Pharmacological properties of faster-acting insulin aspart vs insulin aspart in patients with type 1 diabetes receiving continuous subcutaneous insulin infusion: A randomized, double-blind, crossover trial

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Aim: To evaluate the pharmacological characteristics of faster-acting insulin aspart (faster aspart) compared with insulin aspart (IAsp) during continuous subcutaneous insulin infusion (CSII).

Methods: In this randomized, double-blind, crossover trial, 48 men and women aged 18 to 64 years with type 1 diabetes mellitus (T1DM) received faster aspart and IAsp as a 0.15 U/kg bolus dose via CSII, on top of a basal rate (0.02 U/kg/h), in a glucose clamp setting (target 5.5 mmol/L).

Results: After a CSII bolus dose, the pharmacokinetic/pharmacodynamic profiles for faster aspart were left-shifted compared with those for IAsp. For faster aspart vs IAsp, the early glucose-lowering effect (area under the curve for glucose infusion rate [GIR]0-30min) was approximately 2-fold higher (least squares means 24.9 vs 11.4 mg/kg; estimated ratio faster aspart/IAsp 2.18, 95% confidence interval [CI] [1.33; 5.04]; P = .002), onset of glucose-lowering effect (time to early 50% of maximum GIR) occurred 11.1 minutes earlier (41.1 vs 52.3 minutes; 95% CI faster aspart – IAsp [−15.4; −6.9]; P<.001), and offset of glucose-lowering effect (time to late 50% of maximum GIR) occurred 24.0 minutes earlier (214.7 vs 238.7 minutes; 95% CI [−38.9; −9.1]; P=.002). Likewise, significantly greater early exposure and significantly earlier onset and offset of exposure were observed for faster aspart vs IAsp. Faster aspart and IAsp were both well tolerated.

Conclusions: In patients with T1DM using CSII, faster aspart better mimics the endogenous prandial insulin secretion and action than does IAsp. Faster aspart therefore has the potential to provide clinical benefits over current rapid-acting insulins in the insulin pump setting.

KEYWORDS
insulin pump therapy, pharmacodynamics, pharmacokinetics, type 1 diabetes

1 | INTRODUCTION

Postprandial glycaemic control plays a substantial role in reaching recommended glycated haemoglobin (HbA1c) goals in diabetes. Compared with regular human insulin (RHI), rapid-acting insulin analogues (insulin aspart, insulin lispro and insulin glulisine) have provided better postprandial glucose control through an earlier and greater peak glucose-lowering effect. Nevertheless, absorption of current rapid-acting insulins occurs too slowly to adequately replicate endogenous prandial insulin action. Consequently, optimum postprandial glucose control remains a challenge in patients with diabetes.

Continuous subcutaneous insulin infusion (CSII) is increasingly used in people with diabetes. CSII therapy is associated with improved glycaemic control, lower risk of hypoglycaemia and greater patient convenience compared with multiple daily injection therapy. An insulin with a fast onset and fast offset of glucose-lowering effect might be particularly important in a CSII setting to further improve postprandial glucose control without the risk of late

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post-meal hypoglycaemia, and could also be a key factor in improving the performance of closed-loop, artificial pancreas systems.\textsuperscript{7}

Faster-acting insulin aspart (faster aspart) is insulin aspart in a new formulation that contains 2 well-known excipients, niacinamide and L-arginine. These are both listed in the US Food and Drug Administration inactive ingredient database, in products for injection, at higher concentrations than used in faster aspart.\textsuperscript{10} With faster aspart, niacinamide is responsible for faster initial absorption after subcutaneous administration and L-arginine serves as a stabilizing agent. In subjects with type 1 diabetes mellitus (T1DM), faster aspart administered by subcutaneous injection had a twice-as-fast onset of appearance, a 2-fold higher early exposure, and >50% greater early glucose-lowering effect compared with insulin aspart.\textsuperscript{11}

Current rapid-acting insulin analogues showed significant improvements over RHI in the pump setting with respect to glycaemic control and reduced hypoglycaemia.\textsuperscript{12,13} Likewise, faster aspart may provide benefits over current rapid-acting insulin analogues when used in a CSII regimen. In the present trial we evaluated, for the first time, the pharmacokinetic and pharmacodynamic properties of faster aspart compared with insulin aspart, using CSII in a euglycaemic clamp setting in subjects with T1DM.

