |
| | | | | | | | noCHO-B | 9.4 ± 0.4 after noCHO-B | Mean blood Glu (mmol/L)—5 h after B |
| | | | | | | | The diets were consumed on 2 sequential days, 3 identical meals with CHO 3 meals, no CHO breakfast, lunch and dinner with CHO—similar to other group | 8.4 ± 0.5 after CHO-B |
| | | | | | | | no sig. differences in Glu measurements after lunch and dinner | 7.5 ± 0.4 after noCHO-B | no sig. differences in gastric emptying |
Table 2. Cont.

| Study                                      | Health Status Age (Years) | Duration & Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|--------------------------------------------|----------------------------|-------------------------------------------|-------------|----------------------|----------------------|---------------------------|
| Rabinovitz, et al., 2014 [300]              | Overweight/obese with T2DM | 3 months                                  | 46          | SB (small breakfast)  | At B: 12–18% PRO, 14–22% FAT, 60–70% CHO, 13% of total E was recommended in the SM, Lunch and Dinner had 33% of total daily E, 2–3 snacks same as in BB | BW no sign. difference HbA1c (%) −0.58 ± 0.18 in BB, −0.13 ± 0.08 in SB Estimated average glucose (mg/dL) −16.6 ± 5.2 in BB, −3.43 ± 2.4 in SB no sign. changes in Glu, Ins, C-peptide, total Chol, TG, CRP, IL-6, TNF-α |
|                                            | 60.7 ± 6.35                | Randomized, treatment-controlled, open clinical trial |             | BB (big breakfast)   | At B: 23–30% PRO, 29–37% FAT, 37–48% CHO 33% of total E was recommended in the BB, Lunch and Dinner had 25% of total daily E, 2–3 snacks same as in SM |
|                                            | 32.37 ± 3.7                |                                           |             |                      |                      |                           |

Abbreviations: B: Breakfast; L: Lunch; D: Dinner; YesB: had breakfast; NoB: skipped breakfast; T2DM: type 2 diabetes; ppd: postprandial; GI: Glycemic Index, GL: Glycemic Load; BSD: breakfast skipping day; DSD: dinner skipping day; PRO: protein; CHO: carbohydrates; FAT: fat/lipids; HOMA-IR: homeostatic model assessment for insulin resistance; AUC: area under the curve; iAUC: incremental area under the curve; Ins: insulin concentrations; Glu: glucose concentrations; HbA1c: glycated hemoglobin A1c; ISI: insulin sensitivity index; WB: body weight; E: total energy intake; SKP: skipping breakfast. An arrow pointing upwards or downwards indicates an increase or decrease.
Two studies from our group, an acute [302] and a short-term (8 weeks duration) [143] cross-over RCT, and a 2013 meta-analysis of studies ranging from 4–24 weeks, reported that high-protein eating plans (25–32% of total energy vs 15–20%) resulted in 2 kg greater weight loss, better maintenance of muscle mass, and 0.5% greater improvements in HbA1c, without significant decreases in fasting glucose [303]. However, it should be noted that the protein sources that seem to have beneficial effects on postprandial hyperglycemia and IR are possibly either lean animal proteins or plant proteins, although this remains to be elucidated. Similar contradictory results on postprandial hyperglycemia and IR have been reported in respect to fats, with some supporting monounsaturated fats, such as extra-virgin olive oil and nuts [103], others supporting saturated fat from dairy products, coconut oil and palm kernel oil [304] and others reporting that saturated fat intake was associated with higher risk of T2DM [305].

4.2.2. The Effects of Consuming Most Food and Calories Earlier in the Day on Glycemia

Due to the daily oscillations of various hormones and the two-way interaction between food consumption and metabolism, a lot of attention has been given on what the energy and food intake distribution in a day should be, the number of meals consumed, the consistency of meal timing, and the time of day that these meals should be consumed, in order to ameliorate postprandial hyperglycemia and IR. Meal timing has been reported to synchronize and boost the peripheral circadian clocks that control downstream metabolic pathways (Figure 3) [306]. Earlier efforts using mixed meals or glucose infusion demonstrated circadian responses of reduced glucose tolerance and insulin sensitivity in healthy participants for the evening hours rather than in the morning [260]. So, the same foods distributed differently throughout the day, appear to have different effects on glycemic control in subjects with prediabetes and/or T2DM. Recently more studies have been conducted to address this issue. Table 3 describes the effects of lunch and dinner on indices of glycemic control. Having a caloric rich breakfast was found to result in increased postprandial insulin secretion and GLP-1 responses, and smaller plasma glucose peaks after breakfast consumption [301]; these effects were not present when the majority of calories was consumed at the evening dinner. The effects of the earlier increases in plasma insulin levels persisted after lunch aiding the glycemic management of the subsequent meals (the “second meal effect”). Results from a 7-days/randomized/open-label/crossover trial with 18 subjects with T2DM, performed in two separate testing days, each over the course of 14 h, showed that eating the majority of calories at breakfast (at 08:00) vs dinner (at 19:00) at home for 6 days before each testing day (30–50% of daily calories at breakfast, i.e., 704 kcal breakfast, 605 kcal lunch and 205 kcal dinner vs 205 kcal breakfast, 605 kcal lunch and 704 kcal dinner), with the “large” meal (breakfast or dinner respectively, containing 22% fat, 47% protein), or the “small” meal (dinner or breakfast, respectively, with 30% fat, 27% carbohydrates, 43% protein), led to reduction in overall postprandial hyperglycemia [301]. Results from another study with 12 healthy young females examining the effects of a late suppertime (18:00 vs 23:00) on gastrointestinal activity the following morning (efficiency of digestion/absorption of dietary carbohydrates ingested at the usual suppertime), showed that a late supper was associated with a worse effect on postprandial glucose profiles the following morning [307]. A pertinent question arising is how the body recognizes calories. Maybe it doesn’t in the short term in which case a high calorie breakfast is a meaningless way to describe a meal being investigated for acute metabolic effects.

On the other hand, skipping the evening dinner (Table 3) may improve glucose intolerance, and lower IR [286,308]. In contrast, eating a rich or a late dinner may aggravate IR and hyperglycemia and deteriorate blood glucose levels the following morning [307,309]. It has also been shown that moving the evening dinner to an earlier time may improve glucose tolerance due to a causal role of endogenous melatonin in the impairment of glucose tolerance, particularly in MTNR18 carriers [310]. Unfavorable effects on postprandial glycemia and overall glycemic control have been reported during Ramadan fasting, where people abstain from eating/drinking during daylight and consume all energy at night [311].
Likewise, consuming lunch later rather than early (after 15:00–16:00 h) in the day, has also been associated negatively with glycemic control (Table 2). Eating lunch late increased IR (higher HOMA-IR) in overweight and obese subjects [312] and increased postprandial blood glucose in healthy individuals [309]. Also, one acute study showed that consuming a snack in the afternoon, but not right after lunch, improved the mean amplitude of glycemic excursions [313]. Furthermore, consuming more carbohydrates at lunch time (Table 4) and more fats later rather than earlier, reduced fasting blood glucose, insulin, and GLP-1 responses. Whole day blood glucose fluctuations were also maintained lower when carbohydrates were consumed earlier than later in the day [314]. Additionally, a short-term trial in healthy individuals demonstrated that consumption of high GI foods was better managed during daytime hours, whereas high GI food intake at dark hours (late afternoon-evening hours) resulted in greater blood glucose peaks and total blood glucose concentrations after meals [315]. Together, these data suggest that carbohydrates should be consumed primarily at lunch and early afternoon hours. Early in the day is a fasting state in which the glycemic responses to carbohydrates are high (poor due to IR), whereas later in the day glycemic responses are better (lower) since preceding meals may sensitize the metabolic and incretin systems to the following ones, thereby improving glucose tolerance [31]. In this regard, although some carbohydrates must be consumed at breakfast to improve glucose tolerance in the following meals, placing the majority of carbohydrates at breakfast may not be a good idea; actually, breakfast and evening dinner should have the minority of carbohydrates with lunch taking the most. Such effect is consistent with the circadian biology of glucose metabolism and does support that following dietary patterns in accordance with the diurnal bodily rhythms can be a useful approach in the management of glycemia. Moreover, consuming most of the food and calories at lunch and early afternoon hours, and not during the night hours may be beneficial for satiety and hunger regulation. Hunger is intrinsically modulated and has circadian periodicity with peaks in the evening, meaning that there is higher tendency for food later in the day. Ghrelin, an orexigenic hormone, increases just before a meal and especially at night [316], but is being suppressed by increases in insulin. Therefore, having a large, rich breakfast may also contribute to hunger, cravings, and postprandial ghrelin reduction [266].
Table 3. Effect of Lunch and Dinner on glycemia in healthy and individuals at high risk for developing or with type 2 diabetes.

| Study                      | Health Status Age (Years) BMI (kg/m²) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|----------------------------|---------------------------------------|---------------------------------------------|-------------|------------------------|----------------------|--------------------------|
| Effect of Lunch on Glycemia|                                       |                                             |             |                        |                      |                          |
| Bandin, et al., 2015 [309] | Healthy 26 ± 4 22.54 ± 2.05           | 2 weeks Randomized and crossover (Protocol 1: metabolic study) | 10 (subjects on protocol 1) | EE group (Early Eating) LE group (Late Eating) | L at 13:00 h, L at 16:30 h Same B (at 8:00 h), D (at 20:00 h), and L as indicated | 1868 ± 234 Kcal/day 15% PRO, 50% CHO, 35% FAT +46% AUC\textsubscript{Glu} after L in LE vs EE +1 mmol/L Glu 90 min after L in LE vs EE +0.6 mmol/L Glu 120 min after L in LE vs EE |
| Garaulet, et al., 2013 [312] | Overweight/obese 42 ± 11 31.4 ± 5.4 | 20 weeks                                    | 420         | Early Lunch Eaters (EL) Late Lunch Eaters (LL) | Early eaters: Lunch before 15:00 h Late eaters: Lunch after 15:00 h Weight loss diet of similar composition Total E ~ 1400 Kcal/day 19% PRO 48% CHO 33% FAT | Fasting Glu (mg/dL) 81.28 ± 15.97 (EL) vs 83.65 ± 16.27 (LL)—non-sign Fasting Ins (mU/L) 5.72 ± 4.71 (EL) vs 6.98 ± 11.66 (LL)—non-sign HOMA 1.17 ± 0.14 (EL) vs 1.57 ± 0.13 (LL)—significant |

