Long-term safety and efficacy of insulin degludec in the management of type 2 diabetes

Philippe Thuillier1
Zarrin Alavi2
Véronique Kerlan1

1Department of Endocrinology, Diabetology and Metabolic Diseases, 2French Institute of Health and Medical Research CIIC 1412, Medical University Hospital of Brest La Cavale Blanche, Brest, France

Abstract: Insulin degludec (IDeg) is a novel antihyperglycemic agent belonging to the therapeutic class of ultra-long duration basal insulin analogs. Its half-life and duration of action are 25 hours and 42 hours, respectively. This pharmacodynamic profile leads to a strict dosing schedule, ie, IDeg is injected at the same time each day to ensure optimal biological action and consistent glycemic control. According to the literature, IDeg provides glycemic control and nocturnal hypoglycemia reduction comparable with other long-acting analogs in type 2 diabetes mellitus. The risk of severe hypoglycemic episodes seems also to be reduced when using IDeg therapy; however, long-term follow-up is warranted for monitoring of possible but relatively infrequent adverse events. IDeg is also available in combination with aspart insulin and with liraglutide. The above preparations have been approved by the European Medicines Agency and other national health authorities. In 2012, the US Food and Drug Administration asked for a complementary study on IDeg-associated cardiovascular risk. Future prospective evaluation of large cohorts of patients with type 2 diabetes mellitus treated with IDeg, with long-term follow-up, can provide further relevant information on the safety of IDeg therapy.

Keywords: degludec insulin, hypoglycemia, HbA1c, safety, type 2 diabetes mellitus, basal insulin analog

Introduction

The European Medicines Agency and other national health authorities have recently given marketing authorization for a new ultra-long duration basal insulin analog, insulin degludec (IDeg), and its various preparations in combination with aspart insulin (IDegAsp) and with liraglutide (IDegLira) for the treatment of adults with diabetes mellitus. Currently, the most efficient glucose-lowering treatments for type 1 diabetes (T1DM) and type 2 diabetes (T2DM) in the event of oral antidiabetic drug failure are insulin glargine (IGla) and insulin detemir (IDet). Basal insulin analogs are in fact preferred to neutral protamine Hagedorn insulin because of their longer duration of action1 and flatter action profile.2 They are also distinguished by their lower intrapatient variability in hypoglycemic action.2 Although to date there has been no randomized trial on the efficacy of long-acting basal insulin analogs with regard to reduction of severe hypoglycemia, long-acting basal insulin analog therapy has been demonstrated to confer comparable glycemic control of both overall and nocturnal hypoglycemia.1 Furthermore, their long-acting effect improves quality of life in patients with T1DM, allowing glycemic control through a single injection compared with the two injections needed when using neutral protamine Hagedorn insulin. Thus, use of these basal insulin analogs should become routine in clinical practice.
Reduction of cardiovascular morbidity and mortality, along with efficacy in HbA1c reduction, are presently regarded as the main criteria for diabetes therapeutics according to the American Diabetes Association/European Association for the Study of Diabetes (ADA/EASD) consensus algorithm for the management of T2DM. Indeed, in accordance with the ADA/EASD guidelines, in addition to HbA1c efficacy, each new antiglycemic molecule must be assessed and approved for criteria such as tolerance and cardiovascular morbidity and mortality.

At present IDeg is not widely used in clinical practice, but IGla and IDet are routinely used according to the ADA/EASD efficacy and safety criteria for long-term management of T1DM and T2DM patients. After giving a brief summary of the pharmacokinetic and pharmacodynamic properties of IDeg, this review analyzes the currently available data on the long-term efficacy and safety of IDeg to help promote its optimal use in the treatment and management of T2DM patients.

**Degludec insulin: a novel slow human insulin analog**

IDeg is a novel antiglycemic agent belonging to a therapeutic class of slow insulin analogs. These molecules are characterized by delayed absorption and action. The molecular structure of IDeg is similar to that of the human insulin amino acid sequence except for a modified beta chain, ie, deletion of threonine at position 30 and addition of a 16-carbon fatty diacid to lysine at position 29 (Figure 1).^5^

**Ultra-flat action profile and low intrapatient variability**

The action of IDeg on the insulin receptor is delayed by two mechanisms. On the one hand, its development is based on the principle of a multi-soluble hexamer. In its pharmaceutical form, before injection, IDeg takes the form of a phenol and zinc formulation containing dihexamers. Once injected, the phenol is quickly eliminated, resulting in formation of multihexamer chains that are released into the subcutaneous tissue. Slow removal of zinc then allows parallel degradation of the multihexamers into monomers, and their gradual entry into the bloodstream. On the other hand, due to the 16-carbon fatty diacid attached at position 29 of the beta chain, the monomers show a high affinity for albumin, further delaying the antiglycemic action. Noteworthy, IDeg differs from IDet in the fatty acid added at position 29, ie, its longer carbon chain (16 in IDeg versus 14 in IDet) as well as its binding to lysine via a glutamic acid spacer in IDeg. This dual mechanism confers IDeg with a slower absorption rate than that of IGla and IDet. The half-life and duration of action of IDeg are 25 hours and 42 hours, IDeg.^7^