2 MATERIALS AND METHODS

2.1 Study design

This was a randomized, single-centre (Profil, Neuss, Germany), double-blind, two-period, crossover trial in people with T1DM. The trial protocol was reviewed and approved by the local health authority (Bundesinstitut für Arzneimittel und Medizinprodukte) and by an independent ethics committee (Ärztekammer Nordrhein). The trial was performed in accordance with the Declaration of Helsinki and Good Clinical Practice, and registered at ClinicalTrials.gov (NCT01992588).

2.2 Participants

Eligible participants were men and women aged 18 to 64 years, who had been diagnosed with T1DM for ≥12 months and treated with multiple daily insulin injections or CSII for ≥12 months (total daily insulin dose <1.2 IU/kg/d and total daily bolus insulin dose <0.7 IU/kg/meal, with HbA\textsubscript{1c} <8.5%, body mass index (BMI) 18.5 to 28.0 kg/m\textsuperscript{2}, and fasting C-peptide ≥0.3 nmol/L). Individuals were excluded if they had clinically significant concomitant diseases, abnormal values in clinical laboratory screening tests, were smokers, or were currently treated with drug(s) that may interfere with glucose metabolism. Written informed consent was obtained before initiation of any trial-related activity.

2.3 Procedures

The trial consisted of a screening visit, 2 dosing visits separated by 3 to 12 days wash-out, and a follow-up visit.

Participants were randomized (1:1) to receive faster aspart (100 U/mL; 3 mL Penfill; Novo Nordisk, Bagsværd, Denmark) followed by insulin aspart (NovoRapid; 100 U/mL; 3 mL Penfill; Novo Nordisk) or vice versa. The trial was double-blind and the computer-generated randomization scheme was prepared by Clinical Supplies Coordination, Novo Nordisk A/S (Måløv, Denmark). Based on the randomization scheme, trial products were packed, in a double-blind manner, in boxes specific for each randomization number before delivery to the clinical site.

The trial products were administered by CSII using a MiniMed Paradigm Veo\textsuperscript{TM}754 insulin pump with a Paradigm Reservoir 3.0 mL and a Quick-set infusion set with a 6-mm cannula and a 23-inch tube (Medtronic MiniMed, Northridge, California). The cannula was inserted subcutaneously in the region of the lower abdominal wall above the inguinal area. One infusion site was used throughout the first dosing visit, while another infusion site within the same region was used throughout the second dosing visit.

At the dosing visits, participants came to the clinical site at 5:00 PM, after fasting since 10:00 AM (water and ≤20 g of rapidly absorbable carbohydrate to prevent hypoglycaemia were allowed). Participants using multiple daily insulin injections were not allowed to use insulin degludec ≤72 hours pre-dose and insulin detemir or insulin glargine ≤48 hours pre-dose (NPH insulin could be used instead). The last injection of NPH insulin or other intermediate-acting insulin had to occur at least 22 hours pre-dose, the last injection or bolus administration by CSII of insulin aspart had to occur at least 12 hours pre-dose (RHI could be used instead) and the last injection of RHI or other short-acting insulin (other than insulin aspart) had to occur at least 6 hours pre-dose. Participants using CSII had to discontinue the basal rate at least 3 hours pre-dose (or at least 8 hours pre-dose if using insulin aspart).

Participants were excluded from the dosing visit if they had experienced hypoglycaemia (plasma glucose ≤ 3.9 mmol/L) ≤24 hours pre-dose.

At 7:00 PM, participants received faster aspart or insulin aspart at an initial priming dose of 0.08 U/kg, followed by a basal rate of 0.02 U/kg/h for 27 hours, on top of which a single bolus dose of 0.15 U/kg was given after 13 hours (i.e. after 22 hours of fasting: Figure S1, Appendix S1). The duration of bolus infusion administered by the insulin pump was 3.7 to 8.3 minutes, depending on absolute dose.