Effect of Dinner on Glycaemia

- Dinner timing & composition
Table 3. Cont.

| Study                        | Health Status | Age (Years) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|------------------------------|---------------|-------------|---------------------------------------------|-------------|------------------------|----------------------|---------------------------|
| Nas, et al., 2017 [286]      | Healthy       | 24.6 ± 3.3  | 3 test days Randomized crossover nutritional intervention | 17          | Control (C) (3 meals) BSD (B skipping) DSD (D skipping) | Isocaloric diets 55% CHO, 30% FAT, 15% PRO BSD-washout-C-DSD or DSD-washout-C-BSD | HOMA IR 1.96 ± 0.82 (C), 2.07 ± 0.91 (BSD), 1.96 ± 1.05 (DSD) Glycemia $\text{AUC}_{\text{glu}}$ (mg/dLx24 h) 2360 ± 111 (C), 2425 ± 131 (BSD), 2374 ± 165 (DSD) MAGE 3.90 ± 1.32 (C), 3.65 ± 1.52 (BSD), 3.28 ± 1.75 (DSD) C-peptide (µg/day) 74 ± 38 (C), 86 ± 40 (BSD), 75 ± 42 (DSD) $i\text{AUC}_{\text{ins}}$ (µU/mLx2 h) after L 211 ± 74 (BSD), 144 ± 74 (DSD) $i\text{AUC}_{\text{glu}}$ (mg/dLx2 h) after L 114 ± 41 (BSD), 62 ± 40 (DSD) HOMAapp after L 59 ± 44 (BSD), 27 ± 23 (DSD) |
Table 3. Cont.

| Study | Health Status Age (Years) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|-------|---------------------------|---------------------------------------------|-------------|-----------------------|---------------------|-----------------------------|
| Jakubowicz, et al., 2015 [296] | T2DM 57.8 ± 4.7  28.1 ± 2.9 | 5 mo, 14 exp. days Randomized open-label crossover-within-subject clinical trial | 18 | Bdiet (HE Breakfast) Ddiet (HE Dinner) | Bdiet: B—2946 kJ, 31% PRO, 47% CHO, 22% FAT  L—2523 kJ, 27% PRO, 50% CHO, 23% FAT  D—858 kJ, 43% PRO, 50% CHO, 23% FAT  Ddiet: B—858 kJ, 43% PRO, 50% CHO, 23% FAT  L—2523 kJ, 27% PRO, 50% CHO, 23% FAT  D—2946 kJ, 31% PRO, 47% CHO, 22% FAT  Composition of diets: 6276 ± 105 kJ  31% PRO  46% CHO  23% FAT  B at ~08:00 h  L at ~13:30 h  D at ~19:30 h | −20% total day AUC$_{Glu}$ for Bdiet vs Ddiet  +20% total day AUC$_{Ins}$ for Bdiet vs Ddiet  +10% total day integrated AUC$_{Ins}$ for Bdiet vs Ddiet  −24% peak Glu (mmol/L × min) at 180 min after HE B vs HE D  +10–19% peak Ins (pmol/L × min) at 30–180 min after HE B vs HE D  Faster Ins peak (60 min) after B in Bdiet  +17% C-peptide (nmol/L × min) after HE B vs HE D  +35% IGLP-1 (pmol/L × min) at 30 min after HE B vs HE D  27% IGLP-1 (pmol/L × min) at 30 min after HE B vs HE D  −13–25% Glu (mmol/L × min) after L in Bdiet vs Ddiet  50% higher, and more rapid early prandial Ins in Bdiet after L |
| Study                  | Health Status | Age (Years) | BMI (kg/m²) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups                                                                 | Dietary Intervention                                                                 | Selected Clinical Outcomes                                                                 |
|-----------------------|---------------|-------------|-------------|---------------------------------------------|-------------|----------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------|
| Grant, et al., 2017   | Healthy       | 25 ± 5.4    | 22.2 ± 1.6  | 4 days Controlled, parallel study           | 11          | Eating at night group/condition (EN) Not eating at night group/condition (NoEN)         | Meals at 19:00 h and 01:30 h B was the tolerance test: ↑CHO (↑GI) at 06:30–07:00 h | EN group: +27% AUC_{Glu} on SW4 +69% AUC_{Glu} on RTDS +11% AUC_{Ins} on SW4 +35% AUC_{Ins} on RTDS NoEN group: +12% AUC_{Glu} on SW4 +2% AUC_{Glu} on RTDS +18% AUC_{Ins} on SW4 +16% AUC_{Ins} on RTDS |
|                       |               |             |             |                                             |             | Tolerance test and measurement were done on 4 different days: PRE—day before the start of the protocol, SW4—days of stimulated night work (sleep between 10:00–16:00 h for all subjects), RTDS—return to daytime schedule | Fasting Glu: No significant effects of condition (p = 0.522), day (p = 0.539) or condition × day (p = 0.228) Fasting Ins: No significant effects of condition (p = 0.380), day (p = 0.056) or condition × day (p = 0.958) for fasting glucose and insulin, respectively |
Table 3. Cont.

| Study | Health Status Age (Years) | BMI (kg/m²) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|-------|--------------------------|-------------|---------------------------------------------|-------------|-----------------------|----------------------|---------------------------|
| Lopez-Minguez, et al., 2018 [310] | Overweight/obese | 42 ± 10 | 2 exp. days, 1 week Randomized, cross-over trial | 40 | LE group (Delayed dinner or Late Eating) EE group (Advanced dinner or Early Eating) Also divided groups in MTNR1B risk carriers (GG) and non-risk carriers (CC) | LE: D at 23:00 h, L at 15:00 h, B at 8:00 h EE: D at 20:00 h, L at 12:00 h, B at 8:00 h Fixed menu for all meals on exp. days L was 8 h before D, with energy content of 650 Kcal D was 30–35% of total energy intake, composed of: 15–17% PRO 58–60% CHO/25–27% FAT | In total population: AUC_<sub>Glu</sub> (mmol/L x h) = 284.74 ± 32.67 (LE) vs 269.61 ± 34.8 (EE) Among GG group: AUC_<sub>Glu</sub> (mmol/L x h) = 292.2 ± 33.8 (LE) vs 270.9 ± 30.4 (EE) Among CC group: No significant effect AUC_<sub>Glu</sub> (mmol/L x h) = 277.3 ± 30.5 (LE) vs 268.2 ± 38.2 (EE) Significant interaction between meal timing (EE vs LE) and genotype (GG vs CC) for AUC_<sub>Glu</sub> |
| Jakubowicz, et al., 2013 [317] | Overweight/obese | 45.8 ± 7.1 | 12 weeks Randomized open-label parallel-arm trial | 74 | B group D group | B group (700 kcal B, 500 kcal L, 200 kcal D) D group (200 kcal B, 500 kcal L, 700 kcal D) ~1400 kcal weight loss diets, same macronutrient content and composition B at 6:00–9:00 h, L at 12:00–15:00 h, D at 18:00–21:00 h | Fasting Glu −11.5% (B) vs −4.2% (D) Fasting Ins −51% (B) vs −29% (D) HOMA-IR −57% (B) vs −32.5% (D) HOMA-B −25% (B) vs −17% (D) ISI +163% (B) vs +56% (D) AUC_<sub>Glu</sub> −22% (B) vs −15% (D) AUC_<sub>Insl</sub> −58% (B) vs −30% (D) |

Abbreviations: T2DM: type 2 diabetes; B: Breakfast; L: Lunch; D: Dinner; E: total energy intake; ppd: postprandial; GI: Glycemic Index; Glu: Glucose; Ins: Insulin; AUC: Area Under the Curve; IAUC: incremental area under the curve; CHO: carbohydrates; PRO: proteins; FAT: fats; HOMA: homeostatic model assessment for insulin resistance; ISI: insulin sensitivity index. An arrow pointing upwards indicates high carbohydrates and high GI.
In conclusion, the recommendations for persons wanting to lose weight may be that the fasting interval should be prolonged to 16 h, which prolongs the release of NEFAs and fat oxidation. A late large evening dinner shortens the fasting interval increasing the possibility of morning hyperglycemia. This is relevant also to a potential role of ketones produced during fasting in suppressing hunger and decrease food intake. The oxidation of fat and ketones may spare glucose to maintain a reasonable fasting level in blood to avoid metabolic alarm. More RCTs are needed in this field to identify the optimal meal and snack timing, whether this changes according to health status, such as in diabetes, and which are the exact mechanisms for amelioration of postprandial hyperglycemia and IR.

