INTRODUCTION

The rate-limiting step of intensive diabetes management for patients with diabetes is treatment-induced hypoglycemia. Thus, hypoglycemia remains a significant barrier in optimizing glycemic control and reducing long-term diabetes-related complications including cardiovascular (CV) death, stroke, and all-cause mortality. CV events are the leading cause of death among patients with diabetes. The risk of CV death is twice that of patients without diabetes. Severe hypoglycemia (resulting in cognitive impairment) can occur across a broad spectrum of A1C levels. Patients achieving near-normal glycemia (<6%) and those who were poorly controlled (≥9%) appeared to be at the highest risk for severe hypoglycemia. Hypoglycemia triggers a vascular and inflammatory cascade which can result in vascular constriction, tachycardia, thrombosis, and fatal arrhythmias. Therefore, patients who have existing CV disease should have their glycemic targets relaxed to mitigate the risk of hypoglycemia.

Recurrent hypoglycemia lowers or even eliminates the blood glucose concentration threshold at which patients develop a sympathetic response likely to prompt them to take evasive action in time to reverse an impending event. Elderly patients may lose their balance without warning, falling to the ground. The subsequent confusion after such a fall is likely to be attributed to “the aging process” rather than to deficient glucose counterregulation.

Clinicians should remind patients with diabetes who use insulin to monitor their blood glucose levels before operating a motor vehicle. The single most significant factor associated with driving collisions for drivers with diabetes appears to be a recent history of severe hypoglycemia, regardless of the type of diabetes, or the treatment used. A single
hypoglycemia event can deplete counterregulatory hormone levels which favor recognition and reversal of low plasma glucose, resulting in a lack of awareness of a hypoglycemic event.

Nocturnal hypoglycemia may result in patients arriving late to work or missing entire days at the office. Patients who experience hypoglycemia are likely to use extra test strips for fear of having a recurrence of low blood glucose levels. Calls to doctors for guidance on glucose management are increased after an episode of hypoglycemia, and patients often inappropriately self-titrate their medications to avoid future events.\[6\]

Hypoglycemia increases the risk of CV and all-cause mortality in patients with diabetes.\[1\] Severe hypoglycemia is associated with a macrovascular events hazard ratio (HR) of 2.88 and a microvascular event HR of 1.81. The mortality HR for a hypoglycemic event in patients with type 2 diabetes is 2.69.\[1\] Hypoglycemia during hospital admissions is associated with increased lengths of stay and with increased 1-year mortality and inpatient mortality rates (2.96%) for patients who had at least one hypoglycemic episode during the hospitalization versus 0.82% for patients who had none.\[5\]


Hypoglycemia can increase vascular inflammation, QT prolongation, intravascular coagulation, life threatening arrhythmias, and delayed clot thrombolysis. Therefore, patients with known CV disease must minimize their risk of hypoglycemia.\[7\]

**DEFINITION OF HYPOGLYCEMIA**

The American Diabetes Association and the European Association for the Study of Diabetes (ADA/EASD) updated its standards of care and hypoglycemia definitions in 2017.\[8\] The implications of different blood glucose levels vary from individual to individual, but <70 mg/dL is considered an alert for hypoglycemia and allows patients time to take corrective action. A glucose level of <54 mg/dL is considered clinically significant and unequivocally hypoglycemic. Severe hypoglycemia has no assigned biochemical value and is defined simply and starkly as a glucose level low enough to cause cognitive impairment such that the assistance of another person is required to administer carbohydrates or glucagon to achieve a recovery. Table 1 lists the American Diabetes Association definition of hypoglycemia levels.


