Degludec is superior to glargine in terms of daily glycemic variability in people with type 1 diabetes mellitus

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Abstract. To investigate the differences in glycemic variability between the long-acting insulins glargine and degludec using continuous glucose monitoring, we conducted an open-label, multicenter, prospective, observational study that enrolled 21 participants with type 1 diabetes mellitus currently receiving basal-bolus insulin therapy with glargine. To avoid the potential influence of diet and exercise on glycemic control, all participants were housed and monitored within the hospital for the duration of the study. Once glycemic control was achieved with glargine, glycemic variability was evaluated using continuous glucose monitoring for 3 days. Glargine was then replaced by degludec and glycemic variability again assessed via continuous glucose monitoring. The primary outcome measure of mean amplitude of glycemic excursions was significantly reduced with degludec ($p = 0.028$), as was area under the curve for daily blood glucose level <70 mg/dL ($p = 0.046$). The required insulin dose was reduced up to 25% in the degludec group, although 24-h mean glucose concentrations were not different between groups. In conclusion, once or twice daily glargine was successfully replaced by a daily injection of degludec. When replacing glargine with degludec, a lower dose should be utilized to avoid hypoglycemia. Degludec is an effective and promising long-acting insulin that reduced hypoglycemia and daily blood glucose variability in participants with type 1 diabetes.

Key words: Continuous glucose monitoring, Insulin degludec, Type 1 diabetes mellitus

INSULIN GLARGINE (Gla) is currently the most commonly used long-acting (basal) insulin analog. Recently, insulin degludec (Deg) has been approved as a new long-acting insulin and is now available for daily clinical practice. Deg has longer and flatter effects compared to other conventional long-acting insulins [1]. The durability of circulating Gla is less than 24 h [2, 3], thus a subset of people with insulin-deficient type 1 diabetes mellitus must receive twice-daily injections of Gla. Additionally, it was reported that glycemic control can be improved in people with type 1 diabetes mellitus by administering Gla twice daily as compared to a single daily dose [3]. In contrast, a single injection of Deg had an evenly distributed glucose lowering effect over a 24 h period with a duration of action greater than 26 h [4]. The reduced number of insulin injections is thought to be preferred by people requiring insulin and is considered an improvement in quality of life [5].

Although the phase 3 trials of Deg showed good performance as a new basal insulin in people with type 1 diabetes [6-8], few studies have shown the effect of Deg on the stability of glucose levels using continuous glucose monitoring (CGM) [9, 10]. CGM systems allow visualization of the daily variations in blood glucose levels, facilitating the detection of postprandial hyperglycemia and asymptomatic hypoglycemia. Therefore, these systems can provide a more accurate means to evaluate the efficacy of glucose lowering therapies within individual participants [11]. Blood glucose variability is an important contributing factor to the severity of coronary artery disease independent of HbA1c [12, 13], as well as an independent predictor of mortality [14]. Mean amplitude of glycemic excursions (MAGE) is a marker of daily blood glucose variability relating to higher postprandial blood glucose or hypoglycemia,
and was recently reported to correlate closely with cognition level and oxidative stress *in vivo* [15, 16]. It is worth evaluating the difference in daily blood glucose variability between Deg and Gla using CGM systems.

In a phase 3 clinical trial, Deg decreased the frequency of nocturnal hypoglycemia compared to Gla [6-8]. However, hypoglycemia was defined as when the participants experienced hypoglycemic symptoms and blood glucose levels were less than 56 mg/dL or when severe hypoglycemia was present that required assistance and therefore, asymptomatic hypoglycemia was not included. Previous CGM data indicate that asymptomatic hypoglycemia is unexpectedly common among people with type 1 diabetes mellitus [17]. Furthermore, in the phase 3 trials of Deg, basal insulin dose was titrated to target a fasting blood glucose of 70-89 mg/dL. People with type 1 diabetes mellitus are more susceptible to hypoglycemia because of the severe fluctuation in their blood glucose. Therefore the lower blood glucose target in these trials could increase the frequency of hypoglycemia as compared to that observed in standard practice. The goal of the present study was thus to investigate the superiority of Deg in terms of glycemic variability compared with Gla under conditions more representative of clinical practice (less than 130 mg/dL at premeal and less than 180 mg/dL at 2-hours (h) postprandial) and to assess incidence of asymptomatic hypoglycemia via CGM. In order to minimize the potential influence of factors such as diet and exercise on glycemic control, and compare the intrinsic efficacy of the two basal insulins, all CGM data were collected from people with type 1 diabetes mellitus that were hospitalized and following optimization of glycemic control.