4.2.3. Effects of Meal Frequency on Postprandial Glycemia and IR

Several studies suggest that more frequent meals increase weight gain due to postprandial fat deposition [318,319], thereby worsening hyperglycemia, IR, hyperlipidemia, and appetite [319,320]. In contrast, others support the idea that frequent meals may reduce body weight and produce lower postprandial glycemic/insulinemic responses, lower blood lipids, improve metabolic control, and reduce appetite [321,322]. Those favoring many (5–6) regular smaller meals/day support that this dietary behavior diminishes glucose fluctuations/swings and provides a steadier delivery of nutrients throughout the day, thereby inducing lower glycemic loads and delayed gastric emptying resulting in less insulin requirements for glucose control, decrease in postprandial glucose levels, and in some cases reduction of hunger and the desire to eat [322–325]. Those favoring few (2–3) regular larger meals per day support that this dietary habit is more in line with human’s natural inclination to eat more in the morning and fast in the evening and during the night, and that it improves the expression of biological clocks regulating glucose metabolism and body weight [320,326]. In addition, some claim that having more meals per day can increase the consumption of unhealthier foods and added sugars leading to adverse effects in body weight, glycemia, and lipidemia [327]. Table 4 and Figure 3 describe the effects of meal frequency and energy and macronutrient distribution on postprandial hyperglycemia.

Clinical trials, both short-term (14–28 days) [321,325,328–332] and long-term [323,324] with no caloric restriction in non-obese and obese women with PCO syndrome at an early or late stage IGT or overt early-stage T2DM, and long-term studies with caloric restriction in T2DM without [333] or with antidiabetic treatment [320,326], have produced contradictory results regarding the association of meal frequency with body weight, postprandial glycemia, IR and overall glycemic control. Out of the only four long-term clinical trials investigating the effects of meal frequency in T2DM, the three with caloric restrictive diets provided contradictory results: (a) in T2DM subjects receiving anti-diabetic medications, 2 large vs 6 smaller meals decreased body weight, fasting plasma glucose/C-peptide/glucagon levels, with no differences in HbA1c, insulin, insulin sensitivity and blood lipids [320]. (b) In contrast, another study in T2DM reported that 5 vs 3 meals per day resulted in decreased BMI and HbA1c, without significant differences in fasting plasma glucose, insulin, and lipids [333]. (c) In uncontrolled subjects with diabetes treated with insulin, 3 vs 6 meals per day resulted in decreased body weight, HbA1c, total daily insulin requirements and higher expression of clock genes [326]. However, it is well known that both weight loss and antidiabetic medications have a significant impact on glucose/lipid metabolism [334], making it difficult to determine whether the beneficial effects were attributed to the medications, energy deficit or meal frequency.
Table 4. Effect of Meal, energy, and nutrient distribution in a day on glycemic control in individuals at high risk for developing or with type 2 diabetes.

| Study | Age (Years) | BMI (kg/m²) | Health Status | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|-------|-------------|-------------|---------------|---------------------------------------------|-------------|------------------------|----------------------|--------------------------|
| Kahleova, et al., 2014 [320] | T2DM | 59.4 ± 7.0 | 32.6 ± 4.9 | 24 weeks each regiment Randomized, open, crossover study | 54 | A6 regiment (6 meals/day) B2 regiment (2 meals/day) | B, L, D, and 3 smaller snacks in between B and L | Caloric restriction of 500 kcal/day 50–55% CHO, 20–25% PRO, <30% FAT (≤7% SFAs, <200 mg/day of Chol), and 30–40 g/day of fibers |

- Effect of the Number of meals per day on Glycemia

- BW (signif.)
  - −2.3 kg in A6
  - −3.7 kg in B2
  - HbA1c
  - −0.23% in A6
  - −0.25% in B2

- Fasting plasma Glu (mmol/L)
  - −0.47 in A4
  - −0.78 in B2

- Fasting immunoreactive Ins (pmol/L)
  - −0.69 in A6
  - −0.75 in B2

- Ins secretion at reference level (pmol min⁻¹ m⁻²)
  - +22.9 in A6
  - +20 in B2

- Glu sensitivity (pmol min⁻¹ m⁻² mmol⁻¹ L⁻¹)
  - +5.8 in A6
  - +5.9 in B2

- TG (mmol/L)
  - −0.28 in A6
  - −0.17 in B2

(all above changes per group were significant)
### Table 4. Cont.

| Study                                    | Age (Years) | BMI (kg/m²) | Health Status | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention                                                                 | Selected Clinical Outcomes                                      |
|-----------------------------------------|-------------|-------------|---------------|---------------------------------------------|-------------|-----------------------|---------------------------------------------------------------------------------------|---------------------------------------------------------------|
| Papakonstantinou et al., 2018 [324]     | 2 IGT groups (early and advanced stage) and T2DM | 49.3 ± 1.8 | 32.4 ± 0.8    | 24 weeks Randomized, crossover study         | 47          | IGT-A (PG 140–199 mg/dL at 120 min post OGTT) | Weight maintenance diet: 1900 kcal/day, 45% CHO, 20% PRO, 35% FAT 6 meals/day (B, L, D, and 3 snacks; with CHO: 20% at B, 10% morning snack, 30% at L, 10% at afternoon snack, 20% at D and 10% at bedtime snack) Or 3 meals/day (B, L, and D; with CHO: 20% at B, 50% at L, and 30% at D) | T2D group: ▼▼ post-OGTT Glu and ▼▼ HbA1c with 6 meals  IGT-A group: ▼ 30-min and ▼▼ 60-min post-OGTT plasma Ins with 6 meals  IGT-B group: ▼ peak Glu with 3 meals ▼▼ peak Glu with 6 meals In all groups: ▼ subjective hunger with 6 meals no differences in satiety or lipids no differences in FPG, Glu or Ins iAUC, fasting Ins, HOMA-IR with 3 vs 6 meals |
|                                        | T2DM ≥25 years | 32.4 ± 5.2  | HbA1c: 8.1 ± 1.1% T2DM for ≥5 yrs, treated with insulin ≥1 yr with >25 units for at least 3 months | 3 months Randomized, parallel, treated with insulin, continuous glucose monitoring | 28          | 6 meals | Isoenergetic diets consisting of 3 or 6 meals/day: 3M 700 kcal breakfast, 600 kcal lunch, 200 kcal dinner; 6M same as 3M and addition of 150 kcal snacks | 12 weeks with 3 meals/day vs 6 meals/day – 5.4 kg weight loss, –1.2% total insulin dose—26 units, higher clock gene expression |

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Table 4. Cont.

| Study                  | Age (Years) | BMI (kg/m²) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|------------------------|-------------|-------------|--------------------------------------------|-------------|-----------------------|----------------------|----------------------------|
| Arnold, et al., 1997 [331] | T2DM or IGT | 46–70       | 8 weeks Randomized, crossover study         | 13          | 3 meal regimes         | Isoenergetic diets consisting of 3 or 9 meals/day; 4 weeks with 3 meals/day; Daily E needs: 25% at B, 25% at L, ~50% at D, and ~150 kcal at a snack | Glu (mmol/L) +2% with 3 meals; +4% with 9 meals; Ins (µU/mL) +1% with 3 meals; −2% with 9 meals; Total Chol (mmol/L) +3.5% with 3 and with 9 meals; TG (mmol/L) −13.5% with 3 and with 9 meals; ApoB (mg/dL) +12% with 3 meals; 17% with 9 meals |
| Salehi, et al., 2014 [333] | T2DM       | 35–65       | 3 months RCT                               | 66          | 6-meal group           | Weight loss diets (−300 kcal/day); 6 isocaloric meals; 3 large meals and 2 small snacks (Ctrl) 56% CHO, 16% PRO and 28% FAT | ↓↓ BMI in 6 M; ↓ BMI in usual pattern; ↓↓ HbA1c in 6 M; ↓↓ Ins and 2 h-ppd Glu in 6 M; ↓↓ Ins in usual pattern; no sign. differences in fasting Glu, fasting Ins, and 2 h-ppd serum Glu in both groups |

- Effect of the Distribution of Energy and Macronutrients in a day on Glycaemia
| Study | Age (Years) | BMI (kg/m²) | Health Status | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|-------|-------------|-------------|---------------|---------------------------------------------|-------------|-----------------------|----------------------|---------------------------|
| Pearce, et al., 2008 [82] | T2DM | 61.3 ± 10 | 34.7 ± 9 | 3 days Randomized crossover study | 23 | CARB-E CARB-B CARB-L CARB-D | Even CHO distribution is all meals/day ~70 g CHO CHO mainly in B (~125 g) CHO mainly in L (~125 g) CHO mainly in D (~125 g) 40% CHO, 34% PRO, 26% FAT | Glucose max (mmol/L): 14.2 ± 1.0 CARB-L 14.5 ± 0.9 CARB-E 14.6 ± 0.8 CARB-D 16.5 ± 0.8 CARB-B Glu AUC₂₀₀ (mmol/L·20 h): 10,049 ± 718 CARB-L 10,493 ± 706 CARB-E 10,717 ± 638 CARB-D 10,603 ± 642 CARB-B small but no sig. difference in fasting blood Glu |
| Jakubowicz, et al., 2015 [301] | T2DM | 57.8 ± 4.7 | 28.1 ± 2.9 | 5 mo, 14 exp. days Randomized open-label crossover-within-subject clinical trial | 18 | Bdiet (HE breakfast) Ddiet (HE diner) | Bdiet: B—2946 kJ, 31% PRO, 47% CHO, 22% FAT L—2523 kJ, 27% PRO, 50% CHO, 23% FAT D—858 kJ, 43% PRO, 50% CHO, 23% FAT Ddiet: B—858 kJ, 43% PRO, 50% CHO, 23% FAT L—2523 kJ, 27% PRO, 50% CHO, 23% FAT D—2946 kJ, 31% PRO, 47% CHO, 22% FAT 6276 ± 105 kJ 31% PRO 46% CHO 23% FAT B at 08:00 h L at 13:00 h D at 19:00 h | −20% daily AUC₆₇₆₈ for Bdiet −24% AUC₆₇₆₈ in Bdiet after B Faster plasma Glu level decrease after B +11% AUC₆₇₆₈ in Bdiet after B +12% Ins peak after B in Bdiet Faster Ins peak (60 min) after B in Bdiet −21–25% AUC₆₇₆₈ in Bdiet after L +23% AUC₆₇₆₈ in Bdiet after L 50% higher, and more rapid early prandial Ins in Bdiet after L |
| Imai, et al., 2018 [313] | T2DM | 67.4 ± 9.4 | 23.5 ± 3.1 | 4 days Randomized, crossover clinical trial | 17 | Group 1: Day 2 snack at 12:30 h and day 3 snack at 15:30 h Group 2: Day 2 snack at 15:30 h and day 3 snack at 12:30 | Diet: Same meals in the 2 groups with B at 07:00 h, L at 12:00 h, D at 19:00 h and snack at either 12:30 h (just after lunch—early) or at 15:30 h (mid-afternoon-late) | MAGE (mmol/L): 6.90 ± 0.69 with early snack 5.19 ± 0.48 with late snack iAUC (mmol/L per min⁻¹) 1030 ± 180 with early snack 701 ± 97 with late snack Time of snack did not affect the mean Glu level |
Table 4. Cont.

