A Review of Recent Evidence from Meal-Based Diet Interventions and Clinical Biomarkers for Improvement of Glucose Regulation

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ABSTRACT: In recent decades, the prevalence of diabetes has rapidly increased worldwide. Medical nutrition therapy has been identified as a major therapeutic support for diabetic patients, while preventive strategies in prediabetic or high-risk individuals have mainly focused on supplementation with bioactive compounds. Recently, meal-based interventions have been investigated as novel and safe long-term strategies for improving glucose regulation. However, evaluation of meal-based interventions is difficult since it requires analysis of sensitive markers. Biomarkers can also be used to identify individuals at risk for diabetes, which is important for disease prevention. In this review, we summarize current evidence from meal-based intervention studies conducted with the aim of improving glucose homeostasis in individuals at risk of diabetes using clinical biomarkers currently used to assess diabetic risk. Very low-calorie diets have significantly improved glucose regulation in obese adults and in adults with type 2 diabetes mellitus. In particular, changing the ratios of macronutrients through calorie restriction reduces fasting glucose level and hemoglobin A1c levels in patients with diabetes mellitus. However, this effect is limited in both obese and healthy adults. To date, multiple glucose-related markers have been identified as clinical biomarkers of diabetes. Additional clinical biomarkers include cholesterol levels, hematological markers, and inflammatory markers. Taken together, the evidence presented in this review may help for selection of clinical biomarkers for meal-based preventive approaches for non- or pre-diabetic individuals to prevent onset of diabetes.

Keywords: biomarker, clinical, diabetes mellitus, glucose, meals

INTRODUCTION

Diabetes is a metabolic diseases caused by defects in insulin secretion and/or insulin action (American Diabetes Association, 2010). The global prevalence of diabetes has risen from 4.7% to 8.5% over the past 30 years, and affects a higher number of overweight and obese individuals (WHO, 2016). In Korean adults 30 years of age or older, the prevalence of diabetes was 14.4% in 2016 (Korean Diabetes Association, 2018). Diabetes and the associated complications are major causes of mortality (Cowie et al., 2006). Depending on insulin dependency, diabetes can be categorized as type 1 or type 2 (American Diabetes Association, 2013). Type 2 diabetes (T2D) accounts for more than 90% of diabetic cases. In individuals with T2D, insulin is regularly secreted in the body, however target cells exhibit insulin resistance (IR) (American Diabetes Association, 2013). If left untreated, T2D can cause serious damage to nerves and blood vessels due to prolonged hyperglycemia (Kwon and Chung, 2013). Even in nondiabetic individuals, high blood glucose concentrations are considered a risk factor for mortality in middle-aged adults (Balkau et al., 1998). Thus, regulating glucose concentrations is important for maintaining good health.

Multiple genetic and environmental factors are major risk factors for T2D. Of these, obesity is one of the most significant determinants (Hu et al., 2001). Lack of exercise, high stress, and high fat and sugar intake are well-known factors that contribute to onset of T2D, and are characteristics of lifestyles that often lead to obesity. Therefore, interventions aimed at modulating these lifestyles, such as via diet and exercise, are critical for preventing T2D in populations at high risk of diabetes. Dietary intervention has been extensively studied in the context of prevention and treatment of T2D (Ben-Avraham et al., 2009; Schwingshackl et al., 2018). One of the most well-known strategies is modulating glycemic control (Stolar, 2010). High levels of fasting glucose can increase...
the risk of complications in diabetes patients. Thus, improving glycemic control is both a major goal of T2D therapy and a preventive strategy.

The effects of individual nutrients and functional compounds on regulating blood glucose levels for management of T2D have been previously reported (Heer and Egert, 2015; Russell et al., 2016; Alkhatib et al., 2017). For example, high fiber diets and soluble fiber supplements have been reported to significantly improve blood glucose levels in patients with T2D (Silva et al., 2013; Thompson et al., 2017). Recently, meal-based interventions has also emerged as a potential strategy for prevention of and treatment of T2D. Meal-based interventions can be used for long-term management of individuals at risk by modulating meal patterns, and as a short-term approach through supplementing meals with nutrients and bioactive compound. It is recognized that meal-based interventions have great potential for disease prevention, but are limited by practical obstacles associated with providing individuals with whole meals. Consequently, only a few studies have evaluated the efficacy of meal-based interventions on glucose regulation. Moreover, most the sensitivities of most clinical biomarkers for diabetes have not been fully validated in the context of assessing glucose changes in non-diabetic risk groups.

There are multiple clinical biomarkers currently available for the diagnosis and classification of diabetes (American Diabetes Association, 2013). Fasting plasma glucose (FPG) and hemoglobin A1c (HbA1c) levels are the most commonly used glycemic parameters. FPG levels are a key factor for determining threshold glucose levels, whereas HbA1c levels reflect average blood glucose levels over a period of two to three months. Currently, HbA1c is widely used as a marker of chronic glycemia (Kwon and Chung, 2013), and is used as a readout of glycemic control in the management of diabetes (American Diabetes Association, 2013). Further, hyperglycemic indicators, anthropometric measurements, insulin indexes, and lipid profiles are used to diagnose, classify, and monitor diabetes. While these markers are commonly used, it is not clear whether these markers are sufficiently sensitive to analyze changes in glucose homeostasis in nondiabetic individuals, particularly in response to dietary intervention.

In this review, we examined recent evidence regarding meal-based interventions and clinical biomarkers for intervention in both healthy individuals and individuals at risk of diabetes. This information should facilitate selection of appropriate clinical biomarkers and will help establish long-term prevention strategies for patients with T2D and individuals in target risk populations participating in meal-based interventions.

