Minimizing Glycemic Fluctuations in Patients with Type 2 Diabetes: Approaches and Importance

Paresh Dandona, MD, PhD

Abstract

Glycemic fluctuations, characterized by short-term oscillations in plasma glucose, are important when managing type 2 diabetes (T2D) and may be considered a target of glucose-lowering therapies. Continuous glucose monitoring (CGM) has been used to evaluate the effects of different treatments on glycemic fluctuations. This review examines approaches to and the importance of minimizing glycemic fluctuations among patients with T2D. Measures of HbA1c, fructosamine, and glycated albumin reflect a long-term average of plasma glucose, and are therefore unable to provide an accurate measure of short-term glycemic oscillations. CGM provides accurate monitoring of real-time glucose fluctuations and has been used to investigate the effects of lifestyle and treatment on daily glycemic control. Dipeptidyl peptidase-4 inhibitors, sodium–glucose cotransporter 2 inhibitors, and glucagon-like peptide-1 receptor agonists have demonstrated significant improvements in measures such as the mean amplitude of glucose excursions and standard deviation of CGM. Case studies of two patients with T2D utilizing CGM are also included in this review, which demonstrated that CGM was a useful tool for diagnosing unrecognized hypoglycemia and hyperglycemia in situations in which it was impractical to check fingerstick concentrations. Altogether, the evidence suggests that glycemic fluctuations are a potential target to consider when managing T2D. CGM allows for the real-time evaluation of glycemic fluctuations and may assist in the development of an individualized treatment plan to adequately control short-term oscillations in blood glucose levels.

Keywords: Glucose, Glycemic control, Glycemic fluctuations, Glycemic variability, Type 2 diabetes.

Introduction

Effectively controlling blood glucose levels is an important aspect of clinical care in the long-term management of diabetes (both type 1 diabetes [T1D] and type 2 diabetes [T2D]) because it reduces the risk of microvascular complications. However, short-term changes in blood glucose levels, termed glycemic fluctuations, can impact mood and quality of life (QoL). Glycemic fluctuations may also contribute to the development of vascular complications in patients with T2D, which remain the leading cause of morbidity and mortality in this patient population. Glycemic fluctuations have traditionally been studied in the context of insulin-treated T1D, but recent research and technology have demonstrated the benefit of monitoring and controlling glycemic fluctuations in T2D as well. This review discusses the importance of glycemic control and glycemic fluctuations in the management of patients with T2D.

Glycated Hemoglobin: Can We Do Better Than HbA1c?

HbA1c is a measure of the fraction of glycated hemoglobin, which increases with average plasma glucose. As the lifespan of red blood cells is ~ 4 months, HbA1c reflects average blood glucose levels over the past 2–3 months. HbA1c is widely used to assess long-term glycemic control in patients with T2D and has been the focus of many glucose-lowering therapies whereby the therapeutic goal is to achieve target HbA1c levels of <7.0% or ≤6.5%, as recommended by international guidelines. However, HbA1c changes slowly in response to...
treatment, and conditions such as hemolytic or hemorrhagic anemia, kidney disease, liver disease, blood transfusions, and altered red blood cell lifespan can affect its validity.9,10

Fructosamine and glycated albumin provide an index of glucose control over 2–3 weeks, rather than the 3-month average provided by HbA1c.11 Therefore, these measures may be a preferable alternative to HbA1c, particularly in patients with chronic kidney disease who undergo hemodialysis, which has been shown to reduce HbA1c values.12 A study by Selvin et al. demonstrated that glycated albumin and fructosamine were strongly associated with microvascular complications among patients with T2D, and associations persisted after adjusting for HbA1c.13 However, fructosamine and glycated albumin measurements have limitations as well; fructosamine measurements in patients with abnormal albumin turnover and hypoalbuminemia may be inaccurate,14 and glycated albumin levels are affected by dysregulated albumin metabolism and hypoalbuminemia.17

Glycemic Fluctuations: A Better Picture of Glycemic Control?