| Study                        | Age (Years) BMI (kg/m²) Health Status | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|------------------------------|---------------------------------------|---------------------------------------------|-------------|-----------------------|----------------------|--------------------------|
| Kessler, et al., 2017 [314]  | NGT and IGT 45.9 ± 2.5 27.1 ± 0.8     | 8 weeks Crossover trial                     | 29          | HC/HF                 | HF/HC                | Isocaloric diets, with energy intake equally distributed in the day HC/HF for 4 weeks: CHO-rich meals until 13:30 h, and FAT-rich meals 16:30 h–22:00 h HF/HC for 4 weeks: FAT-rich meals until 13:30 h, and CHO-rich meals 16:30 h–22:00 h. | In subjects with impaired Glu tolerance and fasting Glu:  
  Fasting Glucose  
  − 11.4% in HC/HF  
  − 9.6% in HF/HC  
  Fasting Insulin  
  − 21.9% in HC/HF  
  − 27.1% in HF/HC  
  Fasting C-peptide  
  − 42.6% in HC/HF  
  − 50.6% in HF/HC  
  HOMA-IR  
  − 33.8% in HC/HF  
  − 34.7% in HF/HC  
  Fasting GLP-1  
  − 45% in HC/HF  
  − 13.3% in HF/HC  
  Whole day Glu  
  +7.9% in HF/HC vs HC/HF |
| Gibbs, et al., 2014 [315]    | Healthy 25.5 ± 8.8 21.9 ± 1.7         | 4 exp. days Randomized, crossover study     | 10          | Low GI meals (LG)     | LG: GI~37            | Glu peak (mmol/L)  
  7.8 ± 0.4 with LG & HG am  
  8.3 ± 0.2 with LG pm  
  9.54 ± 0.4 with HG pm  
  Glu 2 h-ppd (mmol/L)  
  4.85 ± 0.2 with LG & HG am  
  6.42 ± 0.4 with LG & HG am | Glu peak (mmol/L) but a trend for ↓ iAUC_Glu with am meals |
|                             |                                       |                                             |             | High GI meals (HG)    | HG: GI~73            | no sig. differences in iAUC_Glu |
|                             |                                       |                                             |             |                       |                      | no sig. differences in iAUC_Ins |

Abbreviations: IGT: impaired glucose tolerance; T2DM: type 2 diabetes; NGT: normal glucose tolerance; B: Breakfast; L: Lunch; D: Dinner; E: total energy intake; ppd: postprandial; GI: Glycemic Index, Glu: Glucose, Ins: Insulin, AUC: Area Under the Curve, iAUC: incremental area under the curve; CHO: carbohydrates; PRO: proteins; FAT: fats; M: meals; FPG: fasting plasma glucose; BW: body weight; RCT: randomized controlled clinical trial; HC: high carbohydrate; HF: high fat; HOMA: homeostatic model assessment for insulin resistance. Arrows pointing downward indicate a decrease. Two arrows pointing downward indicate a large decrease in the variables under investigation.
To address these issues, we conducted a long-term crossover study in subjects with early and late-stage IGT and naïve T2DM, who received weight maintenance diets as 6 and 3 meals per day [324]. We demonstrated that 6 vs 3 meals per day improved glycemic control in obese naïve T2DM subjects, resulting in significant reductions in HbA1c, peak glucose, postprandial glycemia and hunger, and stabilized or at least improved glycemic excursions reducing hunger and desire to eat in the subjects with IGT [324].

In conclusion, although there is evidence suggesting benefit from many regular smaller meals on glucose metabolism, it remains a controversial issue and more long-term studies are needed to make an evidence-based recommendation to people with IGT or T2DM.

4.3. Effects of Intermittent Fasting on Postprandial Glycemia

Intermittent fasting (IF) is characterized by interchange between periods of fasting and feeding (Figure 3). Fasting periods last longer than the typical whole night fasting of 8–12 h. The exact hours of the eating window can vary with eating occurring in the morning hours (i.e., 09:00–15:00, 09:00–13:00, etc.), in the middle of the day (i.e., 12:00–17:00, etc.), or extending later to night hours (i.e., 12:00–21:00, 13:00–18:00, etc.). Studies in mice have reported beneficial effects of IF against diabetes and obesity with improvements in glucose tolerance and insulin sensitivity, maintenance of insulinemia within normal levels, and improvements in the phosphorylated cyclic AMP response element-binding protein (CREB), rapamycin complex (mTOR), and AMP-activated protein kinase (AMPK) pathways, even in mice fed high-fat diets [256,335–337]. The concept behind IF is that long-term fasting diets can also lead to ketosis, during which glucose reserves run out and glycogen stores are insufficient to provide energy to the brain and the CNS, using ketone bodies and acetone as alternative fuels [338]. Ketosis has been proposed by some to positively affect metabolism with a reduction in ROS, lipolysis, autophagy, and increases in stress resistance, among others [339], further extending the effects of caloric restriction.

Table 5 describes the effects of eating windows in a day on postprandial hyperglycemia.

There are various types of IF protocols all of which divide the day or week in periods of feeding and fasting achieving energy restriction. In most cases there is no guidance for which foods to consume, and when to consume them. IF can be grouped into alternate-day fasting (ADF) (up to 75% energy restriction), whole-day fasting and time-restricted feeding (TRF), whereas in the weekly restricted feeding protocols energy intake is restricted from 1–3 days in a week up to 5 days in a month with ad libitum intake in the remaining days (i.e., 5:2 diet) [338,340,341].

Few TRF trials have been conducted in humans, and a small number of those focused on the effect of TRF patterns in people at risk for developing or with diagnosed T2DM (Table 5). Two short-term (2-week) trials compared the effects of a 9–10 h early or late eating window and reported that early-eating improved BP and glucose tolerance (decreased blood glucose areas under the curve, mean fasting and postprandial glucose levels), while late-eating increased plasma glucose, insulin, and triglyceride levels during the night [342,343]. A study of 11 weeks duration in which overweight participants followed a 6-h early TRF plan (08:00–14:00) resulted in a decrease in night glycemia, 24-h blood glucose fluctuations, fasting plasma glucose/insulin levels, HOMA-IR, and increased evening plasma insulin levels, without changes in day plasma glucose concentrations [344]. Patients with metabolic syndrome following a 10-h TRF also demonstrated reductions in body weight and improvements in glycemia (reduced HbA1c, plasma glucose/insulin levels and HOMA-IR) [345].
Table 5. Effect of the eating window in a day on Glycemia in healthy individuals and subjects at high risk for developing or with type 2 diabetes.

| Study                        | Health Status Age (Years) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups                                                                 | Dietary Intervention                                                                 | Selected Clinical Outcomes |
|------------------------------|---------------------------|---------------------------------------------|-------------|----------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|----------------------------|
| Carter, et al., 2018 [346]   | T2DM, overweight/obese   | 12 months Parallel randomized clinical trial | 97          | Intermittent energy restriction group (IER) Continuous energy restriction group (CER)   | IER: 500–600 kcal/day for 2 non-consecutive days of the week, usual diet for the other 5 days CER: 1200–1500 kcal/day Similar weekly energy restrictions | BW −5 kg in CER −6.8 kg in IER HbA1c −0.5% in CER −0.2% in IER Ins −0.3% in CER −1.2% in IER |
| Carter, et al., 2019 [347]   | T2DM, overweight/obese   | 12 months—24-month follow-up Parallel randomized clinical trial | 84          | Intermittent energy restriction group (IER) Continuous energy restriction group (CER)   | IER: 500–600 kcal/day for 2 non-consecutive days of the week, usual diet for the other 5 days CER: 1200–1500 kcal/day 24 months Follow-up | At 24 mo.: BW −3.9 kg in CER −3.9 kg in IER HbA1c +0.4% in CER +0.1% in IER TG −0.2 ± 0.3 mmol/L in CER −0.02 ± 0.2 mmol/L in IER |
| Varady, et al., 2013 [348]   | Normal BW/overweight     | 12 weeks Randomized, controlled, parallel-arm feeding trial | 30          | ADF Control (ctrl)                                                                     | ADF: 25% of E needs on fast day (~400–600 kcal at 12:00 h–14:00 h), and ad libitum eating on each alternating feed day ad libitum eating every day | ↓ BW in ADF −13% total Chol in ADF −20% TG in ADF −5% BP (systolic) in ADF −50% CRP in ADF |
Table 5. Cont.