A euglycaemic clamp procedure was initiated 30 minutes before the priming dose using ClampArt (Profil). Participants received a variable intravenous infusion of RHI (15 IU Actrapid [100 IU/mL; Novo Nordisk] in 49 mL saline and 1 mL of the participant’s blood) or 20% glucose to achieve the blood glucose clamp target level of 5.5 mmol/L. The intravenous RHI infusion (if any) was only allowed until 8 hours before the bolus dose of trial product. The clamp continued for 14 hours after the bolus dose. The quality of the conducted clamps is shown in Figure S2, Appendix S1.\textsuperscript{14}

Blood samples for pharmacokinetic assessment were drawn frequently at prespecified time points from before the insulin priming dose until the end of the CSII (27 hours after the priming dose).

2.4 Assessments and endpoints

Free serum insulin aspart concentrations (polyethylene glycol-precipitated) were measured using a validated insulin aspart specific enzyme-linked immunosorbent assay. The intravenous glucose
infusion rate (GIR) needed to keep blood glucose at the clamp target level was recorded every minute during the glucose clamp. Safety assessments included adverse events, local tolerability at the infusion site, hypoglycaemic episodes (defined as "severe" when they were either "severe" as according to the American Diabetes Association, i.e. requiring third party assistance, or verified by a plasma glucose level of <3.1 mmol/L), laboratory safety variables, physical examination, vital signs and ECG.

Endpoints to evaluate onset of exposure and glucose-lowering effect included: time to early 50% of maximum insulin concentration ($t_{Early \;50\% \;C_{max}}$); time to maximum insulin concentration ($t_{max}$); time to early 50% of maximum GIR ($t_{Early \;50\% \;GIR_{max}}$); and time to maximum GIR ($t_{GIR_{max}}$). Endpoints to evaluate early exposure and glucose-lowering effect included: time to early 50% of maximum insulin concentration ($t_{Early \;50\% \;C_{max}}$); estimated difference [95% CI] faster aspart vs insulin aspart $t_{max}$; time to 50% of maximum insulin aspart concentration was also observed for faster aspart vs insulin aspart ($t_{max}$; −25.7 minutes [−34.3; −17.1]; $P < .001$). For graphical reasons, there are some discrepancies between the length of the arrows and the actual mean treatment differences. This is attributable to the fact that the estimated mean treatment differences are derived from all participants’ individual treatment differences, while the mean serum insulin aspart concentration profiles are derived as the mean of all individual serum insulin aspart concentrations at each time point. Faster aspart, $n = 44$; insulin aspart, $n = 46$.

FIGURE 1  Mean serum insulin aspart concentration after a bolus dose of 0.15 U/kg faster aspart or insulin aspart administered by CSII. Variability bands show the s.e.m. The full blue/grey arrows indicate that the estimated onset and offset of exposure occurred earlier for faster aspart vs insulin aspart as reflected by the endpoints time to 50% of maximum insulin aspart concentration in the early part of the pharmacokinetic profile ($t_{Early \;50\% \;C_{max}}$; estimated difference [95% CI] faster aspart vs insulin aspart $−11.8$ minutes [−14.4; −9.2]; $P < .001$) and time to 50% of maximum insulin aspart concentration in the late part of the pharmacokinetic profile ($t_{Late \;50\% \;C_{max}}$; −35.4 minutes [−47.0; −23.8]; $P < .001$). Moreover, as indicated by the dashed arrow, a left shift of the time of maximum insulin aspart concentration was also observed for faster aspart vs insulin aspart ($t_{max}$; −25.7 minutes [−34.3; −17.1]; $P < .001$). For graphical reasons, there are some discrepancies between the length of the arrows and the actual estimated mean treatment differences. This is attributable to the fact that the estimated mean treatment differences are derived from all participants’ individual treatment differences, while the mean serum insulin aspart concentration profiles are derived as the mean of all individual serum insulin aspart concentrations at each time point. Faster aspart, $n = 44$; insulin aspart, $n = 46$.

2.5 | Statistical analysis

Statistical analyses of pharmacokinetic and pharmacodynamic endpoints were conducted using SAS version 9.3 (SAS Institute, Cary, North Carolina) based on all randomized participants receiving at least 1 dose of trial product. Safety endpoints were summarized using descriptive statistics based on all participants receiving at least 1 dose of trial product.