MATERIALS AND METHODS

Search strategy

Two search strategies were used to review publications reporting improvements in glucose regulation. First, we performed a search for meal-based dietary interventions in nondiabetic individuals through searching the PubMed database. The terms, ‘diet, glucose, and intervention’, was used to search publications published between 2014 and 2018. We then performed a second comprehensive literature search for clinical markers used to assess glucose regulation. The terms ‘clinical marker, assessment, and diabetes’ were used to search publications in MEDLINE through PubMed published between 2009 and 2018. Only articles published in English were considered. Relevant literature was extracted based on the title and abstract. Full text articles were subsequently checked according to the criteria listed in Fig. 1 and 2. A total of 29 articles reporting meal-based diet interventions and 17 articles reporting clinical biochemical markers were selected.

Selection criteria

For analysis of meal-based dietary interventions, all included studies were intervention studies that involved human subjects and used glucose regulation markers, such as FPG or HbA1c (Fig. 1). Following identification of relevant papers according to the titles, we excluded non-human studies, review papers, and non-meal-based studies. Next, the abstracts of the remaining papers were reviewed. Papers were excluded if results of glucose control were not provided. Finally, a total of 29 articles were selected for inclusion in this review.

Criteria for selecting articles involving clinical biomarkers of diabetes mellitus were included data derived from T2D patients and cohort studies that included adult with and without diabetes (Fig. 2). We excluded studies which 1) were derived from cross-sectional studies and case control studies, due to insufficient explanation for cause-and-effect relationships and difficulty in obtaining valid indicators to assess disease in time-specific investigations, 2) were identified in individuals with type 1 diabetes, gestational diabetes mellitus, and other diseases, and 3) were based on insufficient information regarding clinical markers. By screening titles and abstracts, articles were excluded based on unrelated diseases, non-human studies, review papers, and studies of individuals with other types of diabetes. Finally, a total of 17 articles were selected for this review.
RESULTS

Meal-based dietary interventions for improving glucose regulation

Initially, we categorized the 29 studies of meal-based dietary interventions for improving glucose regulation according to meal type: low caloric meals (n=5), macronutrient ratio-based meals (n=14), and food pattern studies (n=10). The low caloric meal intervention studies are summarized in Table 1. We observed that changes in FPG in obese individuals were not consistent between the different studies (Lee et al., 2014a; Perichart-Perera et al., 2014). Moreover, in the studies reporting low-calorie diets, 12-week, but not 6-month interventions, were shown to decrease FPG levels. However, very low-calorie restriction diet (e.g. low fat diets with a limitation of \( \sim 700 \text{ kcal/d} \)) significantly reduced FPG and HbA1c levels in both obese and T2D adults regardless of study
duration (Norén and Forssell, 2014; Steven and Taylor, 2015; Steven et al., 2016). Interestingly, the effect of very low-calorie restriction on HbA1c levels was not significant in adults who had diabetes for more than 8 years (Steven and Taylor, 2015), indicating that the effect depends on the severity of glucose impairment.

Recent evidence from multiple meal-based intervention studies examining various macronutrient ratios is summarized in Table 2. The ratio of the macronutrients, rather than calorie intake, determined cardiometabolic health and aging in ad libitum-fed mice (Solon-Biet et al., 2014). In addition, limited carbohydrate diets under calorie restriction, such as low carbohydrate and high protein diets, effectively lowered HbA1c and fasting glucose levels in both pre-diabetic and diabetic adults (Kempf et al., 2014; Tay et al., 2014; Goday et al., 2016); however, this result was not observed in nondiabetic overweight or obese adults (Rajaie et al., 2014). We also observed that these meal-based interventions with various macronutrient ratios affected both glucose markers and serum lipid profiles, including triglycerides (TG), low-density lipoprotein cholesterol (LDL-C), and high-density lipoprotein cholesterol (HDL-C). However, the latter affects are less consistent (Kempf et al., 2014; Tay et al., 2014; Goday et al., 2016; Stentz et al., 2016). Intriguingly, the effects of a two-year intervention involving a low carbohydrate and low saturated fat diet on HbA1c and fasting glucose levels did not significantly differ from effects observed following a high carbohydrate and low fat diet (Tay et al., 2018). It is possible that different ratios of carbohydrate and fat may not be critical to long-term intake. Ratios of macronutrients, even in the absence of calorie restriction, have been found to influence glucose and lipid metabolism (Table 2). Furthermore, low carbohydrate diets have been effective for decreasing HbA1c level in adult diabetes patients, although their effects on fasting glucose levels and serum lipid profiles have been relatively limited (Yamada et al., 2014; Sato et al., 2017; Wang et al., 2018).

The types of carbohydrates consumed, as well as the proportion of energy they represent in a diet, are critical for achieving low fasting glucose levels in gestational diabetic patients. Meanwhile, a high complex carbohydrate diet has been found to reduce fasting glucose levels compared with a control diet containing a relatively low carbohydrate content (Hernandez et al., 2016). Interestingly, the positive effects of a low carbohydrate/high fat diet (50~70% of total calories from fat) were not found to be significant in healthy and obese adults (Numao et al., 2016; Parry et al., 2017; Zinn et al., 2017; Parr et al., 2018).