| Study                          | Health Status | Age (Years) | BMI (kg/m²) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|-------------------------------|--------------|-------------|-------------|---------------------------------------------|-------------|-----------------------|----------------------|-----------------------------|
| Catenacci, et al., 2016       | Obese        | 41.15 ± 8.7 | 37.65 ± 4.85 | 8-week intervention & 24 weeks unsupervised follow-up Randomized trial | 26          | CR (caloric restriction) ADF (alternate day fasting) | CR: −400 kcal/day less than E needs ADF: ad libitum food in fed days, only water, calorie-free beverages and bouillon/stock cube soup on fast (0 kcal) days E distribution in both groups: 20% B, 30% L, 40% D and 10% snack | BW −7.1 kg in CR at week 8 −8.2 kg in ADF at week 8 −5 kg in CR at week 12 −5.7 kg in ADF at week 12 Glu (mg/dL) +3.3 in CR at week 8 +6 in ADF at week 8 +1.7 in CR at week 12 +2.6 in ADF at week 12 Ins (µU/mL) −0.2 in CR at week 8 +3 in ADF at week 8 −2 in CR at week 12 +0.4 in ADF at week 12 TG (mg/dL) −2.8 in CR at week 8 −25 in ADF at week 8 +12 in CR at week 12 +5.1 in ADF at week 12 |
### Table 5. Cont.

| Study | Health Status (Years) | Age (Years) | BMI (kg/m²) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|-------|-----------------------|-------------|-------------|---------------------------------------------|-------------|------------------------|-----------------------|----------------------------|
| Trepanowski, et al., 2017 [350] | Obese | 44 ± 11 | 34 ± 4 | 12 months (6 mo. weight loss & 6 mo. maintenance) RCT | 69 | Alternate day fasting group (ADF) Daily calorie restriction group (DCR) Control—No intervention group (ctrl) | ADF: 25% of energy needs on fast days; 125% of energy needs on alternating “feast days” DCR: calorie restriction to 75% of energy needs every day Ctrl: no-intervention control | ADF vs ctrl: At 6 months: −6.8% BW, −6.3 mg/dL Glu, −7.5 µIU/mL Ins, −2.49 HOMA-IR At 12 months: −6% BW, −3.9 mg/dL Glu, −5.9 µIU/mL Ins, −1.86 HOMA-IR DCR vs ctrl: At 6 months: −6.8% BW, −4.9 mg/dL Glu, −7.0 µIU/mL Ins, −2.56 HOMA-IR At 12 months: −5.3% BW, −9.6 mg/dL Glu, −4.6 µIU/mL Ins, −1.88 HOMA-IR |
| Gabel, et al., 2019 [351] | Insulin resistant subjects | 42 ± 3 y.o. | 35 ± 1 | 12 months (6 mo. weight loss & 6 mo. maintenance) RCT | 100 (43 completed) | Alternate-day fasting group (ADF) Daily calorie restriction group (CR) Control group (ctrl) | 6-months reduced net E intake by 25% Fasting days: 25% of E needs at lunch (12:00 h–14:00 h) Alternating feast days: 125% of E needs over 3 meals/day 6-mo. reduced net E intake by 25% per day over 3 meals every day 6-mo. weight maintenance phase: Both for ADF & CR groups ADF consumed 50% of E needs on fast days and 150% on feast days CR consumed 100% of E needs/day Instructed to maintain body weight | ↓ weight (~18% for ADF, and ~14% for CR) Glu (mg/dL): no significant changes Ins (µIU/mL): −44% ADF 6 mo, −52% ADF 12 mo, −23% CR 6 mo, −14% CR 12 mo HOMA-IR: −48% ADF 6 mo, −54% ADF 12 mo, −19% CR 6 mo, −17% CR 12 mo |
Table 5. Cont.

| Study | Health Status | Age (Years) | BMI (kg/m²) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|-------|---------------|-------------|-------------|---------------------------------------------|-------------|-----------------------|----------------------|-----------------------------|
| Hoddy, et al., 2014 [352] | Obese | 45.5 ± 2.5 | 34.5 ± 1 | 8 weeks Randomized, parallel-arm feeding trial | 59 | ADF-L Alternate day fasting—intake at lunch | 25% of E needs on fast day and ad libitum eating on feed day | BW − 3.5 kg in ADF-L − 4.1 kg in ADF-D − 4.0 kg in ADF-SM Glu − 2% in ADF-L − 1% in ADF-D − 1% in ADF-SM Ins no change in ADF-L − 18% in ADF-D − 12% in ADF-SM HOMA-IR − 10% in ADF-L − 27% in ADF-D − 19% in ADF-SM TG − 6% in ADF-L − 8% in ADF-D − 1% in ADF-SM | |
| | | | | | | ADF-D Alternate day fasting—intake at dinner | | |
| | | | | | ADF-SM Alternate day fasting—intake in small meals | | |
| | | | | | ADF-D: One meal (D) at 18.00 h–20.00 h on each fast day | | |
| | | | | | ADF-SM: divided their fast day meal in 3 mini meals → 100 kcal at 6:00 h–8:00, 300 kcal at 12:00 h–14:00 h and 100 kcal at 18:00 h–20:00 on each fast day | | |
| Study                        | Health Status Age (Years) BMI (kg/m²) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention                                                                 | Selected Clinical Outcomes |
|------------------------------|---------------------------------------|---------------------------------------------|-------------|------------------------|---------------------------------------------------------------------------------------|-----------------------------|
| Harvie, et al., 2011 [353]   | Overweight or obese 40 ± 4 30.6 ± 5.1 | 6 months Randomized trial                   | 89          | Continuous energy restriction group (CER) intermittent energy restriction group (IER) | CER: 25% restriction below estimated requirements for 7 days per week IER: 75% restriction for 2 days per week, with no restriction on the other 5 days per week 25% PRO, 45% low GL CHO, 30% FAT (15% MUFAs, 7% SFAs, 7% PUFAs) | BW −5% in CER −7% in IER  
Ins −15% in CER −29% in IER  
HOMA −19% in CER −27% in IER  
Glu −2% in CER −2% in IER  
Adiponectin no change in CER +10% in IER  
Ghrelin +11% in CER +13% in IER  
TG −23% in CER −17% in IER  
BP (systolic) −6% in CER −3% in IER |
| Study | Health Status | Age | BMI (kg/m²) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|-------|---------------|-----|-------------|---------------------------------|-------------|----------------------|---------------------|--------------------------|
| Harvie, et al., 2013 [354] | Overweight | 47.4 ± 7.7 | 30.9 ± 5.1 | 4 months (3 months weight loss and 1 month maintenance) Parallel randomized clinical trial | 115 | IECR: energy and CHO restriction IECR + PF: allowed ad libitum PRO and FAT DER: daily energy restriction | IECR: 25% overall E restriction, And for 2 d/week restricted CHO (<40 g/day), On restricted days: 20% CHO, 45% PRO and 35% FAT IECR + PF: 25% overall E restriction, And for 2 d/week restricted CHO (<40 g/day) and ad libitum PRO and FAT (MUFA and PUFA), On restricted days: 15% CHO, 35% PRO and 50% FAT DER: 25% overall E restriction, 45–50% CHO, 20–25% PRO and 30% FAT | BW − 5.5 kg in IECR − 5.1 kg in IECR + PF − 3.8 kg in DER Ins − 21% in IECR − 11% in IECR + PF − 10% in DER HOMA-IR − 25% in IECR − 16% in IECR + PF − 11% in DER HbA1c no change in HbA1c in all groups Glu no change in Glu in all groups TG − 9% in IECR − 14% in IECR + PF − 7% in DER BP (systolic) − 3% in IECR − 13% in IECR + PF − 9% in DER |

- Time restricted Feeding (TRF)
| Study                          | Health Status Age (Years) BMI (kg/m²) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|-------------------------------|--------------------------------------|--------------------------------------------|-------------|-----------------------|---------------------|--------------------------|
| Hutchison, et al., 2019 [342] | Overweight/Obese 55 ± 3 33.9 ± 0.8   | 14 exp. days Randomized crossover trial     | 15          | TRFe TRFd             | 2 × 7 days separated by a 2-week washout Eating window: 8:00 h–17:00 h Eating window: 12:00 h–21:00 h Ad libidum water and very low-calorie (<4 kcal/serving) drinks and foods | ↓ BW on day 7 ↓ iAUC<sub>Glu</sub> −36% in TRFe −21% in TRFd No effect on fasting Glu, Ins, (but a ↓ trend) ↓ mean fasting Glu for TRFe vs baseline ↓ 3 h ppd Glu for TRFe vs baseline |
| Wehrens, et al., 2017 [343]   | Normal BW/Overweight 18–30 20–30     | 13 exp. Days Human laboratory trial         | 10          | One group             | Sleeping schedule: ~ 23:00–6:30 h Day 1–3: Wake up at 6:30 h, B at 7:00 h, L at 12:00 h, D at 17:00 h Day 4–5: No treatment, measurements Day 6–11: Wake up at 6:30 h, B at 12:00 h, L at 17:00 h, D at 22:00 h Day 12–13: No treatment, measurements Isocaloric meals: 55% CHO, 15% PRO, 30% FAT | Late meals resulted in: ↑ 5.69 h Glu acrophase (into sleeping time) ↑ 1.5 h Ins acrophase ↑ 1 h TG acrophase |
Table 5. Cont.