To ensure sufficient power to evaluate treatment differences, both for pharmacokinetic and pharmacodynamic endpoints, the sample size calculation was performed not only for the primary pharmacokinetic endpoint, $\text{AUC}_{\text{IAsp,0-15min}}$, but also for the secondary pharmacodynamic endpoint, $\text{AUC}_{\text{GIR,0-1h}}$. The number of completers required was 31 in order to obtain 80% power for detecting a
geometric mean treatment ratio of 1.5 for AUC\textsubscript{\text{IAsp,0-30min}}. This was based on an assumed within-subject standard deviation (on log-scale) of 0.54 (derived from a previous trial with faster aspart\textsuperscript{11} and taking into account that the baseline correction of AUCs in this trial contributed further to the variation) and a significance level of 5%; however, in order to also obtain sufficient power for pharmacodynamic endpoints, the number of completers required in this trial was set to 44 participants. This yielded 80% power for detecting a geometric mean treatment ratio of 1.3 for AUC\textsubscript{\text{GIR,0-1h}}, based on an assumed within-subject standard deviation (on log-scale) of 0.3 and a significance level of 5%.\textsuperscript{11}

Pharmacokinetic and pharmacodynamic endpoints were analysed by means of a linear mixed model, with treatment and period as fixed effects and participant as random effect. For the AUCs, a multiplicative model was used, while for \( t_{\text{Early 50% Cmax}} \), \( t_{\text{max}} \), \( t_{\text{Late 50% Cmax}} \), \( t_{\text{Early 50% GIRmax}} \), \( t_{\text{GIRmax}} \) and \( t_{\text{Late 50% GIRmax}} \) an additive model was used. For endpoints analysed using an additive model, treatment ratios and 95% confidence intervals (CIs) were calculated using Fieller’s method.\textsuperscript{16}

3 | RESULTS

3.1 | Participant disposition and baseline characteristics

A total of 58 individuals were screened, 48 were randomized and treated with trial product, and 46 completed the trial. Two participants were withdrawn (1 as a result of an adverse event of vomiting after insulin aspart treatment and 1 because of a technical problem with a catheter). Participant disposition is presented in Figure S3, Appendix S1.

The mean \((±\text{ standard deviation [s.d.]}\) age of the participants was 46.3 \((±8.6)\) years. The majority of participants were men (66.7%), and all participants were white. The mean body weight was 76.5 \((±11.8)\) kg, mean BMI was 24.5 \((±2.3)\) kg/m\(^2\), mean duration of diabetes was 24.1 \((±12.2)\) years, and mean HbA\(_1c\) at baseline was 7.4\% \((±0.6)\). At entry into the study, 34 participants were receiving multiple daily injection therapy and 14 participants were using CSII.

3.2 | Onset of exposure and glucose-lowering effect

After a bolus dose administered by CSII, both the serum insulin concentration–time profile (Figure 1) and the glucose-lowering effect profile (Figure 2) were shifted to the left for faster aspart vs insulin aspart. With faster aspart, \( t_{\text{Early 50% Cmax}} \) was 11.8 minutes earlier, and \( t_{\text{max}} \) was 25.7 minutes earlier, compared with insulin aspart (Table 1 and Figure 1). Likewise, \( t_{\text{Early 50% GIRmax}} \) was 11.1 minutes earlier, and \( t_{\text{GIRmax}} \) was 18.7 minutes earlier for faster aspart vs insulin aspart (Table 2).

3.3 | Early exposure and glucose-lowering effect

Early insulin exposure and glucose-lowering effect were both greater for faster aspart than for insulin aspart, as shown by the significantly greater partial AUCs and GIR AUCs for faster aspart vs insulin aspart within the first 2 hours of the bolus insulin dose (Figure 3). Early exposure during the first 15 minutes after the bolus dose (AUC\textsubscript{\text{IAsp,0-15min}}) was significantly greater for faster aspart vs insulin aspart (least squares means of 12.5 and 1.8 pmol*h/L, respectively; \( P < .001 \)). Within the first 30 minutes after the bolus dose, a ~3-fold higher insulin exposure (AUC\textsubscript{\text{IAsp,0-30min}}, primary endpoint) and a ~2-fold greater glucose-lowering effect (AUC\textsubscript{\text{GIR,0-30min}}) were seen with faster aspart than with insulin aspart (Figure 3).