Studies comparing IDeg with other long-acting insulin analogs have confirmed that IDeg has more stable plasma concentrations. A pharmacological study by Heise et al showed the intrapatient variability to be four times lower using IDeg than IGla in terms of hypoglycemic effect in T1DM patients. Even more interesting in this study was that the intrapatient variability in the IDeg arm remained stable during 24 hours, while in the IGla arm it
increased significantly 6–8 hours after injection, reaching its maximum between 14 and 16 hours after injection. In respect to the remarkably flat pharmacokinetic profile of IDeg, several studies have focused on verifying its robustness in the various physiological and pathological contexts commonly encountered in T1DM and T2DM patients. A study of 37 T1DM patients (12 children, 13 adolescents, and 12 adults) confirmed the conservation of this ultra-flat pharmacokinetic profile in children and adolescents. Another study comparing the pharmacokinetic and pharmacodynamic profiles of IDeg in a group of “young T1DM” (18–35 years) and a group of “older DT1” (≥65 years) patients after 6 days of subcutaneous injection reported no change in pharmacokinetic and pharmacodynamic properties of IDeg in the older group of patients. Two other studies have investigated the pharmacokinetic profile of IDeg in the presence of chronic renal insufficiency and hepatic impairment and confirmed no changes in these subgroups of patients when compared with healthy volunteers.

Given the particularly stable plasma concentrations of IDeg, two main hypotheses should be be evaluated: first, whether IDeg could limit the frequency of hypoglycemic events while allowing glycemic control comparable with that obtained using other long-acting human insulin analogs (IGla and IDet); and second, whether its longer duration of action could provide patients with greater flexibility of administration and thus better quality of life. To test these hypotheses, the BEGIN® program was conducted in a large T1DM and T2DM population in the form of several Phase III trials to assess the efficacy and safety of IDeg in various indications and in many regimens (Figure 2). This review focuses on the extensive BEGIN program and other trials that have evaluated IDeg in T2DM patients. Additionally, we review the trials comparing different preparations of IDeg, such as IDegAsp and IDegLira.

Figure 2 BEGIN program and other studies focusing on insulin degludec in patients with type 2 diabetes mellitus.
Efficacy of degludec insulin: results of the BEGIN program and other studies

Studies of the efficacy of IDeg are presented in Figure 2, and are mostly substudies in the BEGIN program. These trials were multicenter, controlled, open-label, randomized, and conducted in a “treat to target” design. The latter confers doses of insulin (basal as well as prandial) systematically adjusted for all patients according to the predetermined scheme built for the study.

The protocols and study populations included in each trial are summarized in Table 1. Inclusion criteria were uncontrolled T2DM and no recent history of severe hypoglycemia. Concomitant oral antidiabetic drug therapy was allowed but differed between studies. The baseline characteristics of the T2DM patients in these therapeutic trials indicate a mean age of 60 years. The duration of T2DM ranged from 8 to 13 years and HbA1c levels were mostly between 8.0% and 9.0%. Finally, the reported body mass index was around 30 kg/m², except in Asian studies where it was lower (approximately 25 kg/m²).

Degludec insulin once daily alone versus other long-acting insulin analogs in adult T2DM patients Effects on HbA1c and fasting prebreakfast glycemia

The results of the BEGIN studies are detailed in Table 2. The BEGIN Once Long T2 study13 investigated insulin-naïve patients who had previously been treated only with oral antidiabetic agents. This 1-year study was extended,14 giving 2 years of follow-up, and showed a similar HbA1c reduction in the IDeg arm (−1.06%) and IGla arm (−1.19%), with an estimated treatment difference (ETD) of 95% CI (−0.04, 0.22) between the two groups after 52 weeks of treatment. The fasting prebreakfast glycemia reduction was greater in the IDeg arm than in the IGla arm (ETD IGla-IDeg = −0.43 mmol/L [−0.74, −0.13], *P*=0.005).13 Similar results were

Table 1 Protocols for studies comparing insulin degludec versus a comparator in T2DM patients