| Study                     | Health Status | Age (Years) | BMI (kg/m²) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention | Selected Clinical Outcomes |
|---------------------------|---------------|-------------|-------------|---------------------------------------------|-------------|-----------------------|----------------------|---------------------------|
| Jamshed, et al., 2019 [344]| Overweight    | 32 ± 7      | 30.1 ± 2.7  | 8 days Randomized controlled crossover study | 11          | eTRF Control (ctrl)   | 6 h feeding/18 h fasting  
Feeding: 8:00 h to 14:00 h  
B at 8:00 h, L at 11:00 h, and D at 14:00 h  
12 h feeding/12 h fasting  
Feeding: 8:00 h to 20:00 h  
B at 8:00 h, L at 14:00 h, and D at 20:00 h  
3 daily meals were matched:  
15% PRO, 50% CHO, 35% FAT  
4 days in each condition—3.5–5 weeks washout—4 days in other condition | No significant changes in day Glu  
−7 ± 2 mg/dL night/sleep Glu in eTRF  
−4 ± 1 mg/dL 24-h blood Glu in eTRF  
−12 ± 3 mg/dL MAGE in eTRF  
−2 ± 1 mg/dL morning fasting Glu in eTRF  
2.9 ± 0.4 mU/L morning fasting Ins in eTRF  
−0.73 ± 0.11 HOMA-IR in eTRF  
+25% ± 9% IRS2 gene expression in eTRF  
+4.5 ± 1.6 mU/L evening fasting Ins in eTRF  
+1.09 ± 0.43 evening HOMA-IR in eTRF  
No changes in GLUT1, GLUT4, or IRS1 gene expression at either time of day |
| Wilkinson, et al., 2020 [345]| Metabolic syndrome patients | 59 ± 11.14  | 33.06 ± 4.76 | 12 weeks Single-arm, paired-sample trial | 19          | TRF                   | 10 h feeding/14 h nightly fasting  
Eating at will, subjects reported foods consumed via smartphone app (food and time logs) | TRF vs baseline:  
−3% BW  
−5% Blood Glu (CGM)  
−5% Fasting blood Glu  
−2% HbA1c  
−21% Fasting Ins  
−30% HOMA-IR |
Table 5. Cont.

| Study                     | Health Status Age (Years) BMI (kg/m²) | Duration and Design of Dietary Intervention | Sample Size | Description of Groups | Dietary Intervention                                                                 | Selected Clinical Outcomes                                      |
|---------------------------|--------------------------------------|---------------------------------------------|-------------|------------------------|----------------------------------------------------------------------------------------|-----------------------------------------------------------------|
| Sutton, et al., 2018 [355]| Overweight with IGT 56 ± 9 32.2 ± 4.4 | 5 weeks Randomized, crossover trial (controlled feeding) | 8           | eTRF: 6 h eating window/day, 3 meals, dinner before 15:00 h, ~18 h fasting | eTRF: 6 h eating window/day, 3 meals, dinner before 15:00 h, ~18 h fasting ≤ TRF: 12 h eating period/day, 3 meals, ~12 h fasting between 20:00 h–6:30 h Each plan for 5 weeks, with a 7-week washout period Isocaloric and eucaloric controlled feeding diets (BW maintenance) 15% PRO, 50% CHO, and 35% FAT | No weight loss Fasting Glu no significant changes Mean Glu no significant changes Fasting Ins −3.4 ± 1.6 mU/L Mean Ins −26 ± 9 mU/L Peak Ins −35 ± 13 mU/L Insulinogenic index +14 ± 7 mu/mg IR (iAUC) −36 ± 10 mU/mg |
| Parr, et al., 2020 [356]  | T2DM 50 ± 8.9 34 ± 4.8               | 6 weeks Pre-post non-randomized             | 19          | Habitual diet (2 weeks) | Habitual diet: ~8400 kJ/day; 35% CHO, 20% PRO, 41% FAT, 1% alcohol 9 h feeding/15 h fasting TRE diet: ~8500 kJ/day; 35% CHO, 19% PRO, 42% FAT, 1% alcohol Eating at will | No weight loss −3% HbA1c −3.6% Glu +18% Ins |
| Moro, et al., 2016 [357]  | Healthy athletes 29.21 ± 3.8 BW 84.6 ± 6.2 Kg | 8 weeks Randomized, single blind trial     | 34          | TRF (2 weeks) | TRF Normal Diet group (ND) 8 h feeding/16 h fasting Meals: 13:00 h (40% of total cal), 16:00 h (25% of total cal), 20:00 h (35% of total cal) Meals: 8:00 h (25% of total cal), 13:00 h (40% of total cal), 20:00 h (35% of total cal) | ↓ in fat mass −16.4% in TRF −2.8% in ND Glu −11% in TRF no change in ND Ins −36.3% in TRF −13% in ND |

Abbreviations: B: Breakfast; L: Lunch; D: Dinner; E: energy; ppd: postprandial, Abbreviations: T2DM: type 2 diabetes; IGT: impaired glucose tolerance; B: Breakfast; L: Lunch; D: Dinner; E: energy; ppd: postprandial, GI: Glycemic Index; GL: Glycemic Load; Glu: Glucose; Ins: Insulin; AUC: Area Under the Curve; iAUC: incremental area under the curve; CHO: carbohydrates; PRO: proteins; FAT: fats; TRF: time-restricted feeding; CGM: continuous glucose monitoring; BW: body weight; RCT: randomized controlled clinical trial; HOMA: homeostatic model assessment for insulin resistance; TG: triglycerides; BP: blood pressure. Arrows pointing upwards or downwards indicate an increase or decrease.
Improved insulinemia was observed in overweight subjects with IGT who followed a 6-h TRF, but in this study significant changes in fasting or mean glycemia were not observed [355]. T2DM subjects, who followed a 2-week TRF plan with a 9-h eating window per day demonstrated reductions in hyperglycemia and HbA1c, and increases in plasma insulin levels, even though there was no weight loss observed [356]. TRF’s favorable glycemic and insulinemic effects were reported in healthy athletes performing resistance training [357]. An isocaloric study showed that more body fat was lost with 6-h vs 12-h feeding with significant differences in weight loss (1.5 kg), plasma adiponectin, testosterone, interleukin-1β, and IGF-1 levels, but no changes in fat free mass, resting energy expenditure, and plasma thyroid hormone/glucose levels [357]. A study demonstrated that after one year of following IF (2/5 days per week consuming 500–600 kcal) the weight loss achieved was accompanied by reduction in insulinemia and HbA1c [346]. Weight loss and decreased insulin responses were also observed in a group that received a caloric restricted diet, but these reductions were smaller than those of the IF group; two years after that intervention, both groups had regained some of the weight, HbA1c was lower in the IF group, and both groups now presented a reduction in plasma TG levels [347].

Smaller duration studies from 8 to 12 weeks examining ADF have shown that it is a safe and tolerable approach to weight loss, although it provided conflicting evidence regarding markers for CVD protection [348,349]. In a 12-month study, ADF (25% of daily energy needs on fasting days: one day on, one day off) was compared with a calorie-restricted (75% of energy needs every day) and a control diet [350]. It was demonstrated that although the two intervention groups lost weight, insulin sensitivity, blood lipid levels and blood glucose were not improved. Analysis of insulin-resistant participants from that study, revealed similar weight loss between IF and calorie-restricted interventions after 12 months [351]. Both diet interventions had similar effects on reductions in fat mass and BMI. However, ADF resulted in greater decreases in fasting insulin and insulin sensitivity despite a similar decrease in body weight with the other diet plans [351]. In contrast, another study showed that allowing people to consume the fast day meal at the evening dinner (or dividing it into smaller meals) produced similar weight loss and CVD protection as consuming the meal at lunch, thus allowing for more flexibility in meal timing and possible between long-term adherence to ADF protocols [352].

A 5-week crossover study in men with prediabetes showed the early-time IF (6 h feeding period, with dinner before 15:00) to be in alignment with circadian rhythms in metabolism compared to a control schedule (12 h feeding period), improved insulin sensitivity, β-cell responsiveness, BP, oxidative stress, and appetite, independently of weight loss [355]. Early vs delayed TRF improved glucose tolerance in men at risk for T2DM [342]. Greater weight loss was observed in early eaters (before 15:00) vs later eaters (after 15:00) [312]. Three isocaloric studies showed that front loading (high caloric intake at breakfast) vs evening dinner improved weight loss [317,320,358]. A TRF pattern, with a long fast (>12–14 h) and a limited eating window not extending late in the night, with ~4–10 h in which meals were distributed and consumed, seemed to have several health benefits independent of energy restriction [359]. It has been reported that ADF studies in normal weight, overweight, and obese subjects with a duration of 3–12 weeks were effective in achieving an approximately 3–7% weight loss, 3–5.5 kg body fat loss, lowering blood lipids (approximately 10–21% for total cholesterol and 14–42% for triglycerides) [360]. Whole day fasting studies with a duration of 12–24 weeks also achieved a 3–9% body weight loss, body fat loss and lowering of blood lipids (approximately 5–20% for total cholesterol and 17–50% for triglycerides) [360].

Many people have time-restricted windows, they do not eat their first meal before 14:00 and stop eating at 02:00. Feeding periods after 16:00 had either no effect or unfavorable effects on postprandial glycemia, β-cell responsiveness, BP, and blood lipids [360–362]. It has been shown that fasting until 12:00 results in increased postprandial hyperglycemia and impaired insulin responses after lunch and evening dinner in healthy and T2DM subjects [72], due to effects directly on clock genes and time-controlled genetic expression [285].
To examine whole week feeding restriction regimens, one study [363] randomized 54 obese individuals with T2DM into 3 groups: (a) a very low-calorie diet (VLCD) for 5 days and then 1 day/week VLCD for 15 weeks; (b) a VLCD for 5 days and then 5 days VLCD every 5 weeks consuming 1500–1800 kcal per day without fasting; and (c) a control group consuming 1500–1800 kcal per day for the duration of the study. This study showed that both VLCD groups lost significantly more body weight with the second group losing >5 kg [363]. Another study with 63 overweight and obese T2DM subjects randomized to consume 400–600 kcal/day 2 days/week and then, typical energy intake for 5 days/week vs whole day energy restriction to 1200–1550 kcal/day showed that after 12 weeks there were similar reductions in HbA1c, body weight, antidiabetic medications, body composition, and appetite in both groups [364]. The issues arising with IF is that there is a variety of definitions making hard to interpret the scientific results in a systematic manner: the number of fasting days and the energy intake vary in each study, the management of hunger during fasting days and the days of regular feeding are in most cases unknown, the probability of overconsumption in days of typical feeding is high and social life is affected, particularly in countries of Southern Europe (i.e., Italy, Greece, Spain, etc.) where eating late dinner after 19:00 is very common. Two studies showed that intermittent continuous energy restriction (IER) was a similarly effective alternative to continuous energy restriction (CER) regarding weight loss [353,354].