3.4 | Offset of exposure and glucose-lowering effect

The offset of insulin exposure after a bolus dose administered by CSII occurred earlier for faster aspart than for insulin aspart, as shown by a 35.4 minutes earlier \( t_{\text{Late 50% Cmax}} \) for faster aspart vs insulin aspart (Table 1 and Figure 1). Likewise, the offset of glucose-lowering effect occurred faster for faster aspart than for insulin aspart, as shown by a 24.0 minutes earlier \( t_{\text{Late 50% GIRmax}} \) for faster aspart vs insulin aspart (Table 2).

3.5 | Overall exposure and glucose-lowering effect

The total insulin exposure (AUC\textsubscript{\text{IAsp,0-1h}}) was similar for faster aspart and insulin aspart, while the \( C_{\text{max}} \) was slightly greater for faster aspart than for insulin aspart (Table 1). The total glucose-lowering effect (AUC\textsubscript{\text{GIR,0-1h}}) and the \( GIR_{\text{max}} \) were both similar for faster aspart and insulin aspart (Table 2).

3.6 | Safety

Both faster aspart and insulin aspart were well tolerated, and no safety issues were identified during the trial. No infusion site reactions were reported when administering faster aspart or insulin aspart by CSII. No confirmed hypoglycaemic episodes occurred during the trial.
suppression of hepatic glucose production, which appears to play a major role in controlling postprandial glucose. 

The key findings of the present study, which is the first to investigate the pharmacokinetic and pharmacodynamic properties of faster aspart in a pump setting, were that in individuals with T1DM, faster aspart provided faster onset, greater early exposure and early glucose-lowering effect compared with insulin aspart. In addition, faster aspart was well tolerated. Faster aspart may therefore have the potential to provide improved postprandial glycaemic control over current rapid-acting insulins in a pump setting. Indeed, in a double-blind, randomized, crossover study in 43 individuals with T1DM using CSII for 14 days, faster aspart provided a 25% greater plasma glucose-lowering effect during the first 2 hours of a standardized meal test compared with insulin aspart. Furthermore, the mean postprandial increment in interstitial glucose across all meals over 14 days, measured by blinded continuous glucose monitoring in the outpatient setting, was ~50% lower for faster aspart than for insulin aspart, both at 60 and 120 minutes post-meal. Reduction of postprandial glucose excursions may play an important role in improving overall glycaemic control, especially at lower HbA₁c levels as suggested by some studies.

Although a prominent contribution of postprandial glucose to HbA₁c levels was not confirmed by a later randomized controlled trial, all studies show that postprandial glucose has some impact on HbA₁c. This was also confirmed by an improvement in HbA₁c of 0.15% with faster aspart compared with insulin aspart in individuals with T1DM using multiple daily injection therapy. It remains to be seen if further improvements can be achieved with faster aspart in a CSII setting as the current pharmacokinetic and pharmacodynamic data suggest.

In healthy individuals, early endogenous insulin secretion in response to a meal induces a prompt suppression of hepatic glucose production, which appears to play a major role in controlling postprandial glucose levels. Accordingly, the concept of CSII in patients with diabetes relies on immediate action of the mealtime insulin administered. Current rapid-acting insulin analogues have shown some improvements over RHI in terms of accelerated insulin absorption. There is, however, still room for further improvement in mealtime insulin absorption rate. The results of the present trial, as well as those of previous trials, suggest that faster aspart may better reproduce the physiological insulin action profile seen in response to a meal in healthy individuals.

### TABLE 1

| Onset of exposure, min | Faster aspart | Insulin aspart | Treatment ratio (95% CI) | Treatment difference (95% CI) | P  |
|-----------------------|--------------|---------------|------------------------|-------------------------------|----|
| tEarly 50% Cmax       | 20.7         | 32.5          | 0.64 (0.57; 0.71)      | -11.8 (-14.4; -9.2)           | <.001|
| tmax                  | 56.6         | 82.3          | 0.69 (0.60; 0.78)      | -25.7 (-34.3; -17.1)          | <.001|
| Offset of exposure, min |             |               |                       |                               |    |
| tLate 50% Cmax        | 137.4        | 172.9         | 0.80 (0.73; 0.86)      | -35.4 (-47.0; -23.8)          | <.001|
| Overall exposure      |              |               |                       |                               |    |
| AUCGIR₀-t, mg/kg/min  | 606.2        | 622.8         | 0.97 (0.90; 1.05)      | NA                            | .477|
| Cmaxₜₐₚₜ, pmol/L       | 278.9        | 252.1         | 1.11 (1.03; 1.19)      | NA                            | .010|

NA, not applicable; t, time of first non-positive baseline infusion corrected insulin aspart concentration in the terminal part of the profile (however, no longer than 12 hours).