Recently, intervention studies with various food patterns have been reported (Table 3). For example, in a four-year intervention study of adults with T2D adhering to a Mediterranean diet, HbA1c levels were markedly re-
Table 2. Intervention studies of various macronutrient ratio-based meals on blood glucose regulation

| Meal types of intervention (carbohydrate : fat : protein) | Subjects (country) | Duration of intervention | Results | Comments | Reference |
|----------------------------------------------------------|--------------------|--------------------------|---------|----------|-----------|
| Calorie restriction                                      |                    |                          |         |          |           |
| Moderately restricted carbohydrate diet (45:40:15)       | Overweight/obese adults | 6 weeks per diet (cross-over) | FPG: no change | 350–700 kcal less than the energy requirement | Rajaie et al. (2014) |
| Low carbohydrate diet (14:58:28) vs. high carbohydrate diet (53:30:17) | T2DM adults       | 6 months                 | FPG: no change | Low carbohydrate diet consisting of low-saturated fat | Tay et al. (2014) |
|                                                          |                    |                          | TG, HDL-C, LDL-C, and TC: no change | Calorie restriction with 500–1,000 kcal less than the energy requirement | |
|                                                          |                    |                          |          | Greater increase in HDL-C occurred with the low carbohydrate diet for participants with a baseline HDL-C < 1.3 mmol/L. | |
|                                                          |                    |                          |          | No diet effect in participants with initial HbA1c ≤ 62 mmol/mol | |
|                                                          |                    | 24 months                | FPG and HbA1c: no change | Very low caloric diets with 600–800 kcal/d <50 g carbohydrate from vegetables daily | Goday et al. (2016) |
|                                                          |                    |                          | HDL-C: increase | Low calorie diet with 500–1,000 kcal less than the energy requirement | |
|                                                          |                    |                          | TG: decrease | T2DM duration <10 years | |
|                                                          |                    |                          | TC, LDL-C, and non-HDL-C: no change |          |           |
| High protein diet (40:30:30)                             | Pre-diabetes adults | 6 months                | HbA1c, cholesterol, TG, and LDL-C: decrease | A data after 1.5 years of follow-up was available | Stentz et al. (2016) |
|                                                          |                    |                          | HDL-C: no change | Kemps et al. (2014) | |
|                                                          |                    |                          |          |          |           |
| High protein diet                                        | T2DM adults        | 3 months                 | FPG and HbA1c: decrease |          |           |
|                                                          |                    |                          | HDL-C: increase |          |           |
|                                                          |                    |                          | TG: decrease |          |           |
|                                                          |                    |                          | TC and LDL-C: no change |          |           |
| Very low carbohydrate (ketogenic) diet compared to low calorie diet | T2DM adults       | 4 months                 | FPG, HbA1c, and TG: decrease |          |           |
|                                                          |                    |                          | TC, HDL-C, LDL-C: no change |          |           |
|                                                          |                    |                          |          |          |           |
| Meal types of intervention (carbohydrate : fat : protein) | Subjects (country) | Duration of intervention | Results | Comments | Reference |
|--------------------------------------------------------|------------------|--------------------------|---------|----------|-----------|
| Non-calorie restriction                                 | Healthy adults   | 7 days                    | FPG: increase | 65% total energy from fat | Parry et al. (2017) |
| Low carbohydrate-high fat diet (20:65:15)              |                  |                          | HDL-C: increase |                        |           |
|                                                        |                  |                          | TG: decrease |                        |           |
|                                                        |                  |                          | TC and LDL-C: no change |                        |           |
| Low carbohydrate-high fat diet (20:70:10) compared to control diet (60:30:10) |                  | 3 days per diet (cross-over) | TC, HDL-C, and LDL-C: no change | FPG and TG were significantly decreased comparing baseline to post-intervention. | Numao et al. (2016) |
| Low carbohydrate-high fat diet (40:50:10) compared to control diet (60:30:10) |                  |                          | TG: decrease |                        |           |
| Low carbohydrate-high fat diet (40:50:10)              | Overweight/obese adults | 12 weeks                | HDL-C: increase | Carbohydrate intake less than 45% of total energy | Zinn et al. (2017) |
|                                                        |                  |                          | Serum glucose, TG: decrease | Fat intake more than 33% of total energy |           |
|                                                        |                  |                          | HbA1c and LDL-C: no change |                        |           |
| Low carbohydrate-high fat diet (15:67:18) compared to low fat diet (67:15:18) |                  | 5 days per diet (cross-over) | Mean glucose and AUC: decrease |                        | Parr et al. (2018) |
| High complex-carbohydrate diet (60:25:15)              | Gestational diabetes | 7 weeks                | TG: increase | High carbohydrate was based on complex carbohydrate. | Hernandez et al. (2016) |
|                                                        |                  |                          | FPG: decrease |                        |           |
|                                                        |                  |                          | TC, HDL-C, and LDL-C: no change |                        |           |
| Low carbohydrate diet                                  | T2DM adults (Japan) | 6 months               | HbA1c: decrease | Carbohydrate intake: 130 g/d | Sato et al. (2017) |
|                                                        |                  |                          | TG, HDL-C, and LDL-C: no change | A data after a year of follow-up was available. |           |
|                                                        | T2DM adults      | 6 months               | HbA1c and TG: decrease | Carbohydrate intake less than 130 g/d | Yamada et al. (2014) |
|                                                        | (China)          |                          | FPG: no change |                        |           |
|                                                        |                  |                          | LDL-C and HDL-C: no change |                        |           |
| Low carbohydrate diet (38:42:20)                       | T2DM adults      | 3 months               | FPG and HbA1c: decrease |                        | Wang et al. (2018) |
|                                                        | (China)          |                          | TC: decrease |                        |           |

