Are Glucose Readings Sufficient to Adjust Insulin Dosage?

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Abstract

Aims/hypothesis: Insulin therapy is effective predominantly when dosage is frequently adjusted. However, a controversy surrounds the pertinent clinical parameters required to make effective and safe frequent dosage adjustments. We hypothesize that glucose readings are sufficient to adjust insulin dosage provided that dosage is adjusted every 1–4 weeks.

Methods: To test the hypothesis, we generated several algorithms implemented in software to process glucose readings and recommend insulin dosage adjustments. A post hoc analysis was made on 630 log sheets (2,520 insulin dosage adjustments) from 26 older adults with suboptimally controlled type 2 diabetes. The subjects were followed for a year and treated with intensive insulin therapy that was titrated every 1–4 weeks by a trained study team. More than 88% of subjects attained the treatment goal (hemoglobin A1c <7%) without excessive hypoglycemia. Glucose readings from each log sheet were used as an input to the software, and its recommendations for insulin dosage adjustments were compared to the original ones made by the study team. While the study team could have been exposed to multiple clinical parameters, the software relied solely on glucose readings.

Results: The software recommendations for dosage adjustments were clinically equivalent to the original study team’s recommendations in more than 95% of the cases, unrelated to patients’ insulin sensitivity. The remaining 4.4% (n = 111) were thoroughly examined, yet we did not find any recommendations suggested by the software to be unsafe or unreasonable.

Conclusions/Interpretation: Glucose readings are sufficient to effectively adjust insulin dosage provided that adjustments are made every 1–4 weeks. Therefore, dedicated software can help adjusting insulin dosage between clinic visits.

Introduction

Insulin is the only antidiabetes agent lacking a therapeutic window. In other words, most insulin-treated patients can achieve satisfactory glycemic control provided that appropriate formulations and adequate dosage are prescribed. Yet, almost two-thirds of insulin-treated diabetes patients in the United States fail to reach the therapy goal (hemoglobin A1c [A1C] <7%),1-2 although compliance to antidiabetes medications is generally considered adequate in about three-fourths of the cases.3,4 Most insulin users have type 2 diabetes; some use partial insulin therapy (premixed, biphasic, or long-acting insulin), and some use intensive insulin therapy (basal/bolus). Only clinical trials that reinforce a policy of insulin dosage adjustments every 1–4 weeks (cumulatively >15,000 patient-years) for either partial5-10 or intensive insulin therapy11-13 achieved treatment goals (A1C <7%) among 40% and 80% of the subjects, respectively. Although this finding is hard to separate from the Hawthorne effect, when trials have ended and frequent insulin dosage adjustment strategy was replaced with infrequent conservative care predominantly during clinic appointments, patients’ A1C was unfavorable merely a year after study termination.13,14

The nature of this paradox likely ensues from patients’ metabolic behavior. Because of variability within each patient (Fig. 1), insulin requirements significantly fluctuate over time and thus cannot be met by infrequent dosage adjustments during clinic visits (typically once every 3–6 months).

Most clinicians agree that insulin dosage should be titrated/adjusted far more frequently than during routine clinic visits. However, the clinical parameters as well as the frequency required to make effective frequent dosage adjustments are still controversial. Insulin dosage should not be confused with insulin doses that for some regimens may depend on preprandial glucose reading (sliding scales or correction factors).
and carbohydrate intake. A common convention that diet and exercise habits need to be explored before insulin dosage could be adjusted. Some believe that dosage should be adjusted every 3 days or even daily. Others believe that carbohydrate counting is crucial for the success of the regimen. However, it has been established that for patients with type 2 diabetes treated with intensive insulin therapy, carbohydrate counting is not superior to a sliding scale provided that dosage is frequently adjusted.

Many of the aforementioned studies also published the instructions given to the study team for frequent insulin dosage titrations. These instructions advised on frequent insulin dosage adjustments solely based on glucose readings. Yet, because these instructions encourage considerable amount of flexibility and clinical judgment, it is unclear how faithfully they were followed.