<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Population</th>
<th>Duration</th>
<th>Patient groups</th>
</tr>
</thead>
<tbody>
<tr>
<td>BEGIN Once Long13</td>
<td>2012</td>
<td>Insulin-naïve T2DM</td>
<td>1 years</td>
<td>IDeg (n=773) IGla (n=257)</td>
</tr>
<tr>
<td>BEGIN Once Long (extension)14</td>
<td>2013</td>
<td>Insulin-naïve T2DM</td>
<td>2 years</td>
<td>IDeg (n=289) IGla (n=146)</td>
</tr>
<tr>
<td>Begin Asia15</td>
<td>2013</td>
<td>Insulin-naïve T2DM</td>
<td>26 weeks</td>
<td>IDeg (n=744) IGla (n=248)</td>
</tr>
<tr>
<td>BEGIN Basal-Bolus Type 216</td>
<td>2012</td>
<td>Insulin-treated T2DM</td>
<td>1 year</td>
<td>IDeg (n=228) IGla (n=229)</td>
</tr>
<tr>
<td>BEGIN Flex T217</td>
<td>2013</td>
<td>Insulin-naïve T2DM or insulin-treated T2DM (BI)</td>
<td>26 weeks</td>
<td>IDeg (n=228) IGla (n=229)</td>
</tr>
<tr>
<td>BEGIN Easy AM18</td>
<td>2013</td>
<td>Insulin-naïve T2DM</td>
<td>26 weeks</td>
<td>IDeg (n=233) IGla OD (n=234)</td>
</tr>
<tr>
<td>BEGIN Easy PM19</td>
<td>2013</td>
<td>Insulin-naïve T2DM</td>
<td>26 weeks</td>
<td>IDeg (n=233) IGla OD (n=234)</td>
</tr>
<tr>
<td>BEGIN Low Volume29</td>
<td>2013</td>
<td>Insulin-naïve T2DM</td>
<td>26 weeks</td>
<td>IDeg 200 U/mL (n=228) IGla (n=228)</td>
</tr>
<tr>
<td>BEGIN Compare20</td>
<td>2014</td>
<td>Insulin-treated T2DM (BI)</td>
<td>22 weeks</td>
<td>IDeg 200 U/mL (n=186) IGla 100 U/mL (n=187)</td>
</tr>
<tr>
<td>Onishi et al13</td>
<td>2013</td>
<td>Japanese, insulin-naïve T2DM</td>
<td>26 weeks</td>
<td>IDegAsp (n=147) IGla (n=149)</td>
</tr>
<tr>
<td>BOOST Asia24</td>
<td>2015</td>
<td>Asian insulin-treated T2DM with one Bi or more (premixed or self-mixed insulin)</td>
<td>26 weeks</td>
<td>IDegAsp twice daily (n=282) BIAsp 30 twice daily (n=142)</td>
</tr>
<tr>
<td>BOOST24</td>
<td>2014</td>
<td>Insulin-treated T2DM with one Bi or more (premixed or self-mixed insulin)</td>
<td>26 weeks</td>
<td>IDegAsp twice daily (n=224) BIAsp 30 twice daily (n=222)</td>
</tr>
<tr>
<td>DUAL-II27</td>
<td>2014</td>
<td>Insulin-treated T2DM (BI)</td>
<td>26 weeks</td>
<td>IDegLira (n=199) IDeg (n=199)</td>
</tr>
<tr>
<td>BEGIN Victoza Add-On28</td>
<td>2014</td>
<td>Insulin-treated T2DM from IDeg arm of []</td>
<td>26 weeks</td>
<td>IDeg + liraglutide (n=88) IDeg + Asp (n=89) IDeg (n=236)</td>
</tr>
<tr>
<td>DUAL-II4</td>
<td>2014</td>
<td>Insulin-naïve T2DM</td>
<td>26 weeks</td>
<td>IDegLira (n=834) IDeg (n=414) Liraglutide (n=415)</td>
</tr>
</tbody>
</table>

Abbreviations: Bi, basal insulin; IA, aspart insulin; BIAsp 30, biphasic insulin aspart 30/70; IDeg, degludec insulin; IDegAsp, IDegAsp combination; IDegflex, degludec insulin three times a week in the morning; IDegflex flex, degludec insulin with forced flexible scheme; IDet, detemir insulin; IGla, glargine insulin; Lira, liraglutide; OD, once daily; T2DM, type 2 diabetes mellitus.
found in the extension phase,\textsuperscript{14} i.e., no significant difference in 
HbA\textsubscript{1c} reduction between the two arms (ETD IGla-IDeg to 
0.07% (−0.07, 0.22; P=0.339) and a greater FPG reduction in 
the IDeg arm versus the IGla arm (ETD IDeg-IDeg to −0.36% 
[−0.67, −0.05], P=0.021). When comparing the BEGIN Once 
Long studies, the BEGIN Once Long Asia study also 
compared IDeg versus IGla in insulin-naïve patients, but for 
a shorter 26-week study period.\textsuperscript{15} After 26 weeks of follow-up, 
both groups showed a comparable reduction in 
HbA\textsubscript{1c}, with an 
ETD at 0.11% (−0.03, 0.24). However, reduction of fasting 
plasma glucose was slightly higher in the IGla arm although 
the ETD was not statistically significant (ETD IDeg − IGla, 
−0.09 mmol/L [−0.41, 0.23], P=0.59).\textsuperscript{15}

The BEGIN Basal-Bolus T2 study\textsuperscript{16} included T2DM 
patients who had been treated with insulin for more than 3 
months prior to enrollment. The insulin therapy had been 
delivered as either once-daily basal insulin or in an intensified 
dosing pattern of basal bolus. There was a similar statistically 
significant HbA\textsubscript{1c} reduction in the IDeg group (−1.10%) and 
the IGla group (−1.18%) with an ETD of 0.08% (−0.05, 0.21). 
However, there was a greater (albeit not statistically 
significant) reduction of fasting plasma glucose (ETD IGla − IDeg, 
−0.29 [−0.65, 0.06], P=0.1075) in the IDeg group.\textsuperscript{16}