**CASE PRESENTATION - LISA**

Lisa is 68 years of age and has had poorly controlled type 2 diabetes. She is a former kindergarten teacher who neither smokes or drinks. Lisa had an inferior wall infarction 1 year ago after which she received three stents. She is adherent with her prescribed diabetes treatment regimen yet is unable to achieve her prescribed A1C target of 7.5%. She is currently taking metformin 1 g with breakfast and dinner and liraglutide 1.8 mg/d. Her blood pressure and lipids are at the American Diabetes Association recommended targets. Her A1C is 8.7%. A download of her continuous glucose monitor (CGM) is shown in Figure 1a.

The CGM begins recording interstitial glucose readings 12 h after being inserted into the upper arm of the patient. A handheld “reader” is waved across the sensor site providing the patient with a real-time glucose reading. The glucose level is stored in the reader and can be accessed in the form of charts, graphs, and data points. One important data set is the time within a prescribed targeted range (usually 80–180 mg/dL) and the percentage of the time the patient is hypoglycemic (<70 mg/dL). The dark solid line provides a graphic representation of the median interstitial glucose readings throughout the day as well as during the life of the sensor (10 days). The blue zone which surrounds the median represents the 25th–75th% in which 50% of the interstitial glucose levels are recorded. A separate 10–90% curve represents the glucose levels of 90% of the glucose readings.

These patient’s glucose levels tend to drop overnight increasing Lisa’s risk of experiencing unrecognized nocturnal hypoglycemia if she is intensively treated with basal insulin. Increased glycemic variability following dinner is also associated with hypoglycemia risk. The patient’s
sensor demonstrates a persistent elevation in glucose levels throughout the day which correlates well with her A1C of 8.7%. The clinician should consider intensifying her treatment with the effective agents likely to allow the patient to achieve her targeted glycemic goals while minimizing the risk of hypoglycemia and weight gain.

**LISA CLINICAL FOLLOWUP**

Lisa’s physician prescribes insulin glargine to be used in conjunction with metformin 1 g twice daily and liraglutide. The starting dose of glargine is 10 units at 9 pm daily. She is advised to increase the dose by 1 unit each night until her fasting blood glucose level is <110 mg/dL. The patient continues the use of her CGM and titrates her insulin glargine as directed. Although her interstitial glucose levels improve, she is noted to have developed nocturnal hypoglycemia. [Figure 1b].

After titrating her insulin glargine dose from 10 to 32 units daily, our patient’s glucose levels are noted to have dropped overnight resulting in unrecognized nocturnal hypoglycemia. In addition, her blood glucose levels tend to decline after dinner between the hours of 6 pm and 12 am.

Lisa expresses relief that her glucose levels have improved but are concerned about the frequency of treatment-induced nocturnal hypoglycemia.

A1C, which is a reflection of one’s average blood glucose levels over a 90-day interval, is a poor predictor of hypoglycemia risk. Figure 2 shows the continuous glucose sensor readings of two insulin-requiring patients with type 2 diabetes, and both of whom have A1C levels of 7.8%. Patient A (gray line) demonstrates erratic glucose readings ranging from 40 to 325 mg/dL (dysglycemia), whereas patient B (brown line) has less glycemic variability and no evidence of hypoglycemia. Intensifying the insulin regimen on patient A will result in severe hypoglycemia as well as weight gain and the induction of “o Unger xidative stress” which could promote the development of long-term diabetes-related complications. Dysglycemia is shown in Figure 3. This can be a frustrating problem for patients as well as clinicians.

CGM readings for a single patient were taken over 4 separate days demonstrating significant glycemic variability occurring between the hours of 6–8 am. High glycemic variability reduces one’s quality of life and can promote mood disorders.

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**Figure 1:** Results from a 10 day wear of a freestyle libre continuous glucose monitor. The download from this patient demonstrates the frequency of significant nocturnal hypoglycemia (red coloring) in a patient who tends to have higher blood glucose readings as the day progresses. Increasing the dose of basal insulin for this patient would likely result in severe nocturnal hypoglycemia.
Patients who experience dysglycemia become frustrated with their inability to efficiently regulate blood glucose levels. The clinician may be asked, “I can’t understand why on Tuesday my blood glucose level was 46 mg/dL, yet the following day I was at 278 mg/dL?” All too often these patients are labeled as being “non-compliant” when, in fact, adherence to their prescribed pharmacologic regimen results in dysglycemia. Efficient pharmacotherapy for patients with diabetes must address both the effects of prolonged exposure to hyperglycemia as well as acute daily excursions of glucose levels which could increase one’s risk of developing treatment-emergent hypoglycemia [Figure 4].