1 Data are least squares means.
2 Faster aspart/insulin aspart (for onset and offset of exposure endpoints, the treatment ratio was calculated using Fieller’s method).
3 Faster aspart – insulin aspart.

### TABLE 2

| Onset of glucose-lowering effect, min | Faster aspart | Insulin aspart | Treatment ratio (95% CI) | Treatment difference (95% CI) | P  |
|--------------------------------------|--------------|---------------|------------------------|-------------------------------|----|
| tEarly 50% GIRmax                    | 41.1         | 52.3          | 0.79 (0.72; 0.86)      | -11.1 (-15.4; -6.9)           | <.001|
| tGIRmax                              | 111.9        | 130.6         | 0.86 (0.75; 0.97)      | -18.7 (-34.4; -2.9)          | .021|
| Offset of glucose-lowering effect, min |             |               |                       |                               |    |
| tLate 50% GIRmax                     | 214.7        | 238.7         | 0.90 (0.84; 0.96)      | -24.0 (-38.9; -9.1)          | .002|
| Overall glucose-lowering effect      |              |               |                       |                               |    |
| AUCGIR₀ₜₐₚₜ, mg/kg                   | 1341.5       | 1295.7        | 1.04 (0.95; 1.13)      | NA                            | .427|
| GIRₜₐₚₜₜ, mg/kg/min                  | 7.1          | 6.8           | 1.04 (0.94; 1.14)      | NA                            | .439|

NA, not applicable; t, time of first non-positive baseline infusion corrected GIR in the terminal part of the smoothed GIR profile (however, no longer than 12 hours).

1 Data are least squares means.
2 Faster aspart/insulin aspart (for onset and offset of glucose-lowering effect endpoints, the treatment ratio was calculated using Fieller’s method).
3 Faster aspart – insulin aspart.

# DISCUSSION

The key findings of the present study, which is the first to investigate the pharmacokinetic and pharmacodynamic properties of faster aspart in a pump setting, were that in individuals with T1DM, faster aspart provided faster onset, greater early exposure and early glucose-lowering effect and faster offset compared with insulin aspart. In addition, faster aspart was well tolerated. Faster aspart may therefore have the potential to provide improved postprandial glycaemic control over current rapid-acting insulins in a pump setting. Indeed, in a double-blind, randomized, crossover study in 43 individuals with T1DM using CSII for 14 days, faster aspart provided a 25% greater plasma glucose-lowering effect during the first 2 hours of a standardized meal test compared with insulin aspart. Furthermore, the mean postprandial increment in interstitial glucose across all meals over 14 days, measured by blinded continuous glucose monitoring in the outpatient setting, was ~50% lower for faster aspart than for insulin aspart, both at 60 and 120 minutes post-meal. Reduction of postprandial glucose excursions may play an important role in improving overall glycaemic control, especially at lower HbA₁c levels as suggested by some studies.

Although a prominent contribution of postprandial glucose to HbA₁c levels was not confirmed by a later randomized controlled trial, all studies show that postprandial glucose has some impact on HbA₁c. This was also confirmed by an improvement in HbA₁c of 0.15% with faster aspart compared with insulin aspart in individuals with T1DM using multiple daily injection therapy. It remains to be seen if further improvements can be achieved with faster aspart in a CSII setting as the current pharmacokinetic and pharmacodynamic data suggest.