**Materials and Methods**

**Participants**

This open-label, multicenter, comparative, observational, prospective study enrolled people with type 1 diabetes who had been admitted to Hokkaido University Hospital or KKR Sapporo Hospital. All participants received an explanation of the study procedures and potential risks associated with participation and gave their written informed consent prior to entry. The inclusion criteria were hospitalized people with type 1 diabetes over the age of 20 receiving basal-bolus insulin therapy including Gla as basal insulin. People who had unstable retinopathy of diabetes, a history of anaphylaxis to Deg, were pregnant women, had a persistent elevation of their serum transaminase, or had renal dysfunction were excluded. This study was approved by the Institutional Review Board of Hokkaido University Hospital and was performed in accordance with the Declaration of Helsinki.

**Design of the study**

Enrolled participants were provided a fixed caloric diet that was calculated to adequately meet the nutritional requirements of each participant. The participants were asked to avoid excessive exercise and snacking. Minimal glucose consumption was allowed only when symptomatic hypoglycemia occurred. This study was performed using the following protocol (Fig. 1).

Glycemic control was targeted at less than 130 mg/dL for average premeal blood glucose levels and less than 180 mg/dL for average 2-h postprandial glucose levels, according to the Japan Diabetes Society guidelines. The insulin doses administered were adjusted by expert endocrinologists in order to ensure optimum glycemic control. Administration of other antidiabetic medications during the study period was not allowed. Once glycemic control met the target criteria described above, glycemic variability was evaluated by CGM for 3 days (Day 1-3). Following this 3 day monitoring period, the basal insulin was switched from Gla to Deg. The initial dose of Deg that was administered was 20% less than the dose of Gla the participants had been receiving. The next 3 days (Day 4-7) were used to optimize the dose of bolus insulin and Deg to achieve the same target of glycemic control, and CGM data were then collected for 3 days (Day 7-9). Fasting blood samples were taken before and after the study.

![Fig. 1 Study design. The enrolled hospitalized participants with type 1 diabetes were treated and got the target range of glucose using CGM as described in the text. After 3 days monitoring (day 1-3) using Gla, Gla was replaced to Deg at 20% lower dose than Gla. Insulin dose was adjusted during day 4-6, then the last 3 days monitoring (day 7-9) was done using Deg. CGM, continuous glucose monitoring; Gla, glargine; Deg, degludec.]
After the study, the participants were asked to select which basal insulin they preferred and allowed to continue use of the insulin they had selected.

**Glycemic variability**

“Nighttime” was defined as 0000 h to 0600 h and “early morning,” as 0300 h to 0600 h. In this study, “hypoglycemia” was defined as the area under the curve (AUC) <70 mg/dL; and “hyperglycemia,” as the area under the curve (AUC) >180 mg/dL.

The primary outcome was glycemic variability (MAGE) measured by CGM over a 24-h period for 3 consecutive days (study days 1-3 for Gla and 7-9 for Deg). Secondary endpoints included the SD values during 24-h glucose levels, hypoglycemia over a 24-h period, hypoglycemia during nighttime, hypoglycemia during daytime, hyperglycemia during a 24-h period, mean of daily difference (MODD) for a 24-h period, MODD during early morning, as well as changes in the dose of insulin administered. MODD has been used as an index of day-to-day glucose variability [20].

**Statistical analysis**

Results are expressed as mean ± SD. A paired t-test was employed to compare treatment differences between Gla and Deg. A p-value <0.05 was considered statistically significant. Data were analyzed using Ekuseru-Toukei 2012 (Social Survey Research Information, Tokyo, Japan).

### Results

**Participant characteristics**

A total of 21 hospitalized people with type 1 diabetes were enrolled in the study (5 men, 16 women). Participant baseline data were as follows: age, 55 ± 11 years; duration of disease, 13.9 ± 8.4 years; glycosylated hemoglobin (HbA1c), 67 ± 4 mmol/mol (8.3 ± 2.5%); body mass index, 21.3 ± 3.4 kg/m²; total daily insulin dose, 0.67 ± 0.24 U/kg (36.2 ± 12.8 U); total daily basal insulin dose, 0.24 ± 0.08 U/kg (13.2 ± 5.6 U); the ratio of basal insulin to total daily insulin, 37.6 ± 10.7%. Thirteen participants were receiving Gla twice daily and the others were receiving a single daily injection. The ratio of morning to evening basal insulin dose was approximately 10:9 in participants receiving Gla twice daily (Table 1).

