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ORIGINAL RESEARCH ARTICLE

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Glucose-Lowering Effect of Insulin Degludec is Independent of Subcutaneous Injection Region

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Abstract

Background and Objectives Patients with diabetes mellitus inject insulin in different regions of the body. This study investigated the pharmacokinetic and pharmacodynamic properties of insulin degludec (IDeg), a new-generation once-daily basal insulin with an ultra-long duration

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H. F. Thomsen Novo Nordisk A/S, Aalborg, Denmark of action, after subcutaneous (SC) administration in different injection regions.

Methods In this study, 20 healthy subjects received single SC doses of IDeg (0.4 U/kg; separated by 13–21 days) in the thigh, abdomen and deltoid in a randomised, openlabel, single-centre, single-dose, complete crossover trial. Each dose was followed by a 24-h euglycaemic clamp and 120-h pharmacokinetic blood sampling. The obtained pharmacokinetic/pharmacodynamic profiles were extrapolated to steady state by simulation using a pharmacokinetic/ pharmacodynamic model.

Results Total IDeg exposure [area under the IDeg serum concentration-time curve 0-120 h after a single dose (AUC_{IDeg,0-120h,SD})] and maximum serum concentration [maximum IDeg serum concentration after a single dose (C_{max.IDeg.SD})] were higher (6-7 and 23-27 %, respectively) following a single SC dose in the deltoid or abdomen, compared with the thigh, as also observed with other insulin preparations. No statistical difference was observed in these measures between deltoid and abdominal administration. No pronounced differences were observed in the glucose-lowering effect of IDeg [area under the glucose infusion rate (GIR) curve 0-24 h after a single dose $(AUC_{GIR,0-24h,SD})$ and maximum GIR after a single dose (GIR_{max.SD})] when injected in the thigh, abdomen or deltoid (AUC_{GIR,0-24h,SD} 2,572, 2,833 and 2,960 mg/kg, respectively). Simulated mean steady-state pharmacokinetic and pharmacodynamic profiles supported a flat and stable IDeg exposure and effect regardless of injection region, with comparable total glucose-lowering effects [area under the GIR curve at steady state $(AUC_{GIR \tau SS})$] between the thigh, abdomen and deltoid.

Conclusions These findings support administering IDeg SC in the thigh, upper arm or abdominal wall without affecting IDeg absorption or effect at steady state.

Key Points

Insulin degludec (IDeg) has an ultra-long duration of action that is mediated by the formation of stable multi-hexamers in the subcutaneous (SC) tissue upon injection, resulting in a soluble depot from which IDeg monomers are slowly released into the circulation.

This study found that IDeg has a flat and stable glucose-lowering effect which is independent of injection region (thigh, abdomen, deltoid).

These results support administering IDeg SC in the thigh, upper arm or abdominal wall without affecting IDeg absorption or effect at steady state.

1 Introduction

Diabetes mellitus patients inject insulin in different regions of the body according to personal preference and activity pattern [1-3]. However, differences can exist in the pharmacokinetic and pharmacodynamic properties of insulin following subcutaneous (SC) administration in different regions [4-10].

Insulin degludec (IDeg), a new-generation basal insulin with an ultra-long duration of action, developed for oncedaily administration, has a distinct mechanism of protraction [11]. In contrast to other basal insulin preparations, which form crystals or precipitate upon SC injection [12-15], IDeg stays in solution and forms stable multi-hexamers in the SC tissue from which IDeg monomers gradually separate, producing a slow absorption into the circulation with low day-to-day variability [11, 16, 17]. IDeg has a considerably longer half-life after SC administration than insulin glargine (25 vs. 12 h) [18] and exhibits a duration of action longer than 42 h [11, 16]. The soluble multi-hexamer formation, slow absorption and ultra-long duration of action of IDeg raise the possibility that any differences in pharmacokinetic and pharmacodynamic properties seen between injection regions after a single dose of IDeg may diminish at steady state.