In healthy individuals, early endogenous insulin secretion in response to a meal induces a prompt suppression of hepatic glucose production, which appears to play a major role in controlling postprandial glucose levels. Accordingly, the concept of CSII in patients with diabetes relies on immediate action of the mealtime insulin administered. Current rapid-acting insulin analogues have shown some improvements over RHI in terms of accelerated insulin absorption. There is, however, still room for further improvement in mealtime insulin absorption rate. The results of the present trial, as well as those of previous trials, suggest that faster aspart may better reproduce the physiological insulin action profile seen in response to a meal in healthy individuals.
Another aim when optimizing insulin for use in CSII has been to increase the rate of offset, in order to reduce the risk of late postprandial hypoglycaemia. The duration of mealtime insulin requirement depends on several individual factors, such as food composition and rate of gastric emptying, and there is a limit to the optimum rate of offset because a very fast offset of action could imply insufficient insulin action in the late postprandial period. It was shown in the present study that faster aspart had a faster offset of exposure and action compared with insulin aspart. In standardized meal tests following faster aspart or insulin aspart administration by subcutaneous injection or CSII, plasma glucose for faster aspart was consistently lower than or equal to insulin aspart up to 4 to 6 hours postmeal. This indicates that the rate of offset is not too rapid with faster aspart. Specifically designed studies are needed to further investigate the relationship between rate of offset and the risk of late postprandial hypoglycaemia with faster aspart.

When comparing the present results for faster aspart using CSII with those previously reported for subcutaneous injection, it appears that improvements in onset as well as early exposure and action with faster aspart vs insulin aspart are most pronounced in the pump setting. Treatment ratios of faster aspart vs insulin aspart are most pronounced in the present study as compared with 2.05 and 1.48 after subcutaneous injection or CSII, plasma glucose for faster aspart was consistently lower than or equal to insulin aspart up to 4 to 6 hours postmeal. This indicates that the rate of offset is not too rapid with faster aspart. Specifically designed studies are needed to further investigate the relationship between rate of offset and the risk of late postprandial hypoglycaemia with faster aspart.

When comparing the present results for faster aspart using CSII with those previously reported for subcutaneous injection, it appears that improvements in onset as well as early exposure and action with faster aspart vs insulin aspart are most pronounced in the pump setting. Treatment ratios of faster aspart vs insulin aspart for AUC_{A_{\text{Asp},0-30\text{min}}} and AUC_{GIR,0-30\text{min}} were 2.95 and 2.18 in the present study as compared with 2.05 and 1.48 after subcutaneous injection. It should be noted that this is a comparison between two different studies and ideally a dedicated study needs to be performed to reach firm conclusions regarding the pharmacokinetic and pharmacodynamic properties of faster aspart relative to insulin aspart after subcutaneous injection vs CSII. One of the excipients in faster aspart, niacinamide, promotes the formation of insulin aspart monomers in diluted formulations mimicking subcutaneous conditions after administration, and augments the permeation rate of insulin aspart in human dermal-derived microvascular capillary endothelial cell monolayers. It may be speculated that the continuous presence of niacinamide at the infusion site in the CSII setting due to the basal infusion rate and/or the duration of the bolus infusion may further facilitate the rapid absorption of faster aspart as compared with the single injection setting where dosing occurs almost instantaneously and niacinamide might disappear relatively quickly from the injection site. Another hypothesis could be that the mode of delivery of the bolus infusion when using CSII favours a better diffusion of faster aspart in the subcutaneous tissue until reaching the capillaries. These potential mechanisms are, however, solely speculative at the moment and further investigations are needed within this area.

Differences in duration of the bolus infusion between different insulin pumps have previously been shown to affect the pharmacokinetic and pharmacodynamic characteristics of mealtime insulin. In the present study, the same insulin pump with the same bolus infusion mode was used for the two treatments in a crossover design. Consequently, the duration of bolus infusion was similar for faster aspart and insulin aspart within each participant. Importantly, in the study by Bode et al., reduced postprandial glucose increment was observed with faster aspart compared with insulin aspart across several models of insulin pumps with different speed of bolus delivery.

As a result of the bolus being administered on top of a basal insulin infusion, it was not possible to assess onset of appearance and onset of action in any meaningful way using recognized methodology, which is a limitation of the present study.