T2DM, type 2 diabetes mellitus; FPG, fasting plasma glucose; TG, triglycerides; HDL-C, high-density lipoprotein cholesterol; LDL-C, low-density lipoprotein cholesterol; TC, total cholesterol; HbA1c, hemoglobin A1c; AUC, area under the curve.
Table 3. Intervention studies of various food patterns on blood glucose regulation

| Meal types of intervention | Subjects (country) | Duration of intervention | Change of phenotype | Comments | Reference |
|----------------------------|--------------------|--------------------------|---------------------|----------|-----------|
| Mediterranean diet         | T2DM adults        | 48 months                | HbA1c: decrease     | 1,500 kcal/d for women, 1,800 kcal/d for men | Esposito et al. (2014) |
|                           | Obese children and adolescents | 4 months | FPG: decrease, HDL-C: increase, TC, TG, and LDL-C: decrease | | Velázquez-López et al. (2014) |
| Low GI diet compared to low calorie diet | Obese children | 6 months | FPG: no change, TC, TG, HDL-C, and LDL-C: no change | No change within the group | Visuthranukul et al. (2015) |
| Low GI diet compared to high GI diet | Healthy male adults (China) | 1 day per diet (cross-over) | 24 h glucose iAUC: decrease | | Camps et al. (2017) and Henry et al. (2017) |
| Mexican diet compared to US diet | Healthy female adults (Mexico) (partially overweight) | 24 days per diet (cross-over) | FPG: no change | | Santiago-Torres et al. (2016) |
| Okinawan-based Nordic diet | T2DM adults (Scandinavia) | 3 months | HDL-C: increase in follow-up, FPG and HbA1c: decrease, TG, TC, and LDL-C: decrease | A data after four months of follow-up was available. | Darwiche et al. (2016) |
| DASH diet specifically modified for diabetes | T1DM adolescents | 3 days | Glycemic viability: no change | | Peairs et al. (2017) |
| High-red meat diet compared to high-dairy diet | Overweight/obese adults | 4 weeks per diet (cross-over) | FPG: no change | | Turner et al. (2015) |
| Plant-based burger compared to processed meat burger | T2DM and healthy adults | 1 day (several times) | Postprandial plasma glucose: increased | Increased only one time point | Belinova et al. (2014) |

GI, glycemic index; T2DM, type 2 diabetes mellitus; T1DM, type 1 diabetes mellitus; HbA1c, hemoglobin A1c; FPG, fasting plasma glucose; HDL-C, high-density lipoprotein cholesterol; TC, total cholesterol; TG, triglycerides; LDL-C, low-density lipoprotein cholesterol; iAUC, incremental area under the curve; DASH, dietary approaches to stop hypertension; NHANES, National Health and Nutrition Examination Survey.
duced (Esposito et al., 2014). Low glycemic index (GI) diets, conventionally recommended for diabetes patients, have been shown to significantly lower 24 h glucose incremental area under the curve data for both diabetes patients and healthy adults, as compared with high GI diets (Camps et al., 2017; Henry et al., 2017). However, these diets have exhibited different effects on glucose regulation in obese children and adolescents. The Mediterranean diet has been shown to decrease glucose levels and serum lipid profiles (Velázquez-López et al., 2014), whereas low GI diets have shown no effect on FPG levels and serum lipid profiles in obese children (Visuthranukul et al., 2015). In addition, a dietary approaches to stop hypertension diet did not change glycemic viability in type 1 diabetes adolescents (Peairs et al., 2017), and a Mexican diet did not affect FPG compared to a US diet in healthy adults (Santiago-Torres et al., 2016). Culturally modified diet patterns, such as the Okinawan-based Nordic diet, have exhibited potential for reducing FPG and HbA1c levels in adults with T2D (Darwiche et al., 2016). Meanwhile, a high-red meat diet and a high-dairy diet did not induce differences in fasting glucose level in obese adults (Turner et al., 2015). Finally, plant-based burgers have been shown to induce limited changes to postprandial plasma glucose levels in both healthy adults and adults with T2D (Belinova et al., 2014).

Taken together, calorie restriction and macronutrient ratio adjustments, such as low carbohydrate diets, have been shown to be effective for reducing fasting glucose and HbA1c levels in diabetic patients. However, these effects are limited in both obese and healthy adults. Specifically, HbA1c and fasting glucose levels are altered in obese and healthy subjects following short-term interventions, yet similar results are not shown following long-term interventions. Thus, it appears that clinical biomarkers for glucose regulation may need to be re-evaluated for interventions involving non-diabetic subjects.

### Clinical biomarkers for improved glucose regulation

A total of 17 studies were reviewed to evaluate currently available clinical markers for improved glucose regulation. In these studies, diagnostic criteria for diabetes were mainly based on criteria published by the World Health Organization or by the American Diabetes Association. The former includes FPG $\geq 7.0$ mmol/L (126 mg/dL) or a 2-hr plasma glucose $\geq 11.1$ mmol/L (200 mg/dL) (WHO, 2006), and the latter includes FPG $\geq 126$ mg/dL (7.0 mmol/L), an HbA1c $\geq 6.5\%$, a 2-hr plasma glucose level $\geq 200$ mg/dL, or a random plasma glucose $\geq 200$ mg/dL (11.1 mmol/L) with symptoms of hyperglycemia or hyperglycemic crisis (American Diabetes Association, 2013).

Clinical markers considered in this review are associated with diabetes-related phenotypes, including IR. In addition, conventionally used anthropometric markers were considered, such as height, weight, waist circumference, hip circumference, waist-hip circumference ratio, body mass index (BMI), and blood pressure. Clinical markers of serum parameters [e.g., fasting blood glucose, fasting insulin, homeostatic model assessment (HOMA)-IR, HDL-C, LDL-C, TC, and TG] were used in most of the human intervention studies that assessed glucose regulation. In addition, triglyceride glucose (TyG) indexes and insulinogenic indexes (IGI) were reported. TyG indexes are accepted as a surrogate marker for IR (Roth et al., 2017), whereas IGI represent insulin levels in response to changes in serum glucose (Singh and Saxena, 2010).