HbA1c, fructosamine, and glycated albumin all reflect average blood glucose levels over weeks or months. However, in patients with T2D, blood glucose oscillations can occur within a single day. These short-term oscillations are termed glycemic fluctuations or variability. Specifically, glycemic fluctuations are characterized by variations in blood glucose levels (glycemic excursions) and include hypoglycemic events and postprandial hyperglycemia.14 Since HbA1c does not reflect these short-term oscillations, patients with T2D may achieve a target HbA1c while still experiencing marked glycemic fluctuations.

Short-term changes in blood glucose can impact several functions. During acute hyperglycemia (mean blood glucose of 301 mg/dL), 20 adults with T2D were found to have impaired performance on tests of information processing, working memory, and attention.15 Tension increased during acute hyperglycemia, while hedonic tone and energetic arousal (feelings of happiness and alertness, respectively) decreased. Furthermore, for 45 patients with T2D, the rate of change of postmeal blood glucose was shown to correlate with depressive and anxious mood and symptoms of cognitive dysfunction.16 Glycemic fluctuations are also associated with QoL and patient-reported outcomes. In patients with T1D (n = 82) and insulin-treated T2D (n = 306), reductions in intra-day mean glucose, glycemic fluctuations, and excursions above 140 mg/dL, as measured by continuous glucose monitoring (CGM), were associated with improvements in patient satisfaction and perceived health, independent of concurrent reductions in daily glucose levels and HbA1c.17 In patients with T2D (n = 54) initiating insulin pump therapy, a reduction in glycemic fluctuations was associated with increased treatment preference and improved health utility (a measure of QoL based on mobility, self-care, activities, pain/discomfort, and anxiety/depression), which the authors believe may lead to enhanced treatment adherence.18 Across treatments, some physicians and patients believe that better QoL may potentially improve adherence to treatment,19 whereas others feel that enhanced treatment adherence may contribute to an improved QoL.20,21 Therefore, control of short-term changes in blood glucose has important effects on a patient’s day-to-day life and may contribute to a positive feedback loop of improved glycemic control, treatment adherence, and QoL.

Glycemic fluctuations also play a role in the long-term management of diabetic complications. In a recent editorial, Hirsch22 noted that, as the treatment of T1D (and often patients with a long duration of T2D) shifted from once- or twice-daily insulin to multiple daily injections or a continuous subcutaneous insulin infusion, approaches that more closely mimicked physiologic insulin concentrations and likely reduced glycemic oscillations, the occurrence of microvascular complications declined. Hirsch hypothesized this was an early indicator that glycemic fluctuations may be related to diabetic complications.

This is further supported in a review by Yu et al., which suggests that postprandial hyperglycemia—a key contributor to glycemic fluctuations—is a risk factor for macrovascular complications, and its control is important in reducing the risk of developing cardiovascular disease in patients with T2D.23 Specifically, in a study of over 20,000 patients from 10 prospective European cohort trials, the DECODE study group found an increased risk of death from cardiovascular disease and all causes in patients with high 2-h glucose concentrations (impaired glucose tolerance [140–198 mg/dL] and diabetes [≥198 mg/dL]) following an oral glucose tolerance test.24 Another report of 3370 patients from the Framingham Offspring Study found that the 2-h response to an oral glucose challenge was associated with cardiovascular events, independent of other cardiovascular risks and measures of hyperglycemia.24 In addition, a meta-analysis of 13 studies in patients with T2D found that long-term glycemic fluctuations, measured by the coefficient of variation or standard deviation (SD) of HbA1c, were associated with renal and cardiovascular disease, macrovascular events, and mortality, but not with microvascular events.25

However, contrary to the above-cited studies, some suggest that there is no evidence for a relationship between glycemic fluctuations and diabetic complications. A reanalysis of the Diabetes Control and Complications Trial, which initially suggested the risk for microvascular complications may be related to postprandial hyperglycemia, found these results to be an artifact of the statistical models used.26 Instead, the risk for complications was primarily determined by HbA1c. Although more research is necessary, glycemic fluctuations have been recommended as an emerging target in the management of diabetes27 and may potentially impact the risk of complications in patients who have otherwise achieved HbA1c targets.

