Optimal prandial timing of bolus insulin in diabetes management: a review

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Abstract

The inability to achieve optimal diabetes glucose control in people with diabetes is multifactorial, but one contributor may be inadequate control of postprandial glucose. In patients treated with multiple daily injections of insulin, both the dose and timing of meal-related rapid-acting insulin are key factors in this. There are conflicting opinions and evidence on the optimal time to administer mealtime insulin. We performed a comprehensive literature search to review the published data, focusing on the use of rapid-acting insulin analogues in patients with Type 1 diabetes. Pharmacokinetic and pharmacodynamic studies of rapid-acting insulin analogues, together with postprandial glucose excursion data, suggest that administering these 15–20 min before food would provide optimal postprandial glucose control. Data from clinical studies involving people with Type 1 diabetes receiving structured meals and rapid-acting insulin analogues support this, showing a reduction in post-meal glucose levels of ~30% and less hypoglycaemia when meal insulin was taken 15–20 min before a meal compared with immediately before the meal. Importantly, there was also a greater risk of postprandial hypoglycaemia when patients took rapid-acting analogues after eating compared with before eating.

Introducțion

The importance of optimal glycemic control in preventing the micro- and macrovascular complications associated with diabetes has been well documented [1,2]. Despite this, a significant percentage of people with diabetes do not achieve target glycemic control. The UK National Diabetes Audit 2015–2016 found that HbA1c levels were >58 mmol/mol (7.5%) in 70.8% of people with Type 1 diabetes and 34.3% in those with Type 2 diabetes [3]. Data published in the USA in 2013 estimated that 47.8% of people with diabetes had HbA1c levels of >53 mmol/mol (7%) [4]. The inability to achieve optimal glycemic control in diabetes is multifaceted, as highlighted by Khunti et al. [5]; however, postprandial hyperglycaemia is one likely key contributing factor [6]. High postprandial blood glucose (BG) levels also contribute to greater glycemic variability, another marker of poor glycemic control [7]. Epidemiological studies show an association between impaired glucose tolerance and cardiovascular risk and outcome [8].

Prandial insulin replacement is important. In individuals without diabetes, prandial insulin makes up ~50% of the total daily pancreatic output. Most of the prandial insulin is secreted within the first hour after the meal [9]. The International Diabetes Federation consensus statement recommends that 2-h post-meal glucose levels should not exceed 7.8 mmol/l, as this level is seldom seen in those without diabetes [10]. The American Diabetes Association specifies a postprandial glucose target of 10 mmol/l at 2 h [11]. There is evidence that postprandial glucose excursions beyond these levels increase the risk of retinopathy [12] and greater carotid intima-media thickness, and lead to oxidative stress, inflammation and endothelial dysfunction [13–15]. Furthermore, there is evidence to suggest that post-meal hyperglycaemia is also associated with decreased myocardial blood flow and an increased risk of cancer [16,17]. Meanwhile, the pre-meal target in the intensive arm of the Diabetes Control and Complications Trial was 3.9–6.7 mmol/l and resulted in a significant reduction in vascular complications [18].

In people with either Type 1 or insulin-requiring Type 2 diabetes treated with multiple daily injections (MDI), short- or rapid-acting insulin is given with meals to cover mealtime glucose excursions. The pharmacology of the insulin compared with the glucose profile from the food
ingested govern the extent of post-meal glucose excursions. In the pre-rapid-acting insulin analogue era, regular human insulin (RHI) was the mainstay of bolus insulin therapy; however, recognition of its slow onset of action and delayed peak led to the recommendation to take it ≥30 min pre-meal. In practice, many people did not do this. To address this, rapid-acting insulin analogues were introduced in the 1990s.

There are currently three rapid-acting analogues marketed in the USA and Europe: insulin lispro (Humalog; Eli Lilly, Indianapolis, IN, USA), insulin aspart (Novolog/NovoRapid; Novo Nordisk, Bagsvaerd, Denmark) and insulin glulisine (Apidra; Sanofi, Paris, France). Additionally, a fourth rapid-acting analogue, fast-acting insulin aspart (faster aspart; Fiasp; Novo Nordisk, Bagsvaerd, Denmark), has recently been approved for marketing in Europe and other parts of the world. For insulin lispro, proline and lysine at positions 28 and 29 on the B-chain of human insulin are reversed. With insulin aspart, proline at position 28 on the B-chain of human insulin is replaced with aspartic acid and for insulin glulisine, arginine at position 3 on the B-chain is replaced with lysine, and lysine at position 29 on the B-chain is replaced with glutamic acid. These changes reduce the ability of the insulin molecules to aggregate, and the dimers and monomers are more rapidly absorbed after subcutaneous (s.c.) injection. Next-generation and faster rapid-acting insulin analogues have also been developed that boast superior insulin absorption rates and early glucose-lowering effects when compared with rapid-acting analogues. Faster aspart is insulin aspart set in a new formulation with vitamin B3 (also known as nicotinamide) and arginine.