ADDRESSING GLYCEMIC VARIABILITY USING NOVEL INSULIN FORMULATIONS

Ideal basal insulins should be simple to initiate and titrate, resulting in minimal glycemic variability, as well as provide prolonged duration of action while reducing one’s risk of hypoglycemia and weight gain. In addition, insulins should not increase one’s risk of CV disease, especially in patients who have already experienced a stroke or myocardial infarction. Table 2 provides the coefficients of variability of available basal insulins. The lower the variability, the less likelihood of developing treatment-emergent hypoglycemia.

The coefficient of variability is a predictor of hypoglycemia risk. The lower the coefficient, the less one is likely to experience hypoglycemia from that agent. Figure 5a and b demonstrate the relationship between the coefficient of variability and hypoglycemia risk. Figure 5c depicts ambulatory glucose monitoring (sensor data) from a patient with minimal glycemic variability.

A patient is using an insulin with a high coefficient of variability. Glucose levels demonstrate “dysglycemia” (A). Increasing the basal insulin further is likely to increase the risk of hypoglycemia, which will reduce adherence to the prescribed treatment regimen (B). The use of basal insulin formulations with low glycemic variability will allow patients to achieve their fasting blood glucose targets more efficiently, with less fear of hypoglycemia.[11]

Ambulatory blood glucose monitoring using the freestyle libre sensor demonstrating minimal glycemic variability with an essential flat 24 h glucose profile (black line).

ADVANCES IN INSULIN FORMULATIONS

Advances in basal insulin formulations have provided clinicians and patients with options that provide favorable pharmacokinetic (insulin absorption) and pharmacodynamic (glucose lowering) properties. Newer insulins have flatter, peakless action profiles that demonstrate less variability and a longer duration of action, allowing for flexible dosing. The risk of nocturnal and diurnal hypoglycemia is subsequently reduced. Insulin does not appear to increase CV risk.[12] Patients may also safely combine a glucagon-like peptide-1 RA as either a separate injection or as a component of a fixed-ratio drug. The use of fixed-drug combinations may improve adherence and allow patients to achieve their metabolic targets.[13]

The insulin analog glargine U100 and detemir have longer half-lives than neutral protamine Hagedorn (NPH) insulin and, at similar A1C levels, reduce the frequency of overall and nocturnal hypoglycemia by 42–53%. This reduction in hypoglycemia is most likely secondary to reduced day-to-day variability observed with these insulin analogs versus NPH.[14] Recently approved glargine U-300 and insulin degludec provide patients with a protracted duration of action while reducing the risk of confirmed and nocturnal hypoglycemia in patients with type 2 diabetes. Although insulin is recognized as the most effective blood glucose-lowering therapy, patients who experience hypoglycemia while on insulin are less likely to adhere to their prescribed insulin regimen, thus reducing the
Minimizing one’s risk of experiencing treatment-emergent hypoglycemia is imperative within the primary care practice model.\textsuperscript{[15]}

### INSULIN GLARGINE U-300

Glargine U-300 is a long-acting insulin containing 300 units/mL of insulin glargine. U300 is soluble within the acidic pH of the injection medium (pen injector) but less soluble in the subcutaneous tissue once injected. Once exposed to the physiologic environment of the subcutis, the acidic PH of the pen’s insulin solution forms a microprecipitate, resulting in a slow and prolonged release of insulin from the injection depot. The glargine U300 molecule is identical to the amino acid sequence of U100 insulin. However, on injection, the more concentrated U300 forms a more compact subcutaneous depot with a smaller surface area leading to the protraction of insulin absorption\textsuperscript{[17]} [Figure 6a and b].