Glycemic Fluctuations and Oxidative Stress

Acute hyperglycemic fluctuations can potentially contribute to diabetic complications through increased glycation and activation of oxidative stress.28 The induction of reactive oxygen species (ROS) generation and lipid peroxidation by glucose and saturated fat was first demonstrated by Mohanty et al.29,30 This work was developed further by the demonstration that a high-fat, high-calorie mixed meal induced oxidative and inflammatory stress for several hours.31 Thus, even in healthy subjects, the intake of macronutrients can induce periodic oxidative and inflammatory stress, as reflected in increased ROS generation, lipid peroxidation, and the induction of endotoxemia and proinflammatory cytokines.32 These effects can potentially be amplified in patients
with diabetes and chronic hyperglycemia, who have increased basal levels of oxidative and inflammatory stress.\textsuperscript{33}

Monnier et al. investigated the relationship between chronic hyperglycemia, acute glucose fluctuations, and oxidative stress in patients with T2D and found a significant relationship between acute glucose fluctuations and oxidative stress.\textsuperscript{34} This case-controlled study used CGM to measure fasting plasma glucose and HbA\textsubscript{1c}; mean amplitude of glucose excursions (MAGE) and postprandial area under the curve (AUC) were used to measure acute glucose fluctuations and postprandial hyperglycemia, respectively. Oxidative stress was measured using 24-h urinary excretion of free 8-iso-prostaglandin F\textsubscript{2\alpha}. Oxidative stress is mediated by increased ROS production at the mitochondrial level and membrane NADPH oxidase activity. These mechanisms link persistent hyperglycemia and acute glucose fluctuations with glucose-induced activation of protein kinase C isoforms,\textsuperscript{35} increased production of advanced glycation end products, and greater glucose flow through the aldehyde reductase pathway—all of which lead to damage of vascular endothelial cells in diabetes.\textsuperscript{36} Acute glucose load and hyperglycemia also induce ROS generation through NADPH oxidase-mediated mechanisms.\textsuperscript{37,38}

**Glucose Monitoring Increases Awareness of Glycemic Fluctuations**

Glucose monitoring can increase patients’ awareness of the relationship between their lifestyle and daily glycemic fluctuations. For instance, in a study of 201 insulin-treated patients with T2D, HbA\textsubscript{1c} was significantly decreased by 0.30% and 0.36% after 4 and 8 weeks, respectively, of self-monitored blood glucose (SMBG).\textsuperscript{38} Compliance was a strong predictor of the improvement at 8 weeks, after adjusting for baseline HbA\textsubscript{1c}. However, compliance ranged from 3% to 100%, with an average of 78.4%.

CGM offers a potential alternative to SMBG for patients amenable to the CGM system. Recent advances in CGM technology have made the use of CGM systems easier than past versions, and integration of small CGM systems into closed-loop insulin delivery systems is underway. Future aims for CGM technology include further miniaturization of CGM systems. This easy-access, real-time monitoring system of glucose levels, which also alerts patients to high or low glucose values, can reduce glycemic fluctuations by improving patients’ awareness of hyperglycemia and hypoglycemia.\textsuperscript{39}

**CGM to Evaluate Effectiveness of Treatments on Glycemic Fluctuations**

CGM provides a tool for evaluating the impact of different treatments on glycemic control. In a 4-week study in 33 patients with T2D who had inadequate glycemic control after ≥2 months on metformin, similar HbA\textsubscript{1c} reductions were seen among patients assigned to treatment with the dipeptidyl peptidase-4 (DPP-4) inhibitor sitagliptin (from a mean of 7.0% to 6.6%; \(P < 0.001\)) and those assigned to the sulfonylurea glimepiride (from a mean of 7.3% to 6.9%; \(P < 0.001\)).\textsuperscript{40} However, a significant reduction in MAGE was only seen in the sitagliptin group (from 88 to 67 mg/dL; \(P < 0.001\)). The greater impact of sitagliptin on reducing glucose fluctuations may be related to the glucose-dependent mechanism of action for glucose-lowering effects, whereas sulfonylureas, which are not sensitive to blood glucose concentrations, increase the risk of blood glucose troughs that may contribute to fluctuations (Table 1).