Finally, the BEGIN Flex T2 study\textsuperscript{17} evaluated the 
flexibility of IDeg administration in insulin-naïve T2DM 
patients versus those treated with basal insulin therapy. The 
protocol involved randomizing patients into three treatment 
arms, i.e., IGla, IDeg, and IDeg\textsubscript{Forced-Flex}. IDeg\textsubscript{Forced-Flex} 
was a prespecified, rotating morning and evening dosing schedule 
that created an 8–40-hour interval between doses. The study 
showed similar efficacy with regard to reduction of HbA\textsubscript{1c} 
in the IDeg\textsubscript{Forced-Flex} and IGla arms, i.e., −1.28% and −1.26%, 
respectively (ETD IGla − IDeg\textsubscript{Forced-Flex} 
0.04 [−0.12, 0.20]). Surprisingly, in this study, IDeg seemed lower in efficacy 
compared with the other two arms, with an HbA\textsubscript{1c} reduction 
of −1.07%. The study did not include a statistical comparison 
between IDeg and the two other arms in terms of 
HbA\textsubscript{1c} reduction. However, final difference in HbA\textsubscript{1c}, between 
the IDeg and IDeg\textsubscript{Forced-Flex} arms was not significant (ETD 
IDeg − IDeg\textsubscript{Forced-Flex} −0.13% [−0.29, 0.03]).

### Table 2: Efficacy of insulin degludec 100 U/mL versus glargine insulin on HbA\textsubscript{1c} and fasting blood glucose in patients with type 2 diabetes mellitus

<table>
<thead>
<tr>
<th>Study</th>
<th>Arm</th>
<th>HbA\textsubscript{1c} (% ± SD)</th>
<th>HbA\textsubscript{1c} Reduction (% ± SD)</th>
<th>ETD (%) for HbA\textsubscript{1c} Reduction (CI 95%)</th>
<th>FBG (mmol/L±SD)</th>
<th>FBG Reduction (mmol/L)</th>
<th>ETD (mmol/L) for FPG (CI 95%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BEGIN Once Long\textsuperscript{13}</td>
<td>IDeg (n=773)</td>
<td>7.1</td>
<td>−1.06 (±1.01)</td>
<td>0.09 [−0.04, 0.22]</td>
<td>5.9</td>
<td>−3.81±3.04</td>
<td>−0.43 [−0.74, −0.13]</td>
</tr>
<tr>
<td></td>
<td>IGla (n=257)</td>
<td>7.0</td>
<td>−1.19 (±0.97)</td>
<td>P = NA</td>
<td>6.4</td>
<td>−3.31±2.87</td>
<td>0.005</td>
</tr>
<tr>
<td>BEGIN</td>
<td>IDeg (n=773)</td>
<td>7.0±0.9</td>
<td>NA</td>
<td>0.07 [−0.07, 0.22]</td>
<td>5.56±1.82</td>
<td>−4.17</td>
<td>−0.36 [−0.67, −0.05]</td>
</tr>
<tr>
<td></td>
<td>IGla (n=257)</td>
<td>6.9±0.8</td>
<td>NA</td>
<td>P=0.339</td>
<td>5.93±1.69</td>
<td>−3.56</td>
<td>P=0.021</td>
</tr>
<tr>
<td>BEGIN Once Long\textsuperscript{14}</td>
<td>IDeg (n=289)</td>
<td>7.2</td>
<td>−1.24</td>
<td>0.11 [−0.03, 0.24]</td>
<td>5.5</td>
<td>2.88</td>
<td>−0.09 [−0.41, 0.23]</td>
</tr>
<tr>
<td></td>
<td>IGla (n=146)</td>
<td>7.1</td>
<td>−1.35</td>
<td>P = NA</td>
<td>5.7</td>
<td>2.97</td>
<td>P=0.59</td>
</tr>
<tr>
<td>BEGIN</td>
<td>IDeg (n=744)</td>
<td>NA</td>
<td>−1.10</td>
<td>NA</td>
<td>NA</td>
<td>−2.3</td>
<td>−0.29 [−0.65, 0.06]</td>
</tr>
<tr>
<td></td>
<td>IGla (n=248)</td>
<td>NA</td>
<td>−1.18</td>
<td>P = NA</td>
<td>NA</td>
<td>−2.0</td>
<td>P=0.1075</td>
</tr>
<tr>
<td>BEGIN Basal-Bolus Type 2\textsuperscript{14}</td>
<td>IDeg (n=226)</td>
<td>NA</td>
<td>−1.07</td>
<td>IDeg\textsubscript{Forced-Flex} versus IDeg NA</td>
<td>5.8</td>
<td>NA</td>
<td>IDeg\textsubscript{Forced-Flex} versus IDeg NA</td>
</tr>
<tr>
<td></td>
<td>IGla (n=246)</td>
<td>NA</td>
<td>−1.18</td>
<td>0.08 [−0.05, 0.21]</td>
<td>NA</td>
<td>−2.0</td>
<td>P=0.04</td>
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<tr>
<td>BEGIN Flex T2\textsuperscript{17}</td>
<td>IDeg\textsubscript{Forced-Flex} (n=230)</td>
<td>NA</td>
<td>−1.28</td>
<td>IDeg\textsubscript{Forced-Flex} versus IGla</td>
<td>5.8</td>
<td>NA</td>
<td>IDeg\textsubscript{Forced-Flex} versus IGla</td>
</tr>
<tr>
<td></td>
<td>IGla (n=229)</td>
<td>NA</td>
<td>−1.26</td>
<td>0.04 [−0.12, 0.20]</td>
<td>6.2</td>
<td>NA</td>
<td>−0.42 [−0.82, −0.02]</td>
</tr>
</tbody>
</table>