We hypothesize that glucose readings are sufficient to effectively adjust insulin dosage provided that dosage is adjusted every 1–4 weeks. To test the hypothesis, we generated several algorithms and implemented them in software to recommend insulin dosage adjustments based only on glucose readings. Glucose readings from log sheets of subjects treated with intensive insulin therapy that was titrated every 1–4 weeks by a trained study team were used as an input to the software. While the study team could have been exposed to multiple clinical parameters, the software relied solely on glucose readings. The software and the study team’s recommendations for insulin dosage adjustments were systematically compared using a similarity metric.

Subjects and Methods

Original study design

The original study was designed as a two-center, prospective, randomized, controlled, clinical trial. Inclusion and exclusion criteria are available elsewhere. In brief, subjects were ≥60 years of age, had a clinical diagnosis of type 2 diabetes for at least 1 year, were taking at least one injection of insulin per day (with or without oral antidiabetes medications), and had an A1C ≥7.0%. One hundred seven subjects were randomized, and 98 completed follow-up. Patients were randomized to either multiple daily insulin injections or continue subcutaneous insulin injections. Of the 54 patients randomized to multiple daily insulin injections, data from 28 subjects who were followed at the Michigan site were available for re-analysis. Data from two of the 28 patients were incomplete and insufficient for re-analysis. The remaining 26 subjects were cumulatively followed for 22 patient-years.

The multiple daily insulin injection regimen consisted of once-daily insulin glargine (Lantus®, Aventis, Bridgewater, NJ) and preprandial insulin lispro (Humalog®, Eli Lilly, Indianapolis, IN). All subjects were instructed to monitor their blood glucose levels before meals and at bedtime. At least once a week, subjects were instructed to monitor nocturnal blood glucose levels. Initial basal insulin dosage was calculated as 50% of the total daily insulin dose and administered as glargine before bedtime. The remaining 50% was administered as preprandial lispro boluses and formulated as a sliding scale. Accordingly, dosage was given as a combination of four elements: (1) basal insulin dose, (2) sliding scale for breakfast, (3) sliding scale for lunch, and (4) sliding scale for dinner. Dosage adjustments were made once every 1–4 weeks by the study team, which included three endocrinologists and two nurses with extensive experience in intensive insulin therapy. Subjects submitted log sheets (for an example see Supplementary Appendix; Supplementary Data are available online at www.liebertonline.com/dia) including dosage given by the study team, time-tagged glucose reading, and insulin doses. The study team provided a new recommended dosage on each log sheet, which was later communicated to the subject via the study coordinator.

Efficacy was assessed by A1C measured at the baseline and at the 1-, 2-, 4-, 6-, 8-, 10-, and 12-month visits. Safety was evaluated by the rate of severe hypoglycemia, defined as capillary glucose ≤2.8 mmol/L (≤50 mg/dL) associated with neuroglycopenic symptoms that require assistance by another person. Twenty-three of the 26 subjects successfully and safely reached A1C <7% (more than 88%), and mean A1C improved from 7.8% to 6.3%.

Determination of adherence

Subjects’ adherence served as a marker for quality control to corroborate the weight of the study team’s instructions. For each patient, lispro meal doses over a period of two randomly chosen weeks were individually compared to the sliding scale of the fast-acting insulin boluses. The deviation of each administered dose from the expected dose due to the sliding scale was expressed as a percentage.

Software design

To prove or refute our hypothesis it was mandatory to refrain from human interference while regenerating dosage recommendations and comparing them to the study team’s recommendations. Therefore, we created software with algorithms that processed time-tagged glucose readings and adjusted insulin dosage while ignoring any additional clinical parameters. The software was developed by E.B. and I.H. prior to acquiring the clinical data. W.H.H. and his study team, from whom the data were acquired, were not in-
involved in the development of the software. The algorithms embedded in the software were based on state-of-the-art guidelines for insulin management and on the following four principles:

1. Time-tagged glucose readings were the only input used to adjust the current dosage and create recommendations for the next insulin dosage.
2. Insulin dosage was increased for glucose level above target and decreased for glucose level below target.
3. The intensity of adjustments decreases as glucose readings get closer to target to prevent unstable oscillations of dosage.
4. The ability to detect “outliers.” The software utilizes higher-order statistics to detect outliers and treats them separately from the remainder of the data.