Sitagliptin and vildaglaptin add-on therapy for 8 weeks resulted in similar, significant (\(P < 0.05\)) improvements from baseline in 24-h glucose fluctuations (as indicated by reductions of ~15 mg/dL in MAGE) in patients with T2D inadequately controlled by metformin.\textsuperscript{41} However, only vildaglaptin recipients had significant (\(P < 0.05\)) improvements from baseline in parameters reflecting daily glycemic control (mean 24-h blood glucose level, time spent in the ideal glycemic range [70–140 mg/dL] and above hyperglycemic thresholds [140 and 180 mg/dL], along with corresponding AUC values).

The \(\alpha\)-glucosidase inhibitors are a class of glucose-lowering drugs taken at meals that specifically reduce postprandial glucose by inhibiting carbohydrate digestion in the small intestine and delaying its absorption (Table 1).\textsuperscript{42} A 24-week randomized controlled trial compared the addition of the \(\alpha\)-glucosidase inhibitor acarbose or the sulfonylurea glibenclamide to metformin in patients with poorly controlled T2D (\(n = 55\)).\textsuperscript{43} While both treatments significantly reduced HbA\textsubscript{1c} (\(P < 0.001\)), the addition of acarbose significantly decreased MAGE (from a mean of 101 to 72 mg/dL; \(P < 0.001\)), whereas glibenclamide had no significant effect on MAGE (mean of 112 mg/dL at baseline and 114 mg/dL at study end). Likewise, among 45 patients with T2D who had high CGM MAGE values (≥261 mg/dL) on premixed insulin analogue therapy, the addition of acarbose significantly decreased MAGE (from 77 to 47 mg/dL; \(P < 0.01\)) and the mean of daily differences (from 23 to 20 mg/dL; \(P < 0.05\)), a measure of interday glycemic fluctuations.\textsuperscript{44} The incidence of hypoglycemic events in these patients decreased from 24% to 2% (\(P < 0.01\)).

In addition, CGM was utilized in a 4-week, randomized, placebo-controlled study to evaluate the sodium–glucose cotransporter 2 inhibitor dapagliflozin in 100 adult patients with T2D also being treated with either metformin alone or insulin and up to two oral glucose-lowering drugs.\textsuperscript{45} At day 28, dapagliflozin 10 mg/day significantly reduced MAGE compared with placebo (−10.0 vs. +5.3 mg/dL; \(P < 0.01\)). Furthermore, dapagliflozin significantly reduced the proportion of time with blood glucose >180 mg/dL (−12.6% vs. +3.5%; \(P < 0.001\)) and increased the proportion of time spent with blood glucose levels in the target range of 70–180 mg/dL (+12.2% vs. −2.8%; \(P < 0.001\)). A small increase in the proportion of time with blood glucose <70 mg/dL (+0.3% vs. −0.6%) was driven by the dapagliflozin plus insulin group.

CGM can also uncover hypoglycemia, even when no symptoms are present.\textsuperscript{46} In a study of 29 patients with T2D well controlled on metformin ± sulfonylureas, 18 patients (62%) experienced 65 silent, symptom-free hypoglycemic episodes (interstitial glucose ≤70 mg/dL). CGM can alert patients to these silent changes in blood glucose levels, providing patients with the opportunity to make the necessary adjustments to potentially avoid these episodes.