In diabetic adults, a more conserved set of clinical biomarkers have been used, including BMI as an anthropometric marker, and insulin, glucose, cholesterol, and HOMA as biochemical markers (Ren et al., 2016) (Table 4). In addition, IGI and insulin sensitivity indexes were used (Chung et al., 2012). Novel markers, including activin A, activin B, and follistatin, have also been reported as useful indicators for the severity of T2D and IR (Wu et al., 2012). Activin A is a member of the transforming growth factor $\beta$ superfamily and enhanced activin A activity has been associated with T2D (Ueland et al., 2012). Follistatin is a glycosylated plasma protein that plays a role in IR (Hansen et al., 2013). Both glycoprotein acetylation and leptin have been associated with risk and incidence of T2D (Welsh et al., 2009; Connelly et al., 2016).

Clinical indicators associated with diabetes-related phenotypes in nondiabetic and diabetic subjects was summarized in Table 5. Of the various anthropometric markers, BMI followed by waist circumference is the most widely used for both nondiabetic and diabetic subjects. In addition to conventional clinical markers (e.g., glucose, insulin, HDL-C, LDL-C, TC, TG, and HOMA index), alanine transaminase (ALT), and gamma-glutamyl transferase (GGT) have also been used as biomarkers to assess diabetes risk. Higher levels of both ALT and GGT are detected in diabetic compared with non-diabetic individuals (Nguyen et al., 2011; Kurosaki et al., 2014). In another study, levels of HbA1c, TC, LDL-C, high sensitivity C-reactive protein (hsCRP), HOMA-IR, interleukin (IL)-6, tumor necrosis factor (TNF), and IL-12 were higher in diabetic compared with non-diabetic individuals (Mishra et al., 2011). However, in another study, levels of adipin and IL-1$\beta$ were lower and higher, respectively, in diabetic compared with non-diabetic individuals (Zhou et al., 2018). Adipin is one of the major proteins in adipose cells and is useful for identifying diabetic patients at high risk of beta cell failure. In contrast with GIs, such as HbA1c levels, which are well controlled, levels of adipin have been proposed to provide more sensitive monitoring during the introduction of insulin therapy (Lo et al., 2014). Higher levels of ferritin and IL-2 receptor alpha have been observed in individuals with diabetes, whereas...
| Subjects (country) | Investigated markers | Major outcome markers | Diagnosis standard | Reference |
|-------------------|----------------------|-----------------------|--------------------|-----------|
| T2DM (China)      | Anthropometric markers: Height, weight, WC, HC, WHR, BMI, and BP Clinical markers: ALT, HbA1c, HOMA-IR, HOMA-β, HDL-C, LDL-C, TC, TG, FPG, 2hPG, FINS, 2hPINS, IGI, and ISI | - TG/HDL-C ratio<br>- High TG/HDL-C ratio: ALT, HbA1c, HOMA-IR, HOMA-β, HDL-C, LDL-C, TC, TG, FPG, FINS, IGI, ISI increase<br>- High TG/HDL-C ratio: HDL-C decrease | World Health Organization | Ren et al. (2016) |
| T2DM (Korea)      | Anthropometric markers: BMI and BP Clinical markers: HbA1c, HOMA-IR, HOMA-β, HDL-C, TC, TG, fasting glucose, fasting insulin, 2-h glucose, 2-h insulin, DI, IGI, and ISI comp | - BMI<br>- BMI: TG, FPI, fasting insulin, 2-h insulin, HOMA-IR, HOMA-β, and IGI: increase<br>- BMI: DI decrease | American Diabetes Association | Chung et al. (2012) |
| T2DM (Australia)  | Anthropometric markers: WHR and BMI Clinical markers: HbA1c, HOMA-IR, fasting glucose, fasting insulin, HDL-C, LDL-C, TC, TG, Crc, hscRP, activin A, activin B, and follistatin | - Crc, HDL-C, LDL-C, TC, TG, hscRP, fasting glucose, HbA1c, fasting insulin, and HOMA-IR<br>- Markers of the severity of T2D and insulin resistance: activin A, activin B, or follistatin | American Diabetes Association and World Health Organization | Wu et al. (2012) |
| Diabetes (Netherlands) | Clinical markers: GlycA and hscRP | - High GlycA and low hscRP: incidence of T2D | - One or more of the following:<br>1) FPG ≥7.0 mmol/L (126 mg/dL)<br>2) Random sample FPG ≥11.1 mmol/L (200 mg/dL)<br>3) Self-report of a physician diagnosis of T2DM<br>4) Initiation of glucose-lowering medication use, retrieved from a central pharmacy registry | Connelly et al. (2016) |
| Diabetes (Scotland, Ireland, or Netherlands) | Clinical markers: Leptin | - Leptin | - Based on baseline glucose measurements<br>(>95% fasting) | Welsh et al. (2009) |