The insulin preparation used for preprandial bolus supplementation was either insulin aspart, insulin lispro, or insulin glulisine. Insulin secretion in 17 participants was completely defective and serum CPR was undetectable, while the other 4 participants had CPR concentrations that averaged 0.48 ± 0.17 ng/mL. Six of 16 women were pre-menopausal, but none of them were menstruating during the study. When glycemic control met the predefined criteria prior to initiating the study, the average premeal blood glucose levels were 129.4 ± 35.2 mg/dL and the average 2 h postprandial glucose levels were 163.1 ± 47.1 mg/dL.

**Comparison of CGM findings (Table 2)**

Although thirteen participants were receiving Gla twice a day, a single daily injection of Deg was sufficient to control glucose levels in these participants. The primary endpoint of 24-h MAGE was significantly decreased from 144.4 ± 56.6 mg/dL to 121.7 ± 42.2 mg/dL (p = 0.028) when Gla was replaced by Deg (Fig. 2a), while no significant change was observed in 24-h mean glucose levels (153.9 ± 31.8 mg/dL and 153.6 ± 26.2 mg/dL respectively, p = 0.959). Early morning
Table 2  Comparison of 3 days’ CGM data of 21 participants between Gla and Deg

<table>
<thead>
<tr>
<th></th>
<th>Gla</th>
<th>Deg</th>
<th>p value</th>
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<tr>
<td>24-h mean glucose levels (mg/dL)</td>
<td>153.9 ± 31.8</td>
<td>153.6 ± 26.2</td>
<td>0.959</td>
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<tr>
<td>MAGE (mg/dL)</td>
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<tr>
<td>24-h</td>
<td>144.4 ± 56.6</td>
<td>121.7 ± 42.2</td>
<td>0.028</td>
</tr>
<tr>
<td>early morning</td>
<td>41.8 ± 32.6</td>
<td>28.7 ± 11.4</td>
<td>0.048</td>
</tr>
<tr>
<td>24-h SD values glucose levels (mg/dL)</td>
<td>51.6 ± 18.1</td>
<td>43.7 ± 13.7</td>
<td>0.031</td>
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<tr>
<td>AUC &lt;70 (mg/dL-h)</td>
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<tr>
<td>24-h</td>
<td>0.8 ± 1.1</td>
<td>0.2 ± 0.5</td>
<td>0.046</td>
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<tr>
<td>Nighttime</td>
<td>0.5 ± 0.9</td>
<td>0.2 ± 0.3</td>
<td>0.090</td>
</tr>
<tr>
<td>Daytime</td>
<td>0.6 ± 1.0</td>
<td>0.2 ± 0.3</td>
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<tr>
<td>24-h AUC &gt;180 (mg/dL-h)</td>
<td>17.0 ± 16.7</td>
<td>14.0 ± 13.6</td>
<td>0.321</td>
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<tr>
<td>MODD (mg/dL)</td>
<td></td>
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<tr>
<td>24-h</td>
<td>50.2 ± 16.2</td>
<td>44.1 ± 10.2</td>
<td>0.084</td>
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<tr>
<td>Early morning</td>
<td>51.2 ± 28.2</td>
<td>38.7 ± 18.0</td>
<td>0.089</td>
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<tr>
<td>Insulin dose (U/kg)</td>
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<tr>
<td>Twice a day</td>
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<tr>
<td>Basal insulin</td>
<td>0.27 ± 0.10</td>
<td>0.20 ± 0.07</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Bolus insulin</td>
<td>0.48 ± 0.28</td>
<td>0.49 ± 0.28</td>
<td>0.785</td>
</tr>
<tr>
<td>Once a day</td>
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<td></td>
<td></td>
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<tr>
<td>Basal insulin</td>
<td>0.21 ± 0.04</td>
<td>0.15 ± 0.05</td>
<td>0.007</td>
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<tr>
<td>Bolus insulin</td>
<td>0.38 ± 0.11</td>
<td>0.36 ± 0.10</td>
<td>0.609</td>
</tr>
</tbody>
</table>

MAGE, mean amplitude of glycemic excursions; MODD, mean of daily difference; 24-h, all day; early morning, 3 to 6 am; nighttime, 0 to 6 am. Values are expressed as means ± SD.