**Similarity between the software and study team’s recommendations**

The database consisted of 630 log sheets (2,520 insulin dosage adjustments) containing time-tagged glucose readings, previous dosage prescribed by the study team, and a new set of adjusted dosage (see Supplementary Appendix). Each log sheet’s glucose readings were processed by the software. The software generated new recommended dosage for long-acting insulin and three mealtime fast-acting insulin sliding scales. Because of inherent dependencies, only the 4.4–6.7 mmol/L (80–120 mg/dL) glucose range of each sliding scale, i.e., each meal bolus dose, was used for comparison. These four components generated by the software per log sheet were compared to the original study team’s recommendations using a similarity metric.

**Similarity metric**

Differences between the study team recommendations and the software recommendations were classified into six categories:

1. **Identical**: the software and study team made the same dosage recommendation.
2. **Within 10%**: the dosage recommended by the software was within 10% of the dosage recommended by the study team. In addition, the two dosage modifications were in the same direction (i.e., the software did not recommend increasing the dosage, whereas the study team recommended decreasing the dosage, or vice versa).
3. **Within 10–20%**: the dosage recommended by the software was within 10–20% of the dosage recommended by the study team. In addition, the two dosage modifications were in the same direction.
4. **Different; 10–20%**: The software recommended increasing the dosage by 10–20%, whereas the study team recommended decreasing the dosage.
5. **Different; more than 20%**: The software recommended increasing the dosage by more than 20%, while the study team recommended decreasing the dosage.
6. **Other**: all other cases (not complying with categories 1–5).

Of the above, categories 4 and 5 were defined as “Different” and represented potentially hazardous disagreement between the software and the study team.

We considered the software recommendations to be “clinically equivalent” to the study team’s recommendations if they were classified in categories 1, 2, or 3. We assumed that a 20% difference is a reasonable value that can be seen in routine

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**Table 1. Subjects’ Characteristics Compared to the Original Randomized Study Population**

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Multiple daily insulin subjects for re-analysis</th>
<th>Original randomized population</th>
</tr>
</thead>
<tbody>
<tr>
<td>Subjects (n)</td>
<td>26</td>
<td>107</td>
</tr>
<tr>
<td>Age (years)</td>
<td>67.4 ± 4.9</td>
<td>66.2 ± 5.4</td>
</tr>
<tr>
<td>Sex (male)</td>
<td>13 (50)</td>
<td>62 (58)</td>
</tr>
<tr>
<td>Race</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Caucasian</td>
<td>26 (100)</td>
<td>92 (86)</td>
</tr>
<tr>
<td>Black</td>
<td>0</td>
<td>6 (5.6)</td>
</tr>
<tr>
<td>Hispanic</td>
<td>0</td>
<td>6 (5.6)</td>
</tr>
<tr>
<td>Other</td>
<td>0</td>
<td>3 (2.8)</td>
</tr>
<tr>
<td>Duration of diabetes (years)</td>
<td>18.3 ± 9.9</td>
<td>15.9 ± 9.0</td>
</tr>
<tr>
<td>History of diabetes complications</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Retinopathy</td>
<td>9 (34.6)</td>
<td>39 (36)</td>
</tr>
<tr>
<td>Nephropathy</td>
<td>5 (19)</td>
<td>16 (15)</td>
</tr>
<tr>
<td>Neuropathy</td>
<td>14 (53.8)</td>
<td>69 (64)</td>
</tr>
<tr>
<td>History of cardiovascular complications</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypertension</td>
<td>19 (73)</td>
<td>79 (74)</td>
</tr>
<tr>
<td>Dyslipidemia</td>
<td>19 (73)</td>
<td>72 (67)</td>
</tr>
<tr>
<td>Cigarette smoking (current)</td>
<td>1 (4)</td>
<td>5 (5)</td>
</tr>
<tr>
<td>Ischemic heart disease/heart failure</td>
<td>10 (38)</td>
<td>38 (35)</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>29.9 ± 6.1</td>
<td>32.2 ± 5.7</td>
</tr>
<tr>
<td>A1C (%)</td>
<td>7.8 ± 0.7</td>
<td>8.2 ± 1.0</td>
</tr>
</tbody>
</table>