In this single-dose study, we investigated the pharmacokinetic and pharmacodynamic properties of IDeg after SC dosing in the thigh, abdomen or deltoid (upper arm). In addition, a pharmacokinetic/pharmacodynamic model was developed based on the obtained data to simulate the pharmacokinetic and pharmacodynamic profiles of IDeg at steady state for each injection region.

2 Materials and Methods

2.1 Study Design

This randomised, open-label, five-period, single-centre (Profil, Germany), single-dose crossover trial was conducted in healthy subjects (ClinicalTrials.gov identifier: NCT01151072). The study protocol was reviewed and approved by the health authority (Bundesinstitut für Arzneimittel und Medizinprodukte) according to local regulations and by the ethics committee of Ärztekammer Nordrhein. The study was performed in accordance with the Declaration of Helsinki and its amendments, and in accordance with Good Clinical Practice as defined by the International Conference on Harmonisation. Subjects were informed of the risks and benefits of the trial and were informed that they could withdraw at any time for any reason. Consent was obtained in writing before any trialrelated activities, and the investigator retained the consent forms.

2.2 Subjects

Study subjects were healthy males or females aged 18–55 years, with a body mass index (BMI) of 18.0–27.0 kg/m² and fasting plasma glucose concentrations of ≤ 6.0 mmol/L (≤ 108 mg/dL). Key exclusion criteria for participation in the study included the use of prescription drugs within 3 weeks prior to screening, the use of non-prescription drugs (including over-the-counter medication, non-routine vitamins and herbal products) within 3 weeks prior to screening, and smoking.

2.3 Interventions and Pharmacokinetic Sampling

Following screening (Visit 1), subjects were randomised to predetermined dosing sequences consisting of five single doses of IDeg on five separate dosing visits (Visits 2–6). Dosing was conducted via SC injection of 0.4 U/kg body weight (BW) of IDeg in the thigh, abdomen or deltoid (upper arm); intramuscular (IM) injection of 0.4 U/kg BW of IDeg in the thigh area; or intravenous (IV) injection of 0.04 U/kg BW of IDeg. Only the methods and data from the SC dosing arms are reported here. IDeg was provided in 3 mL Penfill[®] cartridges (100 U/mL) (Novo Nordisk A/S, Bagsværd, Denmark) for dosing and administered (using a syringe and needle) into a lifted skinfold in either the anterior surface of the thigh, the lower abdominal wall (above the inguinal area) or the outer aspect of the deltoid area.

At each dosing visit, IDeg administration was followed by a 24-h euglycaemic clamp procedure (see below for description). Subjects attended dosing visits in a fasted state, and each subject remained in the clinic for 48 h after dosing, during which blood samples for pharmacokinetic analysis were taken frequently. Blood samples were also taken frequently for analysis of blood glucose concentrations. Subjects subsequently returned to the clinic at 24-h intervals. Blood samples were taken at these visits (at 72, 96 and 120 h post-dosing) for pharmacokinetic assessment. Dosing visits were separated by a washout period of 13–21 days. An interval of 7–21 days existed between the last of the five dosing visits and a subsequent follow-up visit (Visit 7).

2.4 Euglycaemic Clamp Procedure

Subjects remained fasted (with water *ad libitum*) and in a supine position for the euglycaemic clamp procedure (Biostator[®], MTB Medizintechnik, Amstetten, Germany); target blood glucose: 4.5 mmol/L (81 mg/dL). One to 6 hours before dosing, subjects received a variable IV infusion of human insulin [15 (I)U Actrapid® (Novo Nordisk A/S, Bagsværd, Denmark), 100 (I)U/mL in 49 mL saline and 1 mL of subject's blood] or glucose (20 %) to obtain the glucose clamp target concentration. The target glucose concentration was maintained for at least 1 h before dosing, without any glucose infusion. After dosing, the rate of insulin infusion, if any, was decreased gradually and terminated when glucose concentrations had declined by approximately 0.3 mmol/L (5 mg/dL). A variable IV glucose infusion was then initiated to maintain the clamp target concentration.