For those patients who require additional therapy beyond oral agents, insulin and other injectables such as glucagon-like peptide-1 receptor agonists (GLP-1RAs) are used. Small pilot and subpopulation studies first provided evidence for glucose fluctuation-lowering effects of GLP-1RAs,\textsuperscript{47,48} whereas larger studies designed to assess the effects of GLP-1RAs on glucose fluctuations have been carried out more recently.
Table 1. Mechanism of Action and Primary Effects of Different Glucose-Lowering Drugs Used in Patients Uncontrolled with Metformin

| Drug class                                      | Examples of drugs approved in the US | Administration and timing of dose | Primary MOA for glucose lowering | Primary effects for glucose lowering |
|------------------------------------------------|--------------------------------------|----------------------------------|---------------------------------|-------------------------------------|
| Not sensitive to plasma glucose concentrations |                                      |                                  |                                 |                                     |
| Sulfonylurea                                    | Glibenclamide, glimepiride, glipizide, gliclazide | Oral; Once or twice daily        | Increase insulin secretion       | Target FPG, with resultant lowering of PPG |
| Thiazolidinedione                               | Pioglitazone, rosiglitazone          | Once daily                       | Enhance insulin sensitivity; reduce free fatty acid release | Target FPG, with resultant lowering of PPG |
| Basal insulin and intermediate insulin          | Insulin glargine, insulin detemir, insulin degludec, insulin NPH | Oral; Once or twice daily        | Exogenous insulin replacement   | Target FPG, with resultant lowering of PPG |
| Rapid-acting insulin                            | Insulin aspart, insulin lispro, insulin glulisine | SC injection; Up to three times daily | Exogenous insulin replacement | Target PPG, with resultant lowering of overall glucose |
| Glucose-dependent MOA                          |                                      |                                  |                                 |                                     |
| α-Glucosidase inhibitors                       | Acarbose, miglitol                   | Oral; Up to three times daily at mealtime | Inhibit carbohydrate digestion in the small intestine and delay its absorption | Target PPG, with resultant lowering of overall glucose |
| SGLT-2 inhibitors                               | Dapagliflozin, canagliflozin, empagliflozin | Oral; Once daily                 | Increase urinary glucose secretion | Target FPG and PPG |
| DPP-4 inhibitors                                | Saxagliptin, sitagliptin, linagliptin | Oral; Once or twice daily        | Increase circulating levels of incretins (e.g., GLP-1 and GIP) to increase insulin secretion and suppress glucagon secretion | Target PPG and lower overall glucose |
| Short-acting GLP-1RAs                           | Exenatide BID, lixisenatide QD       | SC injection; Once or twice daily at mealtime | Stimulate GLP-1R to increase insulin secretion and suppress glucagon secretion | Target PPG and FPGa |
| Longer-acting GLP-1RAs                          | Liraglutide QD, exenatide QW, dulaglutide QW, albglutide QW | SC injection; Once daily (liraglutide) or once weekly | Stimulate GLP-1R to increase insulin secretion and suppress glucagon secretion | Target PPG and FPGa |

*aShort-acting GLP-1RAs result in greater reductions in PPG versus longer-acting GLP-1RAs, whereas longer-acting GLP-1RAs result in greater reductions in FPG versus short-acting GLP-1RAs. BID, twice daily; DPP-4, dipeptidyl peptidase-4; FPG, fasting plasma glucose; GIP, glucose-dependent insulinotropic polypeptide; GLP-1, glucagon-like peptide-1; GLP-1RA, glucagon-like peptide-1 receptor agonist; MOA, mechanism of action; NPH, neutral protamine Hagedorn; PPG, postprandial glucose; QD, once daily; QW, once weekly; SC, subcutaneous; SGLT-2, sodium–glucose cotransporter 2.
FLAT-SUGAR was a 26-week study comparing the effects on glycemic fluctuations (measured using CGM) of the GLP-1RA exenatide twice daily (BID) or rapid-acting insulin as prandial add-on therapy to insulin glargine plus metformin in patients with T2D \( (n=102) \). While exenatide BID and prandial insulin both target postprandial glucose concentrations, the insulinogenic effects of exenatide BID are glucose dependent, in contrast to insulin (Table 1). After 26 weeks of treatment, HbA1c levels were comparable in the exenatide BID and rapid-acting insulin groups (7.1% and 7.2%, respectively). Mean changes in the coefficients of variation of glucose from baseline to 26 weeks (as determined by CGM) were significantly different between the exenatide BID and rapid-acting insulin groups (-2.4 vs. +0.4; \( P=0.047 \)). Furthermore, MAGE was improved in the exenatide BID group compared with the rapid-acting insulin group \( (P=0.049) \).