**Note:** ETD (mmol/L) for FPG in favor of IGla, mean ± SD.

**Abbreviations:** CI, confidence interval; ETD, estimated treatment difference; FPG, fasting plasma glucose; IDeg, insulin degludec; IDeg\textsubscript{Forced-Flex}, insulin degludec with forced flexible scheme; IDet, detemir insulin; IGla, glargine insulin; NA, not available; SD, standard deviation.
Table 3  Efficacy of insulin degludec 100 U/mL versus glargine insulin on hypoglycemia events in patients with type 2 diabetes mellitus

<table>
<thead>
<tr>
<th>Study</th>
<th>Arm</th>
<th>Overall hypoglycemia (episodes per patient/year)</th>
<th>Estimate ratio of overall hypoglycemia [CI 95%]</th>
<th>Severe hypoglycemia (episodes per patient/year)</th>
<th>Estimate ratio of severe hypoglycemia [CI 95%]</th>
<th>Nocturnal hypoglycemia (episodes per patient/year)</th>
<th>Estimate ratio of nocturnal hypoglycemia [CI 95%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>BEGIN Once Long</td>
<td>iDeg (n=773)</td>
<td>1.52</td>
<td>0.82 [0.64–1.04]</td>
<td>0.003</td>
<td>0.14 [0.03–0.70]</td>
<td>0.25</td>
<td>0.64 [0.42–0.98]</td>
</tr>
<tr>
<td></td>
<td>iGla (n=257)</td>
<td>1.85</td>
<td></td>
<td>0.023</td>
<td></td>
<td>0.39</td>
<td></td>
</tr>
<tr>
<td>BEGIN Once Long (extension)</td>
<td>iDeg (n=773)</td>
<td>1.72</td>
<td>0.84 [0.68–1.04]</td>
<td>0.006</td>
<td>0.31 [0.11–0.85]</td>
<td>0.27</td>
<td>0.57 [0.40–0.81]</td>
</tr>
<tr>
<td></td>
<td>iGla (n=146)</td>
<td>2.05</td>
<td></td>
<td>0.021</td>
<td></td>
<td>0.39</td>
<td></td>
</tr>
<tr>
<td>BEGIN Once Asia</td>
<td>iDeg (n=289)</td>
<td>3.0</td>
<td>0.82 [0.60–1.11]</td>
<td>0</td>
<td>NA</td>
<td>0.8</td>
<td>0.62 [0.38–1.04]</td>
</tr>
<tr>
<td></td>
<td>iGla (n=146)</td>
<td>3.7</td>
<td></td>
<td>0</td>
<td>1 episode</td>
<td>1.2</td>
<td>P=0.07</td>
</tr>
<tr>
<td>BEGIN Basal-Bolus Type 2</td>
<td>iDeg (n=744)</td>
<td>11.09</td>
<td>0.82 [0.69–0.99]</td>
<td>0.06</td>
<td>NA</td>
<td>1.39</td>
<td>0.75 [0.58–0.99]</td>
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<tr>
<td></td>
<td>iGla (n=248)</td>
<td>13.63</td>
<td></td>
<td>0.05</td>
<td></td>
<td>1.84</td>
<td></td>
</tr>
<tr>
<td>BEGIN Flex T2</td>
<td>iDeg (n=226)</td>
<td>3.6</td>
<td>iDeg/twpm/ideg</td>
<td>NA</td>
<td>Two episodes in each group</td>
<td>0.6</td>
<td>iDeg/twpm/ideg</td>
</tr>
<tr>
<td></td>
<td>iGla (n=230)</td>
<td>3.6</td>
<td>iDeg/twpm/ideg</td>
<td>NA</td>
<td></td>
<td>0.6</td>
<td>iDeg/twpm/ideg</td>
</tr>
<tr>
<td></td>
<td>iDeg (n=229)</td>
<td>3.5</td>
<td>iDeg/twpm/ideg</td>
<td>NA</td>
<td></td>
<td>0.8</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: iDeg, insulin degludec; iDeg/twpm/ideg, insulin degludec with flexible scheme; iDet, detemir insulin; IGla, glargine insulin; NA, not available; NS, not statistically significant.

The ultra-long duration of action of iDeg allows longer injection intervals while maintaining consistent glycemic control, so it might be beneficial in specific patient populations, especially insulin-naïve T2DM patients. BEGIN Easy AM/BEGIN Easy PM study published their latest results for the BEGIN Basal-bolus T2 cohort. After 78 weeks of follow-up, the overall rate of hypoglycemia was 24% lower (P=0.011) and the rate of nocturnal hypoglycemia was 31% lower (P=0.016) in the iDeg group, with similar glycemic control in both groups.