2.5 Data and Statistical Analyses

The primary objective of this study was to investigate the relative exposure among different SC administration regions in healthy subjects following single-dose administration. Secondary objectives were evaluation of the pharmacokinetic and pharmacodynamic profiles and the safety and tolerability of IDeg.

Serum concentrations of IDeg were measured using an IDeg-specific sandwich ELISA, with a lower limit of quantification of 20 pmol/L. The primary endpoint was the area under the IDeg serum concentration–time curve 0-120 h after a single dose (AUC_{IDeg,0-120h,SD}) given by SC administration in the thigh, abdomen or deltoid area. AUC_{IDeg,0-120h,SD} was derived by non-compartmental analysis using the linear trapezoidal technique based on observed values and actual measurement times between 0 and 120 h, with missing values interpolated. The log-transformed AUC_{IDeg,0-120h,SD} was analysed using an ANOVA method with injection region and treatment period as fixed factors and subject as a random effect. In order to account for potential heteroscedasticity, the error-term

was dependent on the injection region. The maximum IDeg serum concentration after a single dose ($C_{\max,IDeg,SD}$) was also assessed. $C_{\max,IDeg,SD}$ was derived from individual concentration-time curves and analysed using the same approach as for AUC_{IDeg,0-120h,SD}.

Pharmacodynamic endpoints included the area under the glucose infusion rate (GIR) curve 0–24 h after a single dose ($AUC_{GIR,0-24h,SD}$) and maximum GIR after a single dose ($GIR_{max,SD}$). GIR data were smoothed using the Loess smoothing technique (fixed smoothing parameter of 0.25). Pharmacodynamic endpoints were summarised using descriptive statistics.

To predict the steady-state pharmacokinetic and pharmacodynamic profiles of IDeg following SC pharmacokinetic/pharmacodynamic administration, а model using single-dose IDeg data from the current study was applied, with area under the concentrationtime curve (AUC) derived by non-compartmental analysis. The pharmacokinetic component of the model consisted of an absorption part with a depot compartment, a transit compartment, a bioavailability parameter, an absorption rate parameter and a transit rate parameter; and a disposition part with two compartments, two clearance parameters and two volume of distribution parameters. The pharmacodynamic component of the model linked the IDeg concentration to GIR by means of an effect compartment, a turnover parameter, an insulin sensitivity parameter and an underlying GIR baseline parameter. The parameters of the model were estimated in a population pharmacokinetic/pharmacodynamic setting, using a non-linear mixed-effects approach, which allowed individual sets of the ten parameters for each of the subjects included in the trial to be obtained. The bioavailability parameter and the absorption rate parameter were allowed to vary between injection regions for each subject. The values of the absorption rate parameter were subsequently calibrated based on information from the comprehensive clinical pharmacology programme of studies conducted with IDeg. The same calibration factor was applied for all subjects and all injection regions. Using the estimated individual parameters, a simulation of once-daily multiple dosing was conducted to obtain mean steady-state profiles. More specifically, once-daily multiple dosing for 6 days at a dose level of 0.4 U/kg was simulated by extrapolating the profile for each of the subjects, and for each injection region, and subsequently calculating the mean of the profiles on Day 6.

Safety endpoints, including adverse events (AEs), laboratory safety variables, physical examination, vital signs, ECG, hypoglycaemic episodes and local tolerability at injection site, were monitored and summarised using descriptive statistics.

3 Results

3.1 Subjects

Twenty-two subjects were screened; two subjects were screening failures (both had first-degree relatives with diabetes). Twenty subjects (17 males and three females) were randomised, and 19 subjects completed the trial. One subject withdrew from the trial for personal reasons after having completed the first three dosing visits (the subject did not complete SC and IM injection in the thigh). All 20 randomised subjects were included in the full analysis set and the safety analysis set.