In a recent, randomized, double-blind study, patients with T2D treated with exenatide once weekly (QW) and metformin \( (n=60) \) had improvements in daily glycemic control and reductions in glucose fluctuations compared with patients treated with placebo and metformin \( (n=56) \). Exenatide QW was associated with reductions from baseline in 24-h glucose profiles at weeks 4 and 10, whereas changes with placebo were less pronounced (Fig. 1). Compared with placebo, exenatide QW significantly reduced 24-h mean glucose (-30.8 vs. -3.0 mg/dL), MAGE (-15.2 vs. +2.9 mg/dL), and the SD of mean glucose (-6.3 vs. +0.7 mg/dL) at week 10 (all \( P<0.001 \)). In addition, exenatide QW significantly reduced the proportion of time spent with patient blood glucose concentrations in the hyperglycemic range (>180 mg/dL; \( P<0.001 \)) and significantly increased the proportion of time spent in the target glycemic range (70–180 mg/dL; \( P<0.001 \)) compared with placebo, without increasing time spent in the hypoglycemic range (<70 mg/dL).

In an analysis of three randomized controlled trials comparing insulin glargine (two studies) or insulin detemir (one study) with exenatide QW, treatment with exenatide QW had a greater reduction in HbA1c than either insulin (\(-1.5\% \) vs. \(-1.3\% \ [P<0.02] \) and \(-1.1\% \) vs. \(-0.7\% \ [P<0.001] \) for the two studies of insulin glargine; \(-1.3\% \) vs. \(-0.9\% \ [P<0.001] \) for the study of insulin detemir). Furthermore, treatment with exenatide QW resulted in a significantly greater reduction in MAGE, as compared with insulin glargine (\(-12 \) vs. \(-4 \) mg/dL \( [P=0.002] \); \(-27 \) vs. \(-1 \) mg/dL \( [P<0.001] \)), but not compared with insulin detemir (\(-16 \) vs. \(-8 \) mg/dL \( [P=0.080] \)). Similar results were found for blood glucose range and blood glucose variability.

The GLP-1RA liraglutide has also been found to reduce glycemic fluctuations in patients with T2D. For instance, IDegLira—a fixed-ratio combination of insulin degludec and liraglutide—has been shown to reduce prandial glucose. After 26 weeks of treatment \( (n=260) \), a meal test found a significantly (\( P=0.0023 \)) greater reduction in mean \( \pm \) SD postprandial glucose increment with IDegLira (\(-15.7 \pm 29.7 \) mg/dL) as compared with insulin degludec (\(-3.1 \pm 35.7 \) mg/dL); treatment with liraglutide had a comparable decrease (\(-14.1 \pm 29.2 \) mg/dL) to that of IDegLira. When the change from baseline in postprandial interstitial glucose increment was assessed with CGM, IDegLira had a significantly greater effect than insulin degludec (estimated treatment difference: \(-6.1 \) mg/dL; \( P=0.0047 \)) and a comparable effect to liraglutide (estimated treatment difference: \(-1.8 \) mg/dL; \( P=0.4122 \)).

A separate study of 63 patients with T2D compared treatment with liraglutide and neutral protamine Hagedorn (NPH) insulin. After 12 weeks, treatment with liraglutide was associated with a significantly \( (P<0.05) \) greater decrease in MAGE (\(-77.4 \) mg/dL with liraglutide; \(-46.8 \) mg/dL with NPH insulin) and in large amplitude of glycemic excursions (\(-104.4 \) mg/dL with liraglutide; \(-54.0 \) mg/dL with NPH insulin), as assessed by CGM.