As a concentrated insulin, glargine U-300 contains 3 times as much insulin per mL as glargine U-100, allowing for a lower volume of injected insulin. Glargine U-300 was detectable at 32 h post-injection with 0.4 units/kg dosing compared with 28 h with glargine U100 dosing.\textsuperscript{[18]} At 0.4 u/kg, U-300 has 14% less variability than U-100, allowing clinicians to titrate the insulin to target lower fasting glucose levels without risking hypoglycemia.\textsuperscript{[19]} In a study comparing the two glargine insulin formulations, the risk of hypoglycemia was found to be 31% lower with glargine U300 compared with glargine U-100 owing to the superior pharmacokinetic profile of U300.\textsuperscript{[20]} [Figure 7] Thus, insulin-requiring patients at risk for hypoglycemia could have 31% lower risk of hypoglycemia if initiated on glargine U300 instead of glargine U100.

### Table 2: Basal insulin coefficient of variability

<table>
<thead>
<tr>
<th>Insulin</th>
<th>Within-subject variability*</th>
</tr>
</thead>
<tbody>
<tr>
<td>NPH</td>
<td>68</td>
</tr>
<tr>
<td>Glargine U-100</td>
<td>48</td>
</tr>
<tr>
<td>Detemir</td>
<td>27</td>
</tr>
<tr>
<td>Concentrated glargine U-300</td>
<td>34.8</td>
</tr>
<tr>
<td>Degludec</td>
<td>20</td>
</tr>
</tbody>
</table>

\*Percentage within subject variability based on glucose infusion rates and area under the curve. Patients receive 4 single subcutaneous doses of 0.4 U/kg under euglycemic glucose clamp conditions on 4 study days. NPH: Neutral protamine Hagedom

![Figure 4](image1.png)

**Figure 4:** Hypoglycemia risk reduces one’s ability to achieve optimal and targeted glycemic control

![Figure 5](image2.png)

**Figure 5:** (a) Patient’s continuous glucose sensor demonstrates significant diurnal glycemic variability trending toward hyperglycemia. (b) Increasing the dose of basal insulin in this patient would likely result in continuous hypoglycemic events throughout the day. (c) Demonstrates a patient with minimal glycemic variability. Increasing the basal insulin in this patient is less likely to result in significant hypoglycemia.
Note that, both insulin formulations reduce A1C equally at 6 months. However, the glargine U300 insulin reduces the risk of nocturnal hypoglycemia by 31% and also results in less weight gain than in patients taking glargine U100.\(^{[21]}\)

**INSULIN DEGLUDEC**

Within the insulin pen, degludec is formulated as “insulin diheximers.” Once injected, the diheximers form multiheximer chains within the subcutaneous depot held together by zinc and phenol. As the zinc dissociates, the multiheximers form insulin monomers, which pass into the capillaries and are carried through albumin to insulin receptors at target organ sites [Figure 8]. Degludec U-200 contains as much insulin as degludec U-100 in just one-half of the injection volume. The two degludec formulations are bioequivalent, and they lower glucose levels at the same rate. Degludec appears to have the lowest coefficient of variability of all insulins, allowing ambitious dosing to targeted fasting glucose levels, with less likelihood of nocturnal and overall hypoglycemia compared with insulin glargine.\(^{[22]}\)

Due to the prolonged duration of action (42 h), degludec may be dosed at any time of the day, which may improve adherence for patients who are shift workers, travel frequently, or have difficulty remembering to dose their basal insulin.\(^{[23]}\)

Insulin degludec is maintained as diheximers within the insulin pen before injection. Once injected, the diheximers form multiheximers maintained by phenol and zinc. Over time, the phenol and zinc dissociate leaving the pharmacologically active form of insulin (monomers) to pass into the capillaries.
Unger: Management of hypoglycemia in patients with type 2 diabetes