The mean [standard deviation (SD)] age of subjects who were randomised to dosing sequences was 37.4 (9.5) years, the mean (SD) BW was 76.4 (12.0) kg and the mean (SD) BMI was 24.1 (2.4) kg/m². The majority of subjects (17/20) were caucasian, two were African American and one was Asian non-Indian.

3.2 Single-Dose and Simulated Steady-State Pharmacokinetics

Single-dose 24-h mean pharmacokinetic profiles are shown in Fig. 1a. Total exposure of IDeg (AUC_I. $_{Deg,0-120h,SD}$) was 6–7 % higher following a single SC injection in the deltoid or abdomen than in the thigh (Table 1). No difference in AUC_{IDeg,0-120h,SD} was found between the deltoid and abdomen. The maximum concentration of IDeg ($C_{max,IDeg,SD}$) was 23–27 % higher following a single SC injection in the deltoid or abdomen than in the thigh (Table 1). No difference in $C_{max,IDeg,SD}$ was found between the deltoid and abdomen.

Simulated mean pharmacokinetic profiles at steady state showed an even distribution of IDeg exposure across a 24-h dosing interval regardless of injection region (Fig. 1b). Simulated steady-state exposure [area under the insulin degludec serum concentration–time curve at steady state (AUC_{IDeg,\tau,SS})] was predicted to be ~8 % higher following injection in the deltoid or abdomen than in the thigh (Table 1).

The observed differences in $C_{\max,IDeg,SD}$ diminished at steady state; the maximum IDeg serum concentration at steady state ($C_{\max,IDeg,SS}$) was estimated to be 10 % higher following injection in the deltoid or abdomen than in the thigh (Table 1).

3.3 Single-Dose and Simulated Steady-State Pharmacodynamics

Single-dose 24-h mean pharmacodynamic profiles are shown in Fig. 2a. No pronounced differences in the



Fig. 1 a Mean 24-h pharmacokinetic profiles after a single subcutaneous dose of insulin degludec (0.4 U/kg), administered in the thigh, abdomen or deltoid. **b** Mean 24-h pharmacokinetic profiles when simulated to steady state after once-daily subcutaneous administration of insulin degludec (0.4 U/kg) in the thigh, abdomen or deltoid. Because there was no difference in total exposure (AUC_{IDeg,0-120h,SD}) between the deltoid and abdomen after a single dose, the pharmacokinetic profiles for these two injection regions are superimposed in the steady-state simulation. $AUC_{IDeg,0-120h,SD}$ area under the insulin degludec serum concentration–time curve 0–120 h after a single dose, *conc.* concentration, *IDeg* insulin degludec

glucose-lowering effect of IDeg (AUC_{GIR,0-24h,SD} and GIR_{max,SD}) were evident following a single SC injection in the thigh, abdomen or deltoid (Table 2). The glucose-lowering effect extended beyond 24 h in all subjects for all three SC injection regions.

In accordance with the simulated steady-state pharmacokinetic profiles, the simulated mean steady-state pharmacodynamic profiles showed an even distribution of glucose-lowering effect across a 24-h dosing interval (Fig. 2b). The simulated glucose-lowering effect of IDeg at steady state did not show any relevant differences among the three SC injection regions (Table 2). **Table 1** Pharmacokinetic endpoints of insulin degludec following a single subcutaneous injection in the thigh, abdomen or deltoid and simulation-based once-daily steady-state values

Pharmacokinetic endpoint	Mean estimate	Mean ratio [95 % CI]
Single dose		
AUC _{IDeg,0-120h,SD} (pmol·h/L)		
Thigh	76,679	
Abdomen	82,059	
Deltoid	81,135	
Deltoid vs. thigh		1.06 [1.01-1.10]
Abdomen vs. thigh		1.07 [1.03–1.11]
Abdomen vs. deltoid		1.01 [0.96–1.06]
$C_{\max,IDeg,SD}$ (pmol/L)		
Thigh	1,938	
Abdomen	2,388	
Deltoid	2,462	
Deltoid vs. thigh		1.27 [1.08–1.49]
Abdomen vs. thigh		1.23 [1.07–1.42]
Abdomen vs. deltoid		0.97 [0.84–1.12]
Steady state		
AUC _{IDeg, τ,ss} (pmol·h/L)		
Thigh	74,353	
Abdomen and deltoid	80,087	
Abdomen and deltoid vs. thigh		1.08 [NA]
C _{max,IDeg,SS} (pmol/L)		
Thigh	3,367	
Abdomen and deltoid	3,703	
Abdomen and deltoid vs. thigh		1.10 [NA]