Fig. 2  Comparison of MAGE (a, b) and hypoglycemia (c, d) between Gla and Deg. MAGE (a and b) and hypoglycemia, defined as an area under curve <70 mg/dL (c and d), were calculated from 3 days’ CGM data by each long-acting insulin. Early morning was defined during 3-6 a.m. Nighttime was defined during 0-6 a.m. Gla, glargine; Deg, degludec; MAGE, mean amplitude of glycemic excursions; CGM, continuous glucose monitoring.
MAGE was also significantly decreased from 41.8 ± 32.6 mg/dL to 28.7 ± 11.4 mg/dL (p = 0.048) (Fig. 2b). Similarly, the standard deviation of daily blood glucose concentrations was significantly decreased from 51.6 ± 18.1 mg/dL to 43.7 ± 13.7 mg/dL (p = 0.031) by switching from Gla to Deg. Significant reductions in 24-h and daytime hypoglycemia were observed (Fig. 2c), while nighttime hypoglycemia only tended to decrease (p = 0.090) (Fig. 2d). The daytime hypoglycemia values were 0.6 ± 1.0 mg/dL·h and 0.2 ± 0.3 mg/dL·h (p = 0.040) for Gla and Deg, respectively.

A small, nonsignificant (p = 0.321) decrease in hyperglycemia (AUC >180 mg/dL) was observed between Gla (17.0 ± 16.7 mg/dL·h) and to Deg (14.0 ± 13.6 mg/dL·h) groups. MODD during early morning also had a trend to decrease with Deg from 51.2 ± 28.2 mg/dL to 38.7 ± 18.0 mg/dL (p = 0.089).

**Insulin dose (Table 2)**

Previous phase 3 clinical trials using Deg reported a lower dose of Deg was required to achieve similar glucose lowering as Gla [6-8]. Therefore, in this trial, the initial dose of Deg provided when switching from Gla to Deg was 20% less than the Gla dose the participant had been receiving. Hypoglycemia was minimized using this dosing strategy and subsequent doses of Deg were adjusted accordingly. For participants who received Gla twice a day, their basal insulin dose was significantly reduced (25.5%) with Deg from 0.27 ± 0.10 U/kg to 0.20 ± 0.07 U/kg (14.0 ± 6.1 U to 10.4 ± 4.6 U) (p <0.001) (Fig. 3a), and bolus insulin dose was not changed (0.48 ± 0.28 U/kg to 0.49 ± 0.28 U/kg (23.9 ± 11.1 U to 24.3 ± 12.4 U) (p = 0.597)). Similarly, in the participants who received Gla once daily, their basal insulin dose was significantly reduced (24.7%) with Deg from 0.21 ± 0.04 U/kg to 0.15 ± 0.05 U/kg (12.4 ± 5.1 U to 9.4 ± 5.5 U) (p =0.007) (Fig. 3b), and bolus insulin dose was not changed (0.38 ± 0.11 U/kg to 0.36 ± 0.10 U/kg (21.2 ± 8.7 U to 20.0 ± 8.9 U) (p = 0.609).

**Discussion**

Deg is the third long-acting insulin available for daily clinical practice since March 2013 in Japan. Although this new insulin is expected to perform well in phase 3 studies [6-8], there is limited data when it comes to the careful evaluation of its efficacy using CGM. In this study, we used CGM to compare the blood glucose stabilizing effects between long-acting insulin Gla and Deg in people with type 1 diabetes mellitus.

Deg significantly improved MAGE as the primary outcome, and standard deviation of daily blood glucose compared with Gla in people with type 1 diabetes mellitus. Moreover, significant lowering of early morning hyperglycemia (AUC >180 mg/dL) was observed between Gla (17.0 ± 16.7 mg/dL·h) and to Deg (14.0 ± 13.6 mg/dL·h) groups. MODD during early morning also had a trend to decrease with Deg from 51.2 ± 28.2 mg/dL to 38.7 ± 18.0 mg/dL (p = 0.089).
MAGE during sleep is important for people with diabetes. Lower MAGE during the early morning leads to decreased risk of the somogyi effect or dawn phenomenon resulting in lower risk of hypoglycemia as well as hyperglycemia [21]. The longer duration of action, reduced peak, and lower variability in pharmacodynamic action of Deg may explain these observed results [1, 22, 23]. Moreover, lower risk of hypoglycemia during sleep would be beneficial for people with diabetes. Deg has a relatively low intra-subject pharmacodynamic variability in the steady state compared to Gla [1], thus the day-to-day variation of glucose (MODD) is expected to improve. However, there was no significant difference in early morning MODD ($p = 0.089$) in the present study. A sub-analysis was performed that included only the 17 participants whose insulin secretion was completely defective. In this population, a significant improvement in MODD both 24-h and early morning was seen with Deg ($p = 0.022$ and 0.001, respectively). These data suggest that type 1 diabetic patients with complete insulin deficiency could receive more benefits in regulation of day-to-day variation in glucose levels by using Deg as their basal insulin.