T2DM, type 2 diabetes mellitus; WC, waist circumference; HC, hip circumference; WHR, waist-hip ratio; BMI, body mass index; BP, blood pressure; ALT, alanine transaminase; HbA1c, hemoglobin A1c; HOMA-IR, homeostatic model assessment of insulin resistance; HOMA-β, homeostatic model assessment of β-cell function; HDL-C, high-density lipoprotein cholesterol; LDL-C, low-density lipoprotein cholesterol; TC, total cholesterol; TG, triglycerides; FPG, fasting plasma glucose; 2hPG, 2-hour postprandial plasma glucose; FINS, fasting serum insulin; 2hPINS, 2-hour postprandial serum insulin; IGI, insulogenic index; ISI, insulin sensitivity index; DI, disposition index; ISIcomp, insulin sensitivity index composite; Crc, creatinine clearance; hsCRP, high sensitivity C-reactive protein; GlycA, glycoprotein acetylation.
| Subjects (country) | Investigated markers | Major outcome markers | Diagnosis standard | Reference |
|-------------------|-----------------------|-----------------------|--------------------|-----------|
| **Normoglycemic, prediabetic, and diabetic adults (USA)** | **Anthropometric markers** | BMI, BP, and MAP | Clinical markers | HDL-C, LDL-C, TG, glucose, insulin, HOMA-IR, ALT, and GGT | Insulin, LDL-C, TG, and HOMA-IR: high (in diabetic) | American Diabetes Association |
| **Nondiabetic and diabetic subjects (Japan)** | **Anthropometric markers** | BMI | Clinical markers | HbA1c, HOMA-IR, fasting insulin, fasting glucose, HDL-C, LDL-C, RLP-C, TC, TG, pre heparin LPL mass, and VLDL/TG | Fasting glucose, HbA1c, fasting insulin, HOMA-IR, TG, RLP-C, and VLDL/TG: high (in diabetic) | Kurosaki et al. (2014) |
| **Healthy and diabetic subjects (India)** | **Anthropometric markers** | Height, weight, WC, HC, WHR, BMI, SBP, DBP | Clinical markers | FBS, PBP, insulin, HbA1c, HOMA-IR, hsCRP, HDL-C, LDL-C, VLDL-C, TC, TG, NOx, ICAM, VCAM, IL-6, TNF, and IL-12 | HbA1c, FBS, PBP, TC, LDL-C, hsCRP, HOMA-IR, IL-6, TNF, and IL-12: high (in diabetic) | American Diabetes Association |
| **Healthy and diabetic subjects (China)** | **Anthropometric markers** | Height, weight, WC, HC, WHR, BMI, and BP | Clinical markers | FFA, FPG, FINS, HbA1c, hsCRP, HOMA-IR, 2hPG, HDL-C, LDL-C, TC, TG, Adipsin, and IL-1β | FFA, FPG, 2hPG, FINS, HbA1c, HOMA-IR, HOMA-β, TG, hsCRP, and IL-1β: high (in diabetic) | World Health Organization |
| **Healthy and diabetic subjects (Denmark)** | **Anthropometric markers** | Height, weight, HC, BMI, SBP, and DBP | Clinical markers | HbA1c, FPG, fasting insulin, adiponectin, CRP, ferritin, and IL2Ra | HbA1c, FPG, fasting insulin, and CRP: high (in diabetic) | American Diabetes Association |

**BMI, body mass index; BP, blood pressure; MAP, mean arterial pressure; HDL-C, high-density lipoprotein cholesterol; LDL-C, low-density lipoprotein cholesterol; TG, triglycerides; HOMA-IR, homeostatic model assessment of insulin resistance; ALT, alanine transaminase; GGT, gamma-glutamyl transferase; HbA1c, hemoglobin A1c; RLP-C, remnant-like particle cholesterol; TC, total cholesterol; LPL, lipoprotein lipase; VLDL, very low-density lipoprotein; WC, waist circumference; HC, hip circumference; WHR, waist-hip ratio; SBP, systolic blood pressure; DBP, diastolic blood pressure; FBS, fasting blood sugar; PBP, post prandial blood sugar; hsCRP, high sensitivity C-reactive protein; NOx, nicotinamide adenine dinucleotide phosphate hydrate oxidase; ICAM, intercellular adhesion molecule; VCAM, vascular cell adhesion molecule; IL, interleukin; TNF, tumor necrosis factor; FPG, fasting plasma glucose; FINS, fasting serum insulin; FFA, fasting fatty acid; 2hPG, 2-hour postprandial plasma glucose; IL2Ra, interleukin-2 receptor alpha.**
higher levels of adiponectin have been observed in indi-
viduals without diabetes (Lyssenko et al., 2012). A posi-
tive correlation between plasma ferritin and fasting insu-
lin and glucose levels have also been reported (Fumeron et 
al., 2006), with elevated ferritin concentrations associ-
ated with increased risk of diabetes. Thus, serum ferritin 
is considered to be an independent predictor of diabetes 
(Foroohi et al., 2007).

We also reviewed clinical markers as indicators for con-
fiming the incidence of diabetes in non-diabetic groups 
(Table 6). TyG indexes are a valuable surrogate marker 
for the degree of IR and for predicting diabetes incidence. 
Correspondingly, TyG levels are low in non-diabetic sub-
jects (Lee et al., 2014b). Red cell distribution width 
(RDW) is also an easily measured marker and is posi-
tively associated with HbA1c. Thus, it has been proposed 
that RDW potentially represents a biomarker for risk of 
diabetes (Engström et al., 2014). High levels of biochem-
ical parameters, such as blood urea nitrogen (BUN), es-
timated glomerular filtration rate (eGFR), apolipoprotein 
(Apo) A1, leukocyte, sCD163, and soluble plasminogen 
activator urokinase receptor (suPAR), have been asso-
ciated with risk of diabetes (Haugaard et al., 2012; 
Møller et al., 2011; Xie et al., 2018), and are therefore 
useful for predicting diabetes incidence. In contrary, 
eGFR levels have been negatively associated with risk of 
diabetes (Xie et al., 2018).