Data are mean ± SD values or n (%).
No statistically significant differences were noted.
A1C, hemoglobin A1c; BMI, body mass index.
clinical settings. Moreover, it has been demonstrated that identical insulin injections can result in different plasma insulin profiles (can exceed 20%) because of the complex process of insulin absorption and dispersion. For illustration, we believe that it is not unreasonable in clinical settings to have one care-provider increasing lunchtime lispro from 20 units to 21 units and another from 20 units to 24 units, using data from the same patient. The difference between these two examples is almost 20%.

Statistical analysis

Subjects’ characteristics were compared to the original study population by two-tailed Student’s t test for parametric variables and \( \chi^2 \) test for nonparametric variables. Results are presented as mean ± SD values. A \( P \) value < 0.05 was defined as statistically significant.

The process of insulin dosage recommendations, by either the study team or the software, is not a random process. Therefore, we neither performed nor presented statistical comparisons between the software and study team’s recommendations. Instead, the similarity metric was used to measure the distance between the two sets.

Results

The re-analyzed subjects represent the original study population

Data from the 26 subjects treated with multiple daily insulin injections were available for post hoc analysis. Table 1 compares the basic characteristics of this cohort to the original study population that was eligible for
randomization (n = 107). No significant differences were identified.

Mean ± SD A1C of the re-analyzed subjects was 7.8 ± 0.7% before enrollment (median, 7.7%), and following 12 months of intensive insulin therapy it was 6.3 ± 0.8% (median, 6.1%) (P < 0.0001). The mean improvement (±SD) of A1C was 1.5 ± 0.9% (median, 1.6%) and was similar to that of the original study population. Mean total daily insulin dosage of the re-analyzed subjects was 66.2 units (0.7 units/kg) at the beginning of the study and 104.5 units (1.0 units/kg) at the study’s end. Among the re-analyzed subjects 88.5% (23 of 26) achieved A1C <7% at the expense of six episodes of severe hypoglycemia (0.27 events per person-year). Ninety-six percent of the re-analyzed patients had at least one episode of
minor hypoglycemia. The frequency of these events was not statistically different from that observed in the original study population (in the original study 90% of patients incurred minor hypoglycemia, and the frequency of severe hypoglycemia was 0.16 events per person-year) and was comparable to that reported in subjects treated with intensive insulin therapy (most available data derive from patients with type 1 diabetes). Body mass index of the 26 subjects did not significantly change during the study (initial value, 29.9 ± 6.1 kg/m²; final value, 30.9 ± 6.5 kg/m²).

**Subjects were adherent to the study team’s instructions**

To determine dosing accuracy, 1,239 meal boluses were compared to the corresponding sliding scales. Each deviation from a prescribed dose was expressed as a percentage. For instance, if for a pre-lunch glucose level of 10 mmol/L (180 mg/dL) a subject was instructed to administer 24 units of lispro and instead injected 26, this dose was registered as 108.3%. In 77.2% of the instances the reported dose was identical to the prescribed one. In 13.4% of the instances the reported dose deviated from the prescribed dosage by more than 10% (data not shown). We assumed that in these cases, subjects deviated from the prescribed dose because of factors such as decreased food intake or increased physical activity.

Although incorporating only glucose readings, the software recommendations lie in close proximity to those recommended by the study team

A database of n = 2,520 recommendations was available for comparison. As shown in Figure 2A, the software made “Identical” (similarity metric category 1) dosage recommendations (long-acting and fast-acting insulin) in 42.3% (n = 1,067) of the cases, 38% (n = 959) were “Within 10%” (category 2), and 15.2% (n = 383) were “Within 10–20%” (category 3).