**FIGURE 3** Early exposure (A) and early glucose-lowering effect (B) for faster aspart vs insulin aspart after a bolus dose of 0.15 U/kg administered by CSII. n = 44 for faster aspart and n = 45 (AUC_{GIR,0-1.5h} and AUC_{GIR,0-2h}) or 46 (all other endpoints) for insulin aspart. The treatment ratio for AUC_{GIR,0-30min} and AUC_{GIR,0-1h} was calculated using Fieller’s method. LS, least squares.

| Endpoint | Treatment ratio [95%CI] | P-value | LS Mean (pmol·h/L) | P-value |
|----------|-------------------------|---------|--------------------|---------|
| AUC_{A_{\text{Asp},0-30\text{min}}} | 2.95 [2.32;3.73] | <0.001 | 51.4 | 17.4 |
| AUC_{A_{\text{Asp},0-1h}} | 1.52 [1.37;1.69] | <0.001 | 163.0 | 107.3 |
| AUC_{A_{\text{Asp},0-1.5h}} | 1.30 [1.20;1.41] | <0.001 | 275.2 | 212.0 |
| AUC_{A_{\text{Asp},0-2h}} | 1.18 [1.10;1.26] | <0.001 | 363.0 | 306.8 |

| Endpoint | Treatment ratio [95%CI] | P-value | LS Mean (mg/kg) | P-value |
|----------|-------------------------|---------|-----------------|---------|
| AUC_{GIR,0-30min} | 2.18 [1.33;5.04] | 0.002 | 24.9 | 11.4 |
| AUC_{GIR,0-1h} | 1.52 [1.29;1.83] | <0.001 | 165.9 | 109.3 |
| AUC_{GIR,0-1.5h} | 1.34 [1.14;1.57] | 0.001 | 339.6 | 254.3 |
| AUC_{GIR,0-2h} | 1.21 [1.07;1.38] | 0.002 | 543.1 | 448.9 |
treatments, so the interpretation of the pharmacodynamic endpoints \( t_{\text{Early 50\% GIRmax}} \) and \( t_{\text{Late 50\% GIRmax}} \), which were also in favour of faster aspart, is straightforward.

A strength of the present study is that both pharmacokinetic and pharmacodynamic measures were obtained after a bolus insulin dose on top of a basal infusion, thus resembling a clinically relevant dosing scheme during CSII. Furthermore, the study included individuals with T1DM, which is considered the optimum population in glucose clamp studies, as this allows comparison of exogenous insulins with respect to glucose-lowering effect without interference from endogenous insulin. Nevertheless, the strictly controlled conditions of the glucose clamp method, eg, the long fasting period and the wash-out of current insulin, differ from clinical practice and may imply certain challenges in translating to clinical outcomes.

In conclusion, in people with T1DM using CSII, faster aspart provides faster onset and greater early exposure and glucose-lowering effect, as well as faster offset compared with insulin aspart, thereby better mimicking the endogenous prandial insulin secretion and action. Accordingly, faster aspart has been shown to provide improvements in postprandial glucose control over insulin aspart in individuals using CSII. Faster aspart used in an insulin pump setting may represent an advancement in insulin therapy towards optimum postprandial glucose control in patients with diabetes and may also have potential in advanced pumps and closed-loop systems.

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Conflict of interest

T. H.’s institution has received research grants from Adocia, AstraZeneca, Becton Dickinson, Biocron, Boehringer Ingelheim, Danex Biopharm, Eli Lilly, Grünenthal, Gulf Pharmaceutical Industries, Johnson & Johnson, Marvel, MedImmune, Medtronic, Novartis, Novo Nordisk, Roche Diagnostics, Sanofi, Senseonics and Zealand Pharma. In addition, T. H. is a member of advisory boards for Novo Nordisk and has received speakers’ honoraria and travel grants from Eli Lilly, Mylan and Novo Nordisk. E. Z. has received travel grants from Novo Nordisk and Danex Biopharm and speakers’ honoraria from Novo Nordisk and Roche Diabetes Care. T. R. and H. H. are employees and shareholders of Novo Nordisk. L. N. has no conflict of interest to declare.

Author contributions

T. H. contributed to the study design, conduct/data collection, analysis and writing of the manuscript. E. Z. contributed to the study design and writing of the manuscript. L. N. contributed to the study conduct/data collection and writing of the manuscript. T. R. contributed to the data analysis and writing of the manuscript. H. H. contributed to the study design and writing of the manuscript.

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