In conclusion, some practical strategies from IF trials that seem to be effective in body weight and postprandial glycemia support consuming the evening dinner earlier in the day, consuming most of the energy at breakfast and lunch, avoiding snacks late at night, and reducing feeding time by 1–2 h with the last meal before 18:00 or the latest at 20:00. Due to the variable IF protocols, the choice of IF pattern, the short duration of trials, the varied window of energy intake and the heterogeneous populations studied, more well-designed longer duration studies are needed to draw safe conclusions and recommendations.

5. Alternative Dietary Interventions and Postprandial Glycemia

Various plant-derived foods, plant compounds and functional ingredients/foods have been linked with ameliorated postprandial hyperglycemia, and they may be considered “alternative” dietary interventions for diabetes management.

5.1. Vitamins–Minerals

Without underlying deficiency, the benefits of multivitamins or mineral supplements on glycemia for prediabetes or T2DM have not been supported by evidence, and therefore routine use is not recommended [114]. Accordingly, the routine use of chromium, vitamin D, micronutrient supplements for improving glycemia in people with diabetes is not supported by evidence and is also not recommended [114].

5.2. Herbs and Spices

Use of herbs and spices during cooking is encouraged as it is a safe method for flavoring and preservation of food, and for providing antioxidant ingredients, such as phenolic compounds. Cinnamon, one of the most used culinary spices, may be considered an alternative food intervention. It belongs to the family and is found commercially in many forms including sticks of bark, powder, and powder-derived extracts. Cinnamon contains various bioactive compounds such as cinnamaldehyde, eugenol, trans-cinnamic acid, phenolic compounds, tannins, catechins, terpenes, proanthocyanidins, and coumarin. The content of these compounds differs according to the form in which cinnamon is used. Both in vitro and in vivo animal studies have shown insulin-enhancing or insulin-like effects after cinnamon administration [365–367]. Several clinical trials have been conducted since the 2000s that administered cinnamon as extracts, capsules or supplements at varying amounts ranging from 0.5 g to 6 g per day, in subjects with T2DM. Most of the studies have reported a generally favorable effect of cinnamon on postprandial/fasting hyperglycemia and diurnal glucose fluctuations [368–375]. It has been shown that even low amounts
(1 g/day) of cinnamon powder was enough to reduce fasting glucose and improve blood lipid profiles of T2DM patients [375]. Higher doses also confirmed these effects in some studies [369–374]. Other studies reported additional improvements on insulin [369] and HbA1c [369,370,372,373] levels after cinnamon addition in the diet (as extracts or supplements). Glucose and insulin responses to oral glucose tolerance test (OGTT) were also found to be improved in healthy subjects after cinnamon supplementation (3 gr/day) [376]. However, not all studies agree with the favorable glycemic effects of cinnamon. Results from 6 RCTs did not find a difference in blood glucose or insulin levels after cinnamon consumption [368,377–382]. Decreased serum triglycerides, total cholesterol, and LDL-cholesterol were also demonstrated in a few reports [370,372,375]. Interestingly, one study reported that the positive effects of cinnamon consumption on glycemia are acute and do not persist after cessation [376]. Cinnamon intake tended to be beneficial for controlling glycemia in T2DM patients with poor glycemic control, but more research is needed to establish these effects. However, use of any herbal supplements, including cinnamon, curcumin, or aloe vera, for improving glycemia in people with diabetes is not supported by sufficient scientific evidence and is therefore not recommended [114].

5.3. Fermented Foods

Fermented foods and beverages, defined as “foods made through desired microbial growth and enzymatic conversions of food components”, have been used for thousands of years, and are proposed to offer several health benefits due to their bacteria strains, and the presence of certain organic acids in foods generated through fermentation or added, and are responsible for texture, flavour, and better preservation of fermented foods [383,384]. Lactic acid may lead to slower starch absorption due to inhibition of amylolytic enzymes, thus, reducing its bioavailability due to the interaction between starch and gluten, and the delay of gastric emptying, leading to lower postprandial glycemic responses [385,386]. Inclusion of the sodium salt of propionic acid to whole-meal barley bread has also been reported to lower postprandial hyperglycemia [383]. Sourdough fermentation of wheat dough has been repeatedly reported to lower the GI of bread and postprandial glucose excursions, due to the formation of organic acids, leading to reduced rate of starch digestion and delayed gastric emptying [383,385–388]. Results from one study reported that consumption of meals containing beta-glucan, fermented in the colon, resulted in decreased postprandial glucose concentrations due to a delayed and somewhat reduced carbohydrate absorption from the gut not from the effects of fermentation in the colon [389]. However, another study showed that sourdough fermentation is a method able to lower the postprandial glycemic responses to bread, but this did not relate to the starch accessibility or general bioavailability, but rather to bacteria-induced delayed gastric emptying [390]. The latter was also supported by another study reporting that the lactic acid in fermented milk products did not lower the postprandial glycemias and concluded that the presence of organic acids may counteract the insulinotropic effect of milk in mixed meals [391]. Additionally, it has also been reported that fermentation of carbohydrates may regulate postprandial glucose excursions to a second meal by reducing NEFA competition for glucose disposal and to a minor extent by affecting intestinal motility [392].

Vinegar (acetic acid) is an example of fermented food. Vinegar consumption dates back to ancient times with reports of its use by Hippocrates for wound care [393]. In folk medicine, vinegar is considered a natural remedy and is used as such, whereas now it has been included in the “super foods” for its properties and claims of effects on weight loss, digestion, and even skin quality. Most of trial data support beneficial effects of vinegar on postprandial glycemia and overall glycemic control. In healthy subjects, vinegar consumption with a meal (either as dressing on salad or as a drink) resulted in generally lower postprandial glycemia [386,394–396] and in some studies reduced insulinemia [386] as well. In one study, a 55% reduction in post-meal glycemia was reported when a high GL meal was consumed [397]. Another study suggested that the effects of vinegar consumption are prominent when ingested with carbohydrate-rich meals rather than with low GL
meals [397]. A few clinical trials have also been conducted in individuals with prediabetes and/or T2DM. Vinegar intake before or together with a meal was associated with improved glycemia, reduced post-meal area under the curve for glucose, reduced fasting blood glucose, insulin and triglycerides, increased muscle glucose uptake, and reduced the need for fast-acting insulin in subjects with T1DM [398–403]. However, in another study, postprandial plasma glucose or insulin were unaffected by vinegar co-ingestion [404]. Currently, there is also some data explaining the possible mechanisms by which vinegar affects glycemia. Acetic acid consumption has been shown to delay gastric emptying in healthy subjects and in people with diabetes [386,405]. Additionally, digestion of complex carbohydrates is also delayed by inhibition of relevant enzymes such as sucrase and disaccharidase [406,407]. All these effects have been shown to reduce postprandial glucose responses. Vinegar intake at bedtime was associated with lower fasting glycemia by decreasing the rates of hepatic gluconeogenesis and improving insulin secretion in subjects with T2DM [401].

Yoghurt and cultured milk products are other examples of fermented foods and have been shown to reduce postprandial glycemic responses in both healthy individuals, and subjects with prediabetes or overt T2DM [384], beyond the milk from which these products are made [408]. Similar results have been reported in at least one RCT with milk kefir, kimchi, sauerkraut and natto [384]. A recent consensus statement on fermented foods from the International Scientific Association for Probiotics and Prebiotics reported that although the family of fermented foods is large (variable food categories including fermented dairy products and other fermented foods with living versus dead microorganisms; food types including fermented vegetables, fermented soy and yoghurt; and individual fermented food products with well-characterized strains and nutrient compositions), and not all foods are examined or proven for their health benefits, and the mechanisms by which they may lead to lower postprandial glycemic responses have not been fully elucidated, consumption of some of these food products seems promising and their beneficial health effects remain to be established by more and well-designed RCTs and may also be obtained from harvesting information from existing population-based diet and health databases [384]. In conclusion, consumption of fermented foods should be encouraged and some of these foods may lead to amelioration of postprandial hyperglycemia and IR.

5.4. Probiotic Dairy Foods

Another alternative dietary intervention includes probiotic dairy foods. A recent mini-review suggested that matured products (i.e., ripened cheeses), fermented dairy products (kefir), and whey-based products (mainly milk beverages), and the addition of prebiotics and/or plant-derived products have a higher ability to regulate postprandial glycemia due to their probiotic strains with higher proteolytic and exopolysaccharides-forming abilities leading to inhibition of digestive enzymes, such as the α-amylase (1,4-alpha-D-glucan-glucanohydrolase), the enzyme that hydrolyses polysaccharides to glucose and maltose oligosaccharides, and the α-glucosidases, membrane bound enzymes located in the epithelium of the brush borders of the small intestine, that hydrolyze the oligosaccharides at the non-reducing links releasing the bound α-glucose, thus increasing the blood glucose levels [409]. However, a recent systematic review with 27 probiotic interventions (Lactobacillus, Bifidobacterium, Clostridium and Akkermansia) reported contradictory results regarding the effects of certain probiotics on amelioration of IR, suggesting the need for long-term RCTs in people with obesity and cardiometabolic risk [410].