When analyzed separately, both long-acting (glargine) and fast-acting (lispro) gave identical or similar recommendations in 95–97% of the cases, respectively (Fig. 2B). Eight cases (0.3%) were considered “Different; 10–20%” (category 4). Of these eight, seven cases were recorded in a single subject treated with 51 units of insulin/day. These seven cases represented 7.95% of the 88 dosage recommendations for this particular subject. In all of these cases, the study team recommended decreasing the dosage by a single unit, whereas the software recommended increasing the dosage by a single unit. The eighth case was recorded in a patient treated with 55 units of insulin/day. In this particular event, the study team recommended decreasing the patient’s lunch lispro dosage from 11 to 10 insulin units, whereas the software recommended increasing the same dosage from 11 to 13 insulin units. Pre-dinner glucose measurements for the discussed week were 8, 10, 20.3, 4.4, and 18.7 mmol/L (144, 180, 365, 79, and 337 mg/dL). Thus, it is not clear whether the software recommendation was erroneous. No cases of “Different; >20%” (category 5) were noted.

Of the 103 cases categorized as “Other” (category 6), in 40 non-clustered episodes (1.6% of the entire data set), the study team recommended increasing a dosage, whereas the software recommended decreasing it. The rest of the “Other” cases included adjustments differing by more than 20% mostly because of quantization (e.g., if a bolus dosage component of 4 units was increased by the study team to 5 units, whereas it was kept unchanged by the software, the difference was 25%).

In summary, 95.6% of the software recommendations (using only glucose measurements to adjust the prior dosage) yielded clinically equivalent insulin dosage adjustments when compared with those of the study team.

As illustrated by the examples in Figure 2C, in all subjects, insulin dosage adjustments were essential not only for induction of intensive insulin therapy but also for maintaining optimal A1C during the entire study.

**Glucose reading is the only parameter required to frequently adjust insulin dosage, invariant to insulin sensitivity**

To determine whether the similarity between recommendations of the software and the study team was not related to subjects who required either high or low insulin dosage, we divided the 2,520 data points into three groups (820, 776, and 924, corresponding to eight, nine, and nine subjects) according to the subjects’ insulin sensitivity. Insulin sensitivity was calculated by total daily insulin dose (in units) divided by weight (in kg) at the end of the trial. The subjects were divided into the following groups: (A) 0.3–0.6 (21–59 units/day), (B) 0.6–1.0 (52–120 units/day), and (C) 1.0–1.9 (113–232 units/day).

In all three groups, more than 93% of the software dosage recommendations were clinically equivalent to the study team’s dosage recommendations, i.e., no more than 20% apart (Fig. 2D). Similarity between dosage recommendations tended to be lower for patients treated with fewer insulin units per day. This was ascribed to clinically minor differences of 1–2 units between the software and the study team’s recommendations. In conclusion, these results suggest that frequent insulin dosage adjustments can be made based only on glucose readings independent of patients’ insulin sensitivity.

**Other clinical guidelines for insulin dosage adjustments based on glucose readings may reasonably correlate with ones made by trained care-providers**

Our software was designed to imitate the decision making process of a care-provider when adjusting insulin dosage every 1–4 weeks. Yet, different care-providers may have different approaches. Although not initially designed to operate independent of the care-provider input, we implemented in software the guidelines for frequent insulin dosage adjustments published by Bergenstal et al. The same data used before from the 630 log sheets (2,520 insulin dosage adjustments) were fed to “Bergenstal’s guidelines,” and the recommendations were compared to the original recommendations of the study team. In 15.8% of the cases the guidelines gave identical recommendations to the ones made by the study team; in 32.5%, “within 10%”; in 39.2%, “within 10–20%”; no “different 10–20%” or “different >20%” were noted; and 12.5% were categorized as “other” (data not shown).
Discussion

This report explores the vital clinical parameter required to facilitate frequent insulin dosage adjustments. Unfortunately, the growing mismatch between patients’ needs and caregivers’ availability disallows frequent outpatient insulin dosage adjustments, thus making insulin therapy less effective. In the preceding analysis, we compared modifications of insulin dosage (long-acting and mealtime fast-acting) made by a dedicated software to ones made by an experienced study team. The only input the software used to adjust insulin dosage was glucose readings, whereas the study team communicated with the patients and could have been exposed to other clinical parameters. We showed that the software recommendations were clinically equivalent to the ones made by the study team. Concomitantly, the original recommendations of the study team were demonstrated to be safe and effective. Although frequently adjusted insulin therapy has been supported by multiple studies, the optimal frequency of adjustments is yet to be determined. In this study, subjects were contacted every 1–4 weeks for dosage adjustments and confer superior control without excess of hypoglycemia.