A recent meta-analysis of five studies in T2DM showed higher efficacy of iDeg versus IGla in reduction of overall hypoglycemia (estimated rate ratio [ERR] 0.83 [0.74–0.94]) and nocturnal hypoglycemia (ERR 0.68 [0.57–0.82]). However, comparable efficacy was reported for both anti-glycemic agents in reduction of severe hypoglycemia (ERR 0.81 [0.42–1.56]). Finally, the same meta-analysis reported a statistically significant reduction in overall (~25%) and nocturnal (~38%) hypoglycemia rates during the maintenance period. These data suggest a long-term beneficial effect of iDeg after its optimal dose is achieved.

Towards a therapeutic regimen with iDeg three times a week?

The ultra-long duration of action of iDeg allows longer injection intervals while maintaining consistent glycemic control, so it might be beneficial in specific patient populations, especially insulin-naïve T2DM patients. BEGIN Easy AM/BEGIN Easy PM study published their latest results for the BEGIN Basal-bolus T2 cohort. After 78 weeks of follow-up, the overall rate of hypoglycemia was 24% lower (P=0.011) and the rate of nocturnal hypoglycemia was 31% lower (P=0.016) in the iDeg group, with similar glycemic control in both groups.

A recent meta-analysis of five studies in T2DM showed higher efficacy of iDeg versus IGla in reduction of overall hypoglycemia (estimated rate ratio [ERR] 0.83 [0.74–0.94]) and nocturnal hypoglycemia (ERR 0.68 [0.57–0.82]). However, comparable efficacy was reported for both anti-glycemic agents in reduction of severe hypoglycemia (ERR 0.81 [0.42–1.56]). Finally, the same meta-analysis reported a statistically significant reduction in overall (~25%) and nocturnal (~38%) hypoglycemia rates during the maintenance period. These data suggest a long-term beneficial effect of iDeg after its optimal dose is achieved.

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addition, the overall hypoglycemic risk was increased in the IDeg3TW arm versus the IGla once daily arm, with an ERR of 1.58 (1.03, 2.43), and the rate of nocturnal confirmed hypoglycemia was higher in the IDeg3TW group than in the IGla once daily group (ERR 2.12 (1.08, 4.16). The authors concluded that they did not recommend use of IDeg3TW in the morning or evening for insulin-naïve T2DM patients.

**Insulin degludec: which titration algorithm is the most effective?**

The ultra-long duration of action of IDeg, ie, more than 24 hours, might not be relevant in dose adjustment from fasting prebreakfast glyemia, which is commonly used for titration of other long-acting analogs. The BEGIN Once Single Use study aimed to identify the best titration modality for adjusting the IDeg dose, using a “simple” algorithm allowing dose adjustment according to a prebreakfast self-measured plasma glucose versus a “step-wise” algorithm allowing dose adjustment according to the lowest prebreakfast self-measured plasma glucose in the last 3 days. Patients (n=222) were randomly assigned into two arms, ie, IDegSimple and IDegStep-Wise. The HbA1c decreased similarly in both arms (−1.09% in IDegSimple versus −0.93% in IDegStep-Wise) with an ETD of −0.16% (−0.39, 0.07). The overall and nocturnal hypoglycemia rates were not significantly different. The authors concluded that both algorithms showed comparable effectiveness and tolerability, suggesting the possibility of using the two algorithms according to patient preference.

**Other forms of insulin degludec for T2DM patients**

Other forms of IDeg have been developed for use in T2DM patients, including IDeg–IAsp and IDeg–liraglutide (IDeg–Lira) preparations; the former is a soluble coformulation of IDeg and IAsp (70% IDeg and 30% IAsp). In a study by Onishi et al, IDegAsp was compared with IGla alone. After 26 weeks, mean HbA1c was lower in the IDegAsp arm (7.0%) than in IGla arm (7.3%), with an ETD (IDegAsp – IGla) of −0.28% (−0.46, −0.10, P<0.01). Further, IDegAsp was associated with lower rates of overall and nocturnal hypoglycemia versus IGla, but the ERRs of 0.73 (0.50, 1.08) and 0.75 (0.34, 1.64) were not statistically significant.

In the BOOST trials, IDegAsp was compared with premixed biphasic insulin aspart 30, each administered twice daily. Both studies showed comparable HbA1c reduction, with superior reduction in fasting plasma glucose for IDegAsp (ETD −1.06 mmol/L [−1.43, −0.70], P<0.001 in the Kaneko et al trial and −1.14 mmol/L [−1.53, −0.76], P<0.001) in the Fulcher et al trial with a lower total daily insulin dose in IDegAsp trial with a lower total daily insulin dose in IDegAsp and IDegIAsp trial versus biphasic insulin aspart 30. In the former trial, rates of overall and severe hypoglycemia were similar in both groups, while the rate of confirmed nocturnal hypoglycemia was lower with IDegAsp (ERR 0.67 [0.43, 1.06]), but not statistically significant. In the trial by Fulcher et al, rates of overall and confirmed nocturnal hypoglycemia were lower in the IDegAsp group (ERR 0.68 [0.52, 0.89]) P=0.0049 and 0.27 [0.18, 0.41], P<0.0001, respectively). In the maintenance period, rates of severe hypoglycemia were also lower for IDegAsp (P=0.04), with one episode in the IDegAsp group versus 13 episodes in the biphasic insulin aspart 30 group. These two studies suggest an equivalent efficacy of IDegAsp versus biphasic insulin aspart 30 for HbA1c reduction, with a decreased risk of overall and nocturnal hypoglycemia and probably a reduction of severe hypoglycemia in the maintenance period.