Insulin degludec and insulin glargine have demonstrated the equivalent risk of treatment-emergent CV events in patients with type 2 diabetes.\[^{24}\] A randomized, double-blind, two period crossover, treat-to-target trial was conducted in patients with type 2 diabetes with basal insulin with or without oral antidiabetic drugs over 65 weeks.\[^{15}\] Patients were initiated on either insulin glargine or insulin degludec for 32 weeks followed by a blinded crossover to the other insulin. Each 32-week interval consisted of a 32-week titration period which allowed patients to achieve their glycemic targets (fasting glucose 71–90 mg/dL), followed by a 16-week maintenance phase comparing the difference in hypoglycemia events (confirmed blood glucose < 56 mg/dL or an episode requiring 3rd party assistance) when glycemic control and doses were stable. The primary endpoint was the rate of overall symptomatic hypoglycemic events or confirmed hypoglycemia during the 16-week maintenance phase. The secondary endpoints were the rate of nocturnal and symptomatic hypoglycemia occurring between the hours of 12:01 am and 5:59 am. The proportions of patients with hypoglycemic episodes were 22.5% for insulin degludec versus 31.6% for insulin glargine U100. The proportion of patients with nocturnal hypoglycemia events was 9.7% versus 14.7% favoring insulin degludec. 2.4% of patients using glargine U100 experienced severe hypoglycemia versus 1.6% of patients using insulin degludec.

Lisa’s glycemic control has improved significantly. 83% of her interstitial glucose readings are within the target of 80–180 mg/dL. However, 11% of her glucose levels are <70 mg/dL. These events appear to be occurring overnight as well as at 4 pm.

**TARGETING HYPOGLYCEMIA PREVENTION IN HIGH-RISK PATIENTS**

Improving patient education and empowering the patient to take some control over their disease are often very valuable tactics to improve both treatment outcomes and treatment adherence and therefore reduce the frequency of hypoglycemia. Patients prescribed drugs which work through a glucose-independent manner (such as insulin and sulfonylureas) should be instructed to monitor for and report any treatment-emergent hypoglycemia events. Adjustments in therapy will be required to mitigate risk.

Patients who are unable to perceive the symptoms of hypoglycemia should be prescribed continuous glucose sensors which alarm when a rapid decline in interstitial glucose levels is noted. Sensor will allow patients to act to reverse hypoglycemia before they develop cognitive impairment. Spouses and family members should be interviewed as well to determine if a patient has developed abnormal behaviors suggestive of hypoglycemia unawareness such as falls, confusion, disorientation, aggressive behavior, and nightmares [Table 3].

Family members should be instructed on the proper means of reversing severe hypoglycemia in patients with diabetes as shown in Figure 3.
Therapeutic advances are continually being made, and insulins that ever more closely match the physiological profile of human insulin are currently the Food And Drug Administration approved and should be considered as first-line agents in patients at high risk for experiencing treatment-emergent hypoglycemia. Patients may also consider the use of CGM and ambulatory glucose monitoring to mitigate their risk of hypoglycemia.

CONCLUSION

Iatrogenic hypoglycemia represents a much more substantial barrier to the effective control of blood glucose concentrations in patients with diabetes than is currently appreciated. Greater awareness and detection of all hypoglycemic events by careful monitoring, adherence to guidelines, and use of optimal treatment combinations are needed to prevent the serious medical and economic consequences associated with this adverse effect of antihyperglycemic medications. There is a need for improved patient and provider education on optimal detection and understanding of hypoglycemia and the benefits of accurate blood glucose measurement, which together can help to reduce the risk and fear of hypoglycemia, hypoglycemia unawareness, and hypoglycemic events.

ACKNOWLEDGEMENT

The author would like to thank the Primary Care Network for the assistance and support in preparing this manuscript.

REFERENCES