The frequency of severe hypoglycemia both in the reanalyzed cohort and in the original study population was low as been routinely demonstrated elsewhere and considered to be beneficial despite its risk. In more than 80% of the cases, dosage adjustment recommendations made by the software were, at most, within 10% of those made by experienced endocrinologists and nurses. In 15% of the cases dosage modifications made by the software were within 10–20% of the study team’s recommendations. Similar correlation was demonstrated separately for long-acting and fast-acting insulin and among subjects with different insulin sensitivity.

In our metric, the ability of the software to make safe insulin dosage adjustments was assessed by six distance categories (see Similarity metric). Categories 4 and 5 included cases in which the software recommended an increase in dosage whereas the study team recommended decreasing it. These categories were considered as potentially hazardous because an overdose of insulin may result in hypoglycemia. Yet, in the eight cases (0.3%) classified as “Different; 10–20%” (category 4) it was unclear whether following the software recommendations was unreasonable. As outlined in Figure 2A, no events of “Different; >20%” (category 5) were identified.

Category 6, i.e., “Other,” was found in 4.1% of the cases. This category included cases where the study team recommended increasing a dosage component whereas the software recommended decreasing the same component. In these cases it could be inferred that underdosing may lead to subsequent hyperglycemia, followed by overdosing and finally hypoglycemia. Our static comparison could not have unequivocally excluded it. Yet, such episodes occurred rarely (40 non-clustered cases, or 1.6% of all episodes of dosage adjustments) and were therefore unlikely to have lead to hazardous situations if the software were to independently make frequent dosage adjustments.

Although the software is not intended to operate independent of a care-provider’s input, we computed clinical guidelines for insulin dosage adjustments suggested by a different institution. We found reasonable correlation implying that clinical guidelines for frequently adjusted insulin therapy based on glucose readings do not require additional parameters to make the therapy effective.

Our study is limited by the fact that subjects were closely followed and more titrated by an expert study team. Unfortunately, in reality, patients are infrequently seen in the clinic, and their insulin dosage is infrequently adjusted. Still, only clinical studies that incorporate close follow-up allow the type of analysis presented. Although the A1C goal set in our study was questioned by the ACCORD study, the ACCORD patients were treated with multiple diabetes medications and not only with insulin. Recent analysis of the ACCORD data did not find insulin therapy to be an independent risk factor for adverse outcome.

The software discussed here emulates the decision-making process of an expert care-provider, coaching the patient to frequently adjust insulin dosage. Although we fully acknowledge the impact of diet and exercise on diabetes management, the presented data suggest that glucose readings alone are sufficient to enable effective and safe insulin dosage optimization provided that dosage adjustments are made frequently. A prospective clinical study where a healthcare provider uses only glucose data to adjust subjects’ insulin therapy dosage is planned.

In today’s reality, the overwhelming workload in clinics and the deficiency in care-providers trained in insulin titration make frequent insulin dosage adjustments unrealistic. Consequently, only 35% of insulin-treated patients achieve A1C <7% because insulin dosage is seldom adjusted and regimens become too rigid to compensate for the dynamic needs of patients. Dependent on further clinical data, we postulate that such software has the capacity to enable patients to safely realize the full benefits of their insulin therapy by adjusting insulin dosage between appointments in order to achieve optimal glycemic control. This can alleviate the care-providers from their impossible task to frequently optimize insulin therapy, shorten clinic waiting time, and assign more time for patients’ education and management of co-morbidities.

Author Disclosure Statement

Both I.H. and E.B. are co-founders of Hygieia Inc. W.H.H. declares no competing financial interests.

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