The ADA and EASD recently approved the combination of basal insulin and a GLP-1 receptor agonist. IDegLira, a novel combination of basal IDeg and the long-acting GLP-1 analog liraglutide, was then developed for the treatment of T2DM patients as a once-daily, single subcutaneous injection. The pharmacological rationale for use of this combination is that lower fasting blood glucose levels can be obtained using IDeg and liraglutide, while liraglutide can also modestly reduce post-prandial glucose excursions. The DUAL-1 showed noninferiority of IDegLira versus IDeg alone, with an ETD of −0.47% (−0.58, −0.36, P<0.0001). The same trial showed the superiority of IDegLira versus liraglutide (ETD −0.64% (−0.75, −0.53, P<0.0001)). The DUAL-2 trial showed the superiority of IDegLira versus IDeg alone in terms of HbA1c reduction (−1.9% in the IDegLira arm versus −0.9% in the IDeg arm; ETD −1.1% [−1.3, −0.8], P<0.0001) and mean weight reduction. However, the hypoglycemia rate was comparable in both groups.

Finally, the BEGIN Victoza Add-On study compared IDeg + IAsp versus IDeg + liraglutide in a T2DM population from the BEGIN Once Long extension cohort. Patients were randomized to IDeg + liraglutide and IDegAsp if the target HbA1c (≥7.0%) was not reached at 104 weeks. If their HbA1c was <7.0%, patients continued IDeg in a third non-randomized arm. The results showed that IDeg + liraglutide (−0.74%) reduced HbA1c significantly more than IDeg + IAsp (−0.39%), with an ETD of −0.32% (−0.53, −0.12, P=0.0024). Further, patients on IDeg + liraglutide had significantly less overall and nocturnal hypoglycemia, and significantly greater weight loss (−2.8 kg) versus patients on IDeg + IAsp (−0.9 kg)
with an ETD (IDeg + liraglutide – IDeg + IAsp) of −3.75 kg (−4.70, −2.79, \(P<0.0001\)). This study suggested that adding a long-acting GLP-1 analog such as liraglutide may be superior to adding a single daily dose of IAsp in the event of failure of IDeg + metformin to achieve \(\text{HbA1c}\) reduction and weight loss, as well as in prevention of hypoglycemic events.\(^{28}\)

A more concentrated form of IDeg (200 U/mL) has been also developed and tested in two studies, ie, BEGIN Low Volume\(^{29}\) and BEGIN Compare.\(^{30}\) BEGIN Low Volume compared IDeg (200 IU/mL) versus IGla and reported results similar to those of other trials in insulin-naïve T2DM patients, with equal efficacy in terms of \(\text{HbA1c}\) reduction and a comparable hypoglycemia rate.\(^{29}\) The BEGIN Compare study compared IDeg 200 IU/mL with IDeg 100 IU/mL, and the results showed comparable efficacy between the groups with regard to glycemic control and hypoglycemia rates.\(^{30}\)

**Insulin degludec and quality of life in T2DM patients**

In addition to assessment of efficacy and safety criteria, there are at least two hypotheses concerning assessment of quality of life related to the use of an antiglycemic agent. T2DM patients treated with IDeg may feel less stress than those treated with antiglycemic agents administered on a strict schedule. Moreover, the significant reduction in nocturnal hypoglycemia rate achieved by IDeg could be a factor in improving quality of life in T2DM patients. It is known that hypoglycemia-related anxiety is associated with deterioration of quality of life in patients with diabetes.\(^{31}\) A recent meta-analysis\(^{32}\) addressed this issue using the Short-Form 36 questionnaire filled in by patients (T2DM patients in five trials and T1DM patients in one trial. These results were then processed and converted into a EuroQol-5D score. The meta-analysis concluded that there was a moderate but significant improvement in quality of life in patients treated with IDeg when compared with those treated with IGla. This improvement was independent of the flexibility of administration, ie, IDeg was injected at a fixed time once daily. Further studies are needed to better understand the benefits of IDeg on quality of life in T2DM patients.