![FIG. 1. Change from baseline in mean 24-h glucose profiles at weeks 4 and 10 (exenatide QW + MET, \( n=60 \); placebo + MET, \( n=56 \)). Light blue line = exenatide QW + MET at week 4; navy blue line = exenatide QW + MET at week 10; light pink line = placebo + MET at week 4; magenta line = placebo + MET at week 10. MET, metformin; QW, once weekly. Reproduced with permission from Frías et al.](image-url)
37 patients with T2D treated with either liraglutide plus intensive high-dose insulin therapy or standard insulin uptitration for 6 months, CGM again showed reduced glycemic fluctuations, as measured by the within-day SD of mean daily glucose (22.5% reduction with liraglutide plus insulin [P < 0.0001]; no significant change with standard insulin therapy alone). An uncontrolled study of 20 Japanese patients with T2D also found liraglutide to significantly reduce MAGE (from 94 to 44 mg/dL; P < 0.001 vs. baseline), as well as the proportion of time spent in hyperglycemia, without any increased time spent in hypoglycemia.

Although exenatide BID and liraglutide are both GLP-1RAs, they can have differing effects on glycemic fluctuations. An open-label, randomized study of 35 patients with T2D inadequately controlled on sulfonylureas compared the effects of 7 days of treatment with either exenatide BID or liraglutide (once daily) on glycemic fluctuations, as measured by 24-h CGM. No differences were found for 24-h mean glucose, SD of blood glucose, MAGE, or early fasting glucose. However, for breakfast and dinner, 3-h postmeal glucose was significantly lower with exenatide BID than with liraglutide (P < 0.01), whereas both the mean and SD of blood glucose between 12:00 am and 6:00 am was significantly higher (P < 0.05) with exenatide BID compared with liraglutide. This difference may reflect the duration of action and the pharmacodynamics of the two drugs (Table 1).

Finally, as part of the AWARD-4 trial, the effects of dulaglutide versus insulin glargine on glycemic fluctuations were assessed. A subgroup of 144 patients taking either once-weekly dulaglutide (1.5 or 0.75 mg) or insulin glargine, both added to prandial insulin lispro, were evaluated using CGM. After 26 weeks, dulaglutide 1.5 mg significantly reduced glycemic fluctuations compared with insulin glargine, as measured by within-patient SD (−12.6 vs. −7.2 mg/dL; P < 0.05) and mean of daily differences (−7.2 vs. +1.8 mg/dL; P < 0.05). At 52 weeks, within-patient SD and mean of daily differences were similar across all groups. At both time points, MAGE was similar across groups.

Data from the above studies demonstrate that multiple glucose-lowering agents that act in a glucose-dependent manner, such as DPP-4 inhibitors and GLP-1RAs, or that inhibit carbohydrate digestion, such as α-glucosidase inhibitors, may reduce the occurrence of hyperglycemia without increasing the risk of hypoglycemia. Together, this limits short-term glycemic fluctuations. In addition, DPP-4 inhibitors and GLP-1RAs have ROS-suppressive and anti-inflammatory effects. As glycemic fluctuations have been linked to oxidative stress and downstream vascular dysfunction and damage, this may provide further benefit for patients in addition to reducing glycemic fluctuations. Insulin treatments were generally not as effective against glycemic fluctuations, although they were effective in reducing the longer-term measure of glycemic control, HbA1c. Additional studies that utilize CGM would be helpful to further establish the effects of these different agents on glycemic fluctuations.

CGM in Clinical Practice: Case Studies

The two cases below demonstrate that CGM is a useful tool for diagnosing unrecognized hypoglycemia and hyperglycemia in situations in which it is impractical to check fingerstick concentrations.

**Case 1**

A 41-year-old male with T2D regularly exercised by running 3–4 miles daily. The patient had good glycemic control with an HbA1c of 6.8% on treatment with metformin and continuous subcutaneous insulin infusion (i.e., an insulin pump). After his daily run, he reported feeling weakness and a lack of energy. He denied having neurogenic symptoms of hypoglycemia, such as sweating, shakiness, palpitations, anxiety, or increased sensation of hunger. He also denied neuroglycopenic symptoms, such as dizziness, inappropriate behavior, difficulty with concentration, confusion, or blurred vision.