In this review, clinical markers of T2D and interven-
tions for preventing diabetes in non-diabetic groups were 
investigated. Currently, clinical markers of diabetes in-
clude well-known indicators of anthropometric measure-
ments, blood samples, and indices. Of these, activin A, 
activin B, and follistatin are useful for determining the se-
verity of diabetes. Hematological and biochemical mark-
ers used to identify the onset of diabetes in non-diabetic 
(healthy subjects) include RDW, hemoglobin (Hgb), he-
matocrit (Hct), BUN, eGFR, Apo A1, Apo B, sCD163, 
and suPAR. Finally, fasting glucose and HbA1c levels 
have been confirmed as appropriate markers for identify-
ing development of diabetes in non-diabetic subjects.

**DISCUSSION**

Recent evidence from meal-based interventions supports 
the following results: 1) very low-calorie diets greatly im-
prove glucose regulation in obese and adults with T2D, 
whereas low-calorie diets have no effect on obese adults; 
2) calorie restriction and the ratio of macronutrients, 
such as low carbohydrate ratios, are effective for reduc-
ing fasting blood glucose and HbA1c levels in diabetes 
patients, although the effects are limited in obese and 
healthy adults; 3) glucose-related markers change in 
obese and healthy subjects following short-term interven-
tions, yet these effects are not observed following long-
term interventions; 4) Mediterranean and low GI diets 
are effective for lowering fasting blood glucose and/or 
HbA1c levels in healthy and T2D adults.

For meal-based interventions, it is necessary to consid-
er the ratios used to define high or low carbohydrate and 
fat macronutrient compositions. An additional and im-
portant consideration for interpreting meal-based inter-
vention data is the cultural aspect of the food. Variations 
in food patterns and cooking methods in different cul-
tures may affect the observed results (Knowler et al., 
2002). For example, conventional macronutrient ratios 
vary between Asian and Western cuisines. In Korea, 
meals with a relatively high carbohydrate content are 
common, and the Korean Dietary Reference Intake rec-
ommends that 55∼65% of total energy should be derived 
from carbohydrates and 15∼30% should be derived from 
fat (Korean Nutrition Society, 2015). However, North 
American meals include relatively high proportions of 
protein and fat, and the US Dietary Reference Intake rec-
ommends that 45∼65% of total energy should come 
from carbohydrates and 20∼35% of should come from fat 
(USDA, 2015). Therefore, results from meal-based inter-
ventions for risk groups should be carefully interpreted.

Of note, the observed effects of meal-based interven-
tions on glucose regulation differed between diabetic pa-
tients and healthy or nondiabetic obese subjects. Specifi-
cally, HbA1c and fasting glucose levels were often only af-
fected by short-term meal-based interventions in healthy 
or nondiabetic obese subjects. This limited effect may be 
due to sensitivities of the markers used. Because clinical 
markers for glucose regulation mostly target T2D, more 
sensitive and appropriate markers should be identified 
to evaluate prevention in healthy subjects.

FPG and HbA1c indicators are mainly used to identify 
changes in glycemic control in response to meal-based in-
terventions. These indicators are used as diagnostic cri-
tera for diabetes (American Diabetes Association, 2013) 
and as representative indicators of changes to blood sugar 
control. BMI is an anthropometric marker and a manda-
tory indicator of T2D. In particular, BMI ≥25 kg/m² indi-
cates an increased risk of T2D (Hsu et al., 2015). In gen-
eral, fasting insulin, HOMA-IR, and HOMA of β-cell func-
tion are recognized as major clinical markers of diabetes, 
with high levels of these indicators present in diabetes 
patients (Haffner et al., 1996; Tangvrasititchai et al., 
2010; Wang et al., 2011). In addition, lipid profiles of TC, 
TG, LDL-C, and HDL-C are used as basic markers for 
diabetes. For example, individuals with diabetes exhibit 
higher levels of triglycerides, LDL-C, and cholesterol, and 
lower levels of HDL-C, compared with healthy controls 
(Tangvrasititchai et al., 2010; Uttra et al., 2011); these 
markers are all significant indicators of diabetes. Other 
clinical indicators, such as white blood cell, red blood cell,
| Subjects (country) | Investigated markers | Major outcome markers | Diagnosis standard | Reference |
|-------------------|----------------------|-----------------------|--------------------|-----------|
| Nondiabetic (Canada) | **Anthropometric markers** | Height, weight, WC, BMI, and BP | **Clinical markers** | Hct, Hgb, RBC, WBC, HOMA-IR, fasting glucose, 2-h glucose, fasting insulin, IGI, ISoctt, IGI/HOMA-IR, ISSI-2, normal glucose tolerance, impaired glucose test, and impaired fasting glucose | World Health Organization | Hanley et al. (2009) |
| Adults without diabetes (USA) | **Anthropometric markers** | BMI | **Clinical markers** | BUN and eGFR | - BUN: high incidence of diabetes | Diabetes medication prescription (including insulin and oral hypoglycemic agents); or an HbA1c test result >6.4% | Xie et al. (2018) |
| Adults without diabetes (China) | **Anthropometric markers** | BMI, SBP, and DBP | **Clinical markers** | AST, ALT, creatinine, fasting glucose, ApoA1, HDL-C/ApoA1 ratio, ApoB, HDL-C, LDL-C, TC, and TG | - ALT, fasting glucose, creatinine, and TG: decrease (in nondiabetic) | Fasting glucose ≥7.0 mmol/L or if subjects were receiving active treatment for T2DM | Wu et al. (2017) |
| Nondiabetic subjects (Korea) | **Anthropometric markers** | Height, weight, WC, HC, WHR, BMI, SBP, and DBP | **Clinical markers** | FPG, Fasting insulin, HOMA-IR, HOMA-β, HDL-C, LDL-C, TC, TG, and serum creatinine | - FPG, fasting insulin, HOMA-IR, TC, and TG: low (in nondiabetic) | American Diabetes Association | Lee et al. (2014b) |
| Nondiabetic subjects (Sweden) | **Anthropometric markers** | Height, weight, WC, and BMI | **Clinical markers** | Insulin, HOMA index, TG, RDW, MCV, RBC, and leukocyte count | - High RDW: insulin, glucose, TG, and HOMA index decrease | FPG ≥7.0 mmol/L | Engström et al. (2014) |
| Healthy adults (Denmark) | **Anthropometric markers** | BMI, SBP, DBP, WHR | **Clinical markers** | Glucose, hsCRP, fibrinogen, α1-antitrypsin, orosomucoid, TC, HDL-C, LDL-C, ApoA1, ApoB, TG, and scD163 | - High scD163 (in diabetes): predicted increased risk of diabetes | World Health Organization | Möller et al. (2011) |
| Healthy adults (Denmark) | **Anthropometric markers** | WC, WHR, BMI, SBP, and DBP | **Clinical markers** | Plasma glucose, serum insulin, HOMA-IR, suPAR, CRP, leukocytes, HDL-C, LDL-C, TC, and TG | - Leucocytes, suPAR, and CRP: low (in nondiabetic) | International Classification of Disease or FPG ≥6.9 mmol/L or use of antidiabetic drugs | Haugaard et al. (2012) |