**Insulin degludec safety and adverse effects**

**General adverse effects**

Several studies have evaluated the safety of IDeg. To date, no study has shown either more or serious adverse events in patients treated with IDeg compared with those treated with IGla. Moreover, according to this review, the proportion of patient drop outs was not different among studies. In terms of the IDeg doses used, tests in T2DM patients showed no obvious difference between insulin doses in the IDeg and IGla groups. According to the literature, comparable weight gain results were obtained in T2DM patients treated with IDeg compared with those treated with IGla. It is worth mentioning that immunological studies have found insignificant traces of anti-degludec antibody in T2DM patients.\(^{13,17}\)

As a result, the currently available data on use of IDeg in daily clinical practice seem reassuring. However, the limitations of trial duration and sample size in the current literature on the safety of IDeg preclude any results being able to be considered conclusive. Larger and longer duration randomized prospective trials on IDeg-related adverse events and serious adverse events are needed to obtain evidence-based and conclusive data. Two important criteria regarding the safety of IDeg need to be further evaluated, ie, cardiovascular safety and risk of neoplasia.

**Cardiovascular safety**

This systemic review highlights the limitations of the available published trials, ie, short observation periods and small sample sizes. Consequently, there are no current conclusive data on the cardiovascular safety of IDeg. However, a 2012 US Food and Drug Administration study of randomized Phase III trials in T1DM and T2DM patients reported a potential cardiovascular risk associated with IDeg. Indeed, in most of the trials comparing IDeg/IDegAsp with a comparator arm, the incidence of cardiovascular events was estimated according to composite criteria of MACE (major adverse cardiovascular events) and MACE+. The latter criterion estimated the occurrence of events during follow-up, ie, acute coronary syndrome, including unstable angina and myocardial infarction, stroke, and cardiovascular death. The definition of cardiovascular events according to the MACE criterion was stricter than that by MACE+ and excluded unstable angina. All events occurring 7 days after cessation of treatment were censored. According to the MACE+ criterion, 95/5,794 events in the IDeg/IDegAsp group versus 37/3,461 in the comparator group were observed (hazard ratio 1.30; 95% confidence interval 0.88–1.93). According to the MACE criterion, there were 70/5,794 events in the IDeg/IDegAsp group versus 21/3,461 in the comparator group (hazard ratio 1.67; 95% confidence interval 1.01–2.75). The number of cardiovascular events according to the MACE criterion was then more statistically significant in IDeg/IDegAsp group.

Given the above data, although limited and not representing those in the general diabetes patient population, the US
Food and Drug Administration has delayed the marketing authorization for IDeg, and requested additional prospective randomized studies on the cardiovascular safety of IDeg. In contrast, the European Medicines Agency and other national health authorities have granted marketing authorization for IDeg. Such divergence in marketing authorization approvals has generated debate on the potential cardiovascular risk of IDeg. To date, there is no relevant evidence-based rationale to explain any potential IDeg-associated cardiovascular risk. The DEVOTE (NCT 01959529) trial is in progress, and is aiming to enroll 7,500 T2DM subjects at high cardiovascular risk (age ≥50 years with a history of cardiovascular disease or diabetic nephropathy or age ≥60 years with cardiovascular risk factors) in order to evaluate such an hypothesis. The results of this trial are expected to be available in 2018. However, according to Novo Nordisk, it seems that the US Food and Drug Administration has accepted a resubmitted marketing authorization application based on interim analysis of data from the DEVOTE study.

Risk of neoplasia

In 2009, some cohort studies raised the issue of a potentially increased risk of cancer in T2DM patients treated with IGla. To date, no study has demonstrated the increased risk of cancer in T2DM patients on IGla therapy. The suspicion of a potential risk of neoplasia in T2DM patients treated with IGla has been explained based on the theoretical capacity of IGla to have insulin-like growth factor 1 (IGF-1)-like activity after its interaction with the IGF-1 receptor. Indeed, several studies have shown a greater affinity of IGla compared with that of human insulin for the IGF-1 receptor in vitro. This greater affinity can be explained by the addition of arginine residues at positions 31 and 32 of the beta chain, thus suggesting an increased risk of cancer in patients treated with IGla. In contrast, in vivo metabolism of IGla in blood shows low mitogenic activity due to the low affinity of its primary metabolite for the IGF-1 receptor. In respect to IDeg, data from the study by Nishimura et al showed a lower affinity of IDeg for the IGF-1 receptor in comparison with that of human insulin. These results theoretically suggest the absence of increased risk in patients treated with IDeg. However, clinically, the duration of this study does not allow an evidence-based conclusion to be reached. The results of the BEGIN Once Long trial in insulin-naïve T2DM patients followed for 1 year showed a very low rate of cancer in the IDeg and IGla groups (1.0% [8/766] in the IDeg group and 0.8% [2/257] in the IGla group). A prospective cohort with long-term follow-up is therefore needed to better assess this risk.

Conclusion

In conclusion, the study results suggest that IDeg provides glycemic control and reduction of nocturnal hypoglycemia comparable with that achieved by other long-acting analogs (IGla and IDet) in patients with T2DM. The rate of severe hypoglycemia also seems to be reduced when using IDeg therapy; however, long-term follow-up is warranted for monitoring of possible but relatively infrequent adverse events. On the other hand, a 2012 US Food and Drug Administration study revealed a potential IDeg-associated cardiovascular risk. Future prospective evaluation of large cohorts of T2DM patients treated with IDeg, with long-term follow-up, can provide more relevant information on the safety of IDeg therapy.

Disclosure

We declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of this work.

References