5.5. Other Alternative Dietary Interventions: i.e., Inulin, Polyphenols, Chia Seeds, Nuts and Whey Protein

Results from two systematic reviews and meta-analyses of 33 RCTs in healthy, overweight/obese, prediabetes, T2DM, and hyperlipidemic subjects examining the metabolic effects of inulin-type fructan intake, reported a reduction in blood glucose, total cholesterol, and triglyceride concentrations in people with diabetes, although the mechanisms were
inconclusive without differences from controls on body weight and blood insulin and with data showing large heterogeneity [411,412].

A western diet is able to deliver between 109 and 313 mg of polyphenol per day, Mediterranean diet between 820 mg and 1.3 g per day, whereas red wine, the food with the highest resveratrol content, contains around 3 mg/100 mL [413]. Typical foods containing polyphenols include tea, coffee, chocolate, cocoa, cinnamon, grape, pomegranate, red wine, berries, and olive oil [414]. A systematic review and meta-analysis of 36 RCTs examining the effects of polyphenols (extracts, supplements, and foods), supplemented in doses of 28 mg to 1.5 g, for 0.7 to 12 months on HbA1c levels in healthy subjects and individuals with prediabetes or T2DM, reported significant reduction in HbA1c (~0.5%), in those with T2DM, without significant effects in the healthy subjects and those with prediabetes [413]. Polyphenol rich extracts had a more marked effect in reducing HbA1c among the trials; with 125 mg/day isoflavonoids from soy products, 1–3 g of cinnamon (27 mg/day of coumarin), 250 mg/day resveratrol from extracts, reported to have the highest efficiency in reducing HbA1c [413]. Dietary polyphenols’ beneficial reported effects on postprandial hyperglycemia and IR may be due to inhibition of \( \alpha \)-amylase and \( \alpha \)-glucosidases, inhibition of intestinal glucose absorption by sodium-dependent glucose transporter-1, stimulation of insulin secretion, and reduction of hepatic glucose output [414].

Chia (Salvia hispanica L.) seeds and oil, contain \( \alpha \)-linolenic acid, vegetable protein and dietary fiber [415]. A systematic review and meta-analysis of 12 trials with healthy individuals, athletes, and subjects with T2DM or metabolic syndrome, showed reductions in postprandial blood glucose and HDL cholesterol levels, and BP, only at the high doses, and only in subgroup analysis with the effects being modest and probably not clinically significant, and with the quality of evidence of the studies being low or very low [415].

Although there is increased scientific interest in the metabolic effects of nuts, the family of nuts is heterogeneous, and not all nuts are reported to be beneficial. A systemic review and meta-analysis of 6 RCTs examining the effects of pistachio nuts on glycemic control and insulin sensitivity in people with T2DM, prediabetes, or metabolic syndrome, reported a significant reduction in fasting glucose and IR, without differences from controls on HbA1c and fasting plasma insulin; the reported beneficial effects were possibly related to their high content of antioxidants (beta-carotene, lutein, proanthocyanidins, and vitamin E), anti-inflammatory compounds, fiber, and healthy fats, monounsaturated fatty acids [416]. Pistachio nuts’ high content of antioxidant components has been proposed to be involved in their beneficial effects on insulin sensitivity [417]. A positive impact of a very high daily dose (85 g/day) of pistachio nuts on postprandial insulinemia has been reported, particularly when consumed with a high carbohydrate diet [418,419]. Also, their monounsaturated fatty acids have been suggested to reduce oxidative stress and improve the insulin-signaling pathway and IR, by maintaining membrane translocation of glucose transporters along with buffering \( \beta \)-cell hyperactivity [420,421]. Moreover, it has been suggested that consumption of high daily doses of pistachio nuts (>57 g/day) may have an up-regulatory effect on GLP-1 secretion in healthy subjects [418], thus explaining the improvement in postprandial insulin secretion. One study conducted on almond nuts reported improvements in glycemic control and lipid profiles in T2DM, without differences in IR vs controls [422]. A recent systematic review and meta-analysis of 9 RCTs examining the effects of almonds on gut microbiota, glucose metabolism and inflammatory parameters in T2DM reported that almond-based diets have significant effects in promoting the growth of short-chain fatty acid-producing gut microbiota, and lower HbA1c and body weight, but with no observed differences in the levels of fasting or 2-h postprandial blood glucose, inflammatory markers (C-reactive protein and tumour necrosis factor- \( \alpha \)), GLP-1, fasting blood insulin, and IR [423]. Another systematic review and meta-analysis of 40 RCTs examining the effects of tree nuts on indices of glycemic control, reported that consumption of peanuts or tree nuts significantly decreased IR and fasting insulin, without effects on HbA1c or fasting glucose levels [424]. Walnuts have been shown to improve IR in another study [425]. A recent systematic review and meta-analysis of 16 RCTs examining the effects
of walnuts on markers of blood glucose control, reported that consumption of walnuts did not result in significant changes in fasting blood glucose levels or IR, with available studies having either “some concern” or be “at high risk” of bias [426]. Although the study results regarding peanuts and tree-nuts on indices of glycemic control seem promising, high quality RCTs with larger population sizes and longer duration are needed to determine the exact efficacy, mechanism of action, precise daily dose, duration, and possible adverse effects of an effective intervention with these types of nuts in prediabetes and T2DM.

A recent systematic review and meta-analysis of 22 RCTs examining the effects of whey protein on glycemic status in patients with metabolic syndrome, reported that consumption of whey protein decreased significantly HbA1c, insulin/triglyceride/total cholesterol levels, and IR without effects on HDL cholesterol and fasting blood glucose levels [427]. Whey protein intake has been suggested to improve metabolic parameters due to bioactive substances, including immunoglobulins, glutamine, lactoferrin, and lactalbumin, which have been shown to activate the release of incretin hormones including GIP and GLP-1, whilst peptides from whey hydrolyzation have also been shown to inhibit dipeptidyl peptidase-IV and inhibit degradation of GIP and GLP-1 [427,428], all of which may have an important role in the improvement of IR [427]. Whey protein is an excellent source of branched-chain amino acids (BCAAs) and it has been shown that after its digestion, a rapid increase in amino acids, particularly BCAAs leads to insulin release, which may improve postprandial hyperglycemia [427].

In conclusion, until further well-designed of long duration studies are performed, it is safe and inexpensive to suggest that consumption of herbs, spices, such as cinnamon, fermented foods, such as vinegar, whey protein, peanuts, and tree nuts may offer some beneficial effects on postprandial glycemia and IR, additional to the overall diet implemented. Figures 2 and 3 describe the key available study findings.

6. Conclusions

Postprandial hyperglycemia and IR are complex issues influenced by many factors and causes. Their management is critical for the prevention of T2DM and amelioration of cardiometabolic risk factors. Diet is the cornerstone of glucose metabolism and weight loss is the remedy for IR. The extent and pace of weight loss, the dietary pattern of choice, and the macronutrient composition of the proposed diet remain to be elucidated with large, well-designed, long-term RCTs. Individualization and patient-centered approach should be the primary method of conduct. However, a balanced diet such as a Mediterranean-style, low GL diet may be a suitable approach to achieve both weight loss and maintenance and ameliorate postprandial hyperglycemia and IR. Lifestyle counseling using moderate energy restriction, regular physical activity and dietary behavior modification techniques has proven to be effective in optimally managing glycemic and insulinemic responses. However, dietary plans, foods and ingredients seem to have many differences when other variables are considered, such as glycemic or insulinemic responses or HbA1c. Although much more research is needed, some key points from the available scientific data for amelioration of postprandial hyperglycemia and IR may include the following: (a) lowering the total amount of carbohydrates consumed during the day to 40–50% of daily energy intake, such as in the case of Mediterranean-style diets, (b) consuming the majority of carbohydrates at lunch time, (c) adding lean proteins, plant proteins, and “good” fats (such as olive oil, peanuts and tree nuts, etc.) in meals, (d) following a meal sequence consuming vegetables first, then proteins and fats and then carbohydrates, particularly unprocessed ones, (e) choosing foods that do not lead to augmented glucose excursions and peaks to nadirs, (f) avoiding eating occasions late at night, and (h) consuming meals at consistent/regular times.

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Abbreviations

| Abbreviation | Description |
|--------------|-------------|
| IR           | Insulin resistance |
| T2DM         | Type 2 diabetes |
| HDL          | High density lipoprotein cholesterol |
| IGT          | Impaired glucose tolerance |
| GI           | Glycemic index |
| GL           | Glycemic load |
| NEFA         | Non-esterified free fatty acids |
| CVD          | Cardiovascular disease |
| NAFLD        | Non-alcoholic fatty liver disease |
| CNS          | Central nervous system |
| HGP          | Hepatic glucose production |
| GLP-1        | Glucagon-like peptide 1 |
| GIP          | Glucose-dependent insulino tropic polypeptide |
| HbA1c        | Glycated hemoglobin A1c |
| NGT          | Normal glucose tolerance |
| BP           | Blood pressure |
| RCT          | Randomized Controlled Clinical Trial |
| ADA          | American Diabetes Association |
| VLC          | Very low carbohydrate |
| IRS          | Insulin receptor substrate |
| GLUT         | Glucose transporter |
| HOMA-IR      | Homeostatic model assessment for insulin resistance |
| SCN          | Suprachiasmatic nucleus |
| VLCD         | Very low calorie diet |
| BMI          | Body mass index |
| Per          | Period |
| LDL          | Low density lipoprotein cholesterol |
| PCO          | Polycystic Ovary Syndrome |
| OGTT         | Oral glucose tolerance test |
| IF           | Intermittent fasting |
| ADF          | Alternate day fasting |
| TRF          | Time restricted feeding |
| IGF-1        | Insulin-like growth factor-1 |
| ROS          | Reactive Oxygen Species |

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