WC, waist circumference; BMI, body mass index; BP, blood pressure; Hct, hematocrit; Hgb, hemoglobin; RBC, red blood cell; WBC, white blood cell; HOMA-IR, homeostatic model assessment of insulin resistance; IGI, insulinogenic index; ISoctt, insulin sensitivity index for oral glucose tolerance tests; ISSI-2, insulin secretion sensitivity index-2; BUN, blood urea nitrogen; eGFR, estimated glomerular filtration rate; HbA1c, hemoglobin A1c; SBP, systolic blood pressure; DBP, diastolic blood pressure; AST, aspartate aminotransferase; ALT, alanine transaminase; Apo, apolipoprotein; HC, hip circumference; HDL-C, high-density lipoprotein cholesterol; LDL-C, low-density lipoprotein cholesterol; TC, total cholesterol; TG, triglycerides; T2DM, type 2 diabetes mellitus; WHR, waist-hip ratio; FPG, fasting plasma glucose; HOMA-β, homeostatic model assessment of β-cell function; TyG index, triglyceride and glucose index; MCV, mean corpuscular volume; RDW, red cell distribution width; suPAR, soluble urokinase-type plasminogen activator receptor.
Hgb, Hct, Apo A1, Apo B, sCD163, and suPAR, also exhibit associations with diabetic risk (Hanley et al., 2009; Møller et al., 2011; Haugaard et al., 2012; Wu et al., 2017).

Several markers are regarded as indicators for diagnosing the severity of diabetes and for predicting the incidence of diabetes. For example, activin A, activin B, and follistatin have been reported to be useful markers to assess the severity of T2D (Wu et al., 2012). In addition, TyG indexes have been reported as a novel marker for IR in healthy adults. As such, TyG has been proposed to serve as a surrogate marker for predicting further risk of diabetes (Lee et al., 2014b). It is important to appropriately select clinical indicators that can predict the onset and estimate the severity of diabetes. Recently, non-targeted metabolite profiling (Hanhineva et al., 2015) and continuous 24 h glucose measurements have been used to diagnose risk of diabetes in non-diabetic healthy subjects (Philippou et al., 2008). Such methods profile biological changes, rather than serving as biomarkers, to identify development of diabetes. Despite such limitations, these methods can monitor overall changes and provide strong data regarding dietary intake (Nansel et al., 2016). Thus, these methods are candidates for identifying onset of diabetes in non-diabetic subjects.

The effects of meal-based dietary interventions on glucose regulation can differ according to the characteristics of the subjects and factors such as the type of the intervention (e.g., meal type and duration). For example, FPG levels were not changed following a 6-month low-calorie diet intervention in obese adult women, but were decreased following a 4-weeks very low-calorie diets in obese adults (Norén et al., 2014; Perichart-Perera et al., 2014). However, general biomarkers for glucose regulation, such as Hba1c and HOMA-IR, are higher in diabetic individuals compared with healthy controls in most studies (Mishra et al., 2011; Kurosaki et al., 2014; Zhou et al., 2018). Therefore, more thorough investigation of appropriate clinical biomarkers is required.

This review has several limitations. First, it was not designed as a systematic review so did not produce comprehensive evidence. Second, studies examining meal-based interventions were based on articles published within the last five years to provide recent examples, whereas studies examining clinical biomarkers were screened from an extended period of time, i.e. the last decade. These screening cut-off’s may constrain the wider implications of this review. In future, a systematic and comprehensive review may be needed.

Overall, this review presents recent evidence from the effects of meal-based dietary interventions on glucose homeostasis in diabetic risk groups, and clinical biomarkers used to assess diabetic risk, and to predict the onset and severity of diabetes. It is anticipated that the insights presented in this review will guide further investigations of meal-based preventive approaches in both non-diabetic and prediabetic populations.

AUTHOR DISCLOSURE STATEMENT
The authors declare no conflict of interest.

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