Analyses based on 20 evaluations after injection in deltoid and abdomen, and 19 evaluations after injection in thigh. The steady-state simulation model assumes an individual clearance value for each subject regardless of injection region; thus, no variance estimates can reliably be calculated

 $AUC_{IDeg,0-120h,SD}$ area under the insulin degludec serum concentration-time curve 0–120 h after a single dose, $AUC_{IDeg,\tau,SS}$ area under the insulin degludec serum concentration-time curve at steady state, *CI* confidence interval, $C_{max,IDeg,SD}$ maximum insulin degludec serum concentration after a single dose, $C_{max,IDeg,SS}$ maximum insulin degludec serum concentration at steady state, *NA* not applicable

3.4 Safety

Most (six of nine) AEs were mild, and AEs were evenly distributed among the three SC dosing arms. No severe AEs or injection-site reactions were reported.

4 Discussion

The present study evaluated the pharmacokinetic and pharmacodynamic properties of IDeg following a single SC dose in the thigh, abdomen or deltoid region. A slightly higher total exposure of IDeg (by 6–7 %) was observed following a single SC injection in the abdomen or deltoid region than in the thigh. No pronounced differences were observed in the glucose-lowering effect of IDeg between single-dose SC injection in the thigh, abdomen and deltoid regions. Simulated mean steady-state pharmacodynamic profiles supported a flat and stable glucose-lowering effect of IDeg across a 24-h dosing interval, and were within the same range for all three injection regions.

Several previous examples exist of the pharmacokinetic and pharmacodynamic properties of insulin products being affected by SC injection region. For example, some studies have reported slower insulin absorption and decreased glucose-lowering effects of short-acting insulins when injected SC in the thigh than in the abdomen [4-8] or deltoid [4, 5]. The same finding has also been observed following administration of regular human insulin or neutral protamine Hagedorn (NPH) insulin in the thigh, compared with injection in the abdomen [7]. However, this is in contrast to data from a study comparing SC injection of NPH insulin in the abdomen or the thigh, which found no difference in measured pharmacokinetic parameters between the two regions [19]. A reason for this might be that the latter study (performed in only 11 people with type 1 diabetes) measured elimination rates from radiolabelled NPH insulin rather than pharmacokinetic parameters directly, and therefore might not have been appropriate to detect differences between injection regions.

With the advent of long-acting insulin products it is also important to determine whether any differences between injection regions are likely to occur with the use of these products. Existing data with radio-labelled insulin demonstrate that the absorption characteristics of insulin glargine are similar regardless of whether SC injection is conducted in the arm, abdomen or leg and no differences were seen in plasma exogenous insulin concentration or blood glucose levels between injection regions [20]. In contrast, with appropriate pharmacokinetic measurements, the AUC and maximum concentration of insulin detemir are reported to be higher (by approximately 10 and 20 %, respectively) following SC injection in the abdomen or deltoid than in the thigh [10]. The present study demonstrates a similar finding for IDeg.