Average daily blood glucose concentrations were similar between Gla and Deg. Given this lack of difference, replacement of Gla to Deg would not affect HbA1c values. This study was designed to assess glycemic variability after achieving targeted premeal and 2 h postprandial glucose levels with Gla or Deg. For this reason it is reasonable to assume that average blood glucose concentrations were similar despite differences in glucose variability. Consistent with the reports from Phase 3 trials, the dose of insulin Deg utilized in this trial was approximately 25% lower than that of Gla.

Recently, the same group has published two reports assessing the effects of Deg and Gla using CGM with a test diet [9, 10]. Patients with type 1 diabetes were switched from Gla or detemir to Deg and followed for 12 or 24 weeks. Consistent with our findings, patients required a lower daily dose of Deg compared to their prior insulin dose and the required dose of basal insulin decreased significantly 24 weeks after switching. However, the improvement in MAGE that we report here was not observed in this previous study [10]. The reason for the discrepancy in CGM data is unclear, but may be due to the differences in the number of participants or the inclusion of hospitalized patients in the current study whose diet and activity were closely monitored. The pharmacokinetic profile of Deg is longer and flatter than that of Gla [1]. Following a single injection, Deg remains detectable in circulation for more than 26 h [4], while Gla is no longer detectable by 24 h [2, 3]. As the duration of action of Deg is greater this results in an accumulation of Deg over time and decreases the required daily dose to maintain glucose levels. Our results confirmed the necessity to modify insulin dose when making Deg dosing calculations based on insulin Gla action.

Interestingly, phase 3 trials have reported that Deg lowers the frequency of nocturnal hypoglycemia [6-8] while our data showed a lower risk of daytime but not nocturnal hypoglycemia. There are several plausible explanations for this discrepancy. First, the targeted fasting blood glucose levels were more strictly defined (79–90 mg/dL) in the phase 3 trials [6-8] which would increase the likelihood of nocturnal hypoglycemic events. Second, only symptomatic and self-monitored blood glucose levels under 56 mg/dL were reported as hypoglycemia in the phase 3 trials, while the definition in the current study was any blood glucose value under 70 mg/dL using CGM. Using CGM Chico et al. reported that 62.5% of people experienced hypoglycemia unawareness suggesting that many asymptomatic hypoglycemia events highly likely went undetected during the phase 3 studies [17]. In addition, phase 3 trials are conducted as outpatient studies, while this inpatient study was conducted under the constant supervision of hospital staff and tightly controlled for diet and exercise. The intensive nature of this study design provided a means to accurately evaluate the intrinsic efficacy of each basal insulin while minimizing potential confounding variables.

The potential limitations of this study include the small sample size and short study duration. Additionally, this study was not conducted as a crossover study, as this would increase the required hospitalization of participants due to the additional washout period requirements thereby increasing the difficulty of enrollment. Since Gla was used ahead of Deg in the evaluation period, it is possible that this was advantageous to Deg. Subsequent studies using a more robust study design, with a larger population and conducted over a longer period of time would likely verify and extend the findings of the current study.

In conclusion, people with type 1 diabetes receiving once or twice daily injections of Gla were transitioned to a once daily injection of Deg which was associated with a dose reduction of approximately 25%.
Once an optimum dose of Deg was determined Deg reduced hypoglycemia and daily blood glucose variability (MAGE and SD) which could provide an additional benefit of decreasing cardiovascular events.

**Author Contribution**

C.Y. contributed to the data analysis and wrote the manuscript. H.M. and A.N. contributed to discussion, reviewed and edited the manuscript. H.M. designed and performed the research, and wrote the manuscript.

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**Disclosure Statement**

No potential conflicts of interest relevant to this article were reported.

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A.N. has received honoraria for lectures from Sanofi.


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