To investigate the possibility of unrecognized hypoglycemia, the patient was fitted with a CGM system and asked to check fingerstick blood glucose at the time of his symptoms. CGM revealed hypoglycemia during the patient’s run as well as marked postmeal hyperglycemia. Despite a reduction in basal insulin, he continued to experience hypoglycemia during and after running. However, during weight training (i.e., resistance training exercises), CGM indicated that he maintained normoglycemia. As a result, the patient switched from running to resistance training for his routine exercise. To prevent postmeal hyperglycemia, he increased his dose of prandial bolus insulin, thereby increasing his insulin to carbohydrate ratio factor. Following these changes, the patient’s HbA1c fell to 6.5% without any hypoglycemia or postmeal hyperglycemia. Fluctuations in his blood glucose concentrations were also reduced, leaving the patient feeling more safe and secure while exercising and administering additional bolus insulin doses with meals. Thus, CGM significantly changed the QoL of this patient while enabling him to achieve his HbA1c goal safely, without any hypoglycemia-related complications.

**Case 2**

A 52-year-old female with T2D for 8 years reported waking up on several occasions during the night following nightmares that left her sweaty, nervous, and shaky. The patient reported normal fasting blood glucose concentrations. She was on treatment with metformin (1000 mg BID), insulin glargine (30 U at bed time), and insulin lispro (three times daily before meals). This regimen was initiated in an attempt to intensify her treatment for uncontrolled T2D with an HbA1c of 10.2%. She was ~1 year postmenopausal and occasionally experienced hot flashes during the day. With estradiol therapy, her daytime hot flashes had resolved, but she continued to have nocturnal sweaty episodes. She was referred to our diabetes and endocrinology clinic for further evaluation of her episodes of night sweats and nervousness, often coupled with nightmares. Her HbA1c was 7.8% and a random blood glucose concentration was 180 mg/dL. She had mild neuropathy and microalbuminuria.

In an attempt to elucidate the relationship between the nighttime episodes and glycemic levels, the patient was fitted with a CGM system. CGM revealed a consistent fall in blood glucose concentrations after midnight, lasting from 20 min to 1 h, with the trend beginning at 9:00 pm. One of these hypoglycemic episodes coincided with a nightmare-associated episode of night sweats. A diagnosis of nocturnal hypoglycemia was made, and her doses of insulin lispro with dinner and insulin glargine at bedtime were reduced. The episodes of night sweats and nightmares disappeared. Later, the patient substituted a GLP-1RA for insulin lispro treatment and her
HbA1c fell to 7.0% without clinical or CGM-recorded hypoglycemia.

Through CGM, the patients described above were able to check for hypoglycemia and hyperglycemia in situations, such as running or sleeping, that would otherwise be impractical for fingerstick checks. The insight provided by CGM led to an improved understanding of daily blood glucose fluctuations, allowing for targeted lifestyle and therapeutic changes and an improvement in their QoL.

Conclusions

Glycemic fluctuations, through either chronic sustained hyperglycemia and/or acute glucose variations, may be associated with diabetic complications. HbA1c remains the gold standard for assessing glycemic control and the efficacy of glucose-lowering therapies. However, HbA1c has limitations, such as the inability to reflect the peaks and troughs of blood glucose levels that occur daily in many patients with T2D. Glycemic fluctuations are emerging as an important potential parameter to consider when managing blood glucose levels, with potential effects on the risk of diabetic complications such as cardiovascular disease and QoL.

For certain patients, CGM may be a useful tool for assessing the impact of treatment and lifestyle on daily changes in blood glucose levels. Recent studies have utilized CGM to demonstrate reductions in glycemic fluctuations with glucose-lowering therapies, such as sodium–glucose cotransporter 2 inhibitors, DPP-4 inhibitors, and GLP-1RAs, and future CGM studies would provide further insight into the effects of specific treatments on glycemic fluctuations.

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Address correspondence to:
Paresh Dandona, MD, PhD
MOB, Erie County Medical Center
462 Grider Street
Buffalo, NY 14215
E-mail: pdandona@kaleidahealth.org