When we tried to correlate this to the pharmacodynamics of IDeg, only small differences were seen in the glucose-lowering effect between the three injection regions following a single dose. Further, as IDeg reaches steady state at 2–3 days [21], multiple once-daily injections will contribute to the exposure and effect seen during a dosing interval at steady state, and it is supposed that the flat and consistent pharmacokinetic and pharmacodynamic profiles that are achieved at steady state would be less susceptible to minor differences in absorption rate. Indeed, the present



Fig. 2 a Mean 24-h pharmacodynamic profiles after a single subcutaneous dose of insulin degludec (0.4 U/kg), administered in the thigh, abdomen or deltoid. **b** Mean 24-h pharmacodynamic profiles when simulated to steady-state after once-daily subcutaneous administration of insulin degludec (0.4 U/kg) in the thigh, abdomen or deltoid. Because there was no difference in total exposure (AUC_{IDeg,0-120h,SD}) between the deltoid and abdomen after a single dose, the pharmacodynamic profiles for these two injection regions are superimposed in the steady-state simulation. $AUC_{IDeg,0-120h,SD}$ area under the insulin degludec serum concentration-time curve 0–120 h after a single dose, *GIR* glucose infusion rate

study supports this notion, since the inter-region differences in absorption rate observed after single-dose administration (as exemplified by $C_{\max,\text{IDeg,SD}}$) diminish at steady state (evaluated by means of simulation) and thus are expected to have limited clinical relevance.

The present study was conducted in healthy, young adults (18–55 years), rather than patients with diabetes, in order to include a relatively homogenous cohort of subjects and thereby facilitate the detection of differences between injection regions, in accordance with regulatory standards [22, 23]. With the inclusion of healthy subjects, a multiple-dose study with a clinically relevant dose would not have been acceptable, due to the risk of hypoglycaemia. Therefore, as discussed above, we extrapolated the pharmacokinetic and pharmacodynamic profiles of IDeg to steady state

 Table 2
 Glucose-lowering effect of insulin degludec following a single subcutaneous injection in the thigh, abdomen or deltoid and simulation-based once-daily steady-state values

Pharmacodynamic endpoint	Mean estimate	CV
Single dose		
AUC _{GIR,0-24h,SD} (mg/kg)		
Thigh	2,572	38
Abdomen	2,833	42
Deltoid	2,960	43
GIR _{max,SD} [mg/(kg·min)]		
Thigh	2.7	32
Abdomen	3.0	37
Deltoid	3.0	42
Steady state		
AUC _{GIR, τ, SS} (mg/kg)		
Thigh	4,719	
Abdomen and deltoid	5,005	
GIR _{max,SS} [mg/(kg·min)]		
Thigh	3.5	
Abdomen and deltoid	3.8	

Analyses based on 20 evaluations after injection in deltoid and abdomen, and 19 evaluations after injection in thigh. The steady-state simulation model assumes an individual clearance value for each subject regardless of injection region as well as individual parameters for each subject linking the insulin degludec concentration to GIR regardless of injection region; thus, no variance estimates can reliably be calculated

 $AUC_{GIR,0-24h,SD}$ area under the GIR curve 0–24 h after a single dose, $AUC_{GIR,\tau,SS}$ area under the GIR curve at steady state, CV coefficient of variation, GIR glucose infusion rate, $GIR_{max,SD}$ maximum GIR after a single dose, $GIR_{max,SS}$ maximum GIR at steady state

by means of simulation using a population pharmacokinetic/pharmacodynamic model based on data from the current trial. When including healthy subjects in a euglycaemic glucose clamp study it is important to address the potential confounding influence of endogenous insulin secretion during the glucose clamp. We did this by choosing a rather low clamp glucose target concentration of 4.5 mmol/L (81 mg/dL), which was successful in suppressing the endogenous insulin secretion throughout the 24-h clamp, as assessed by serum C-peptide concentrations (additional data given in the Electronic Supplementary Material: Online Resource 1).

5 Conclusion

The results of the present study show that IDeg has a flat and stable glucose-lowering effect independent of injection region and that IDeg can be administered SC in the thigh, deltoid or abdomen with clinically comparable glucoselowering effects at steady state. Acknowledgments The involvement of the clinical site (Profil, Neuss; Germany) is gratefully acknowledged. Medical writing assistance was provided by apothecom scopemedical, UK, funded by Novo Nordisk. This study was funded by Novo Nordisk.

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