Manufacturers of rapid-acting insulin analogues recommend injecting immediately before food or soon thereafter, and this is common practice for many people who feel more confident of the amount of carbohydrate eaten after they have eaten it [19–21]. Most structured education programmes recommend injections pre-meal, but often the precise timing is not specified and, in clinical practice, we observe many patients injecting their mealtime insulin post-meal. For this reason, we conducted a systematic literature review of studies evaluating the timing of rapid-acting insulin in an attempt to obtain some clarity on this important topic.

Methods

Data for the present review were collected through searches of PubMed: a specific search over the past 30 years and a more general search over the past 10 years. A search of ProQuest was also conducted which captured the Embase and Biosis databases. Search terms included: ‘diabetes’, ‘diabetes mellitus’, ‘Type 2 diabetes’, ‘Type 1 diabetes’, ‘T1D’, ‘T2D’, ‘bolus insulin’, ‘prandial insulin’, ‘insulin analogue’, ‘insulin aspart’, ‘insulin lispro’, ‘insulin glulisine’, ‘postprandial excursions’, ‘postprandial hyperglycaemia’, ‘postprandial administration’, ‘preprandial administration’, ‘post-meal administration’, ‘pre-meal administration’, ‘insulin timing’, ‘time of dose’, ‘timing of bolus’, ‘timing of prandial’, ‘dosing’, ‘flexibility’, ‘pharmacokinetics’ and ‘pharmacodynamics’. Studies on faster aspart were identified by the original search. A study on BC lispro was reviewed and included after the published search. The specific search of PubMed over the past 30 years yielded 1432 results, and the more general 10-year search yielded 1990 results. The ProQuest search yielded 770 results. The authors reviewed the abstracts of all papers produced by the search and evaluated for relevance papers that specifically looked at the glycaemic effect of timing of rapid-acting insulin in Type 1 or Type 2 diabetes. We identified 19 studies that could potentially be relevant to our review, with 11 being included (Fig. 1). Additional studies included in the review were obtained from references of studies identified by the search.

Evidence from pharmacokinetic and pharmacodynamic studies

Pharmacokinetic (PK) studies carried out in people with Type 1 diabetes show that all three rapid-acting insulin
analogue 46.7% and 35% respectively compared with insulin aspart and insulin lispro, the PK properties of insulin glulisine differ slightly in the majority of published studies, with a faster onset of action observed for glulisine. They demonstrate peak plasma analogues have similar PK and pharmacodynamic (PD) profiles (Table 1) [22,23]. They demonstrate peak plasma insulin concentrations approximately double those of RHI, and a time to maximum concentration less than half that of RHI, with concentrations of the analogues falling more rapidly, returning to levels <20% of peak concentrations at about 4 h (Fig. 2) [19–21]. One study showed little difference between the analogues: insulin aspart reached t50% of peak(ins) at 19.6 ± 1.7 min and insulin lispro at 16.7 ± 1.8 min (P = 0.29), and each analogue reached tpeak(ins) at 43.8 ± 3.9 min and 46.7 ± 4.7 min, respectively (P = 0.66) [22]. When compared with insulin aspart and insulin lispro, the PK properties of insulin glulisine differ slightly in the majority of published studies, with a faster onset of action observed for glulisine. Heise et al. [23] showed that the time to 10% of total insulin area under the curve (INS-AUC) was faster with insulin glulisine compared with insulin lispro at either dose (0.2 U/kg: 0.7 ± 0.2 vs 0.8 ± 0.2 h; 0.4 U/kg: 0.8 ± 0.2 vs 0.9 ± 0.2 h; P < 0.001) [23]. When compared with insulin aspart, faster absorption rates were noted with insulin glulisine (shorter times to 10% and 20% of INS (max); P = 0.0005 each) [24].

Heise et al. [25] also investigated the PK properties of faster aspart, comparing it with insulin aspart. A faster initial onset of absorption of faster aspart vs insulin aspart was supported by a significantly earlier onset of appearance (4.9 vs 11.2 min) and time to reach half the maximum concentration (t50%Cmax(ins) 20.7 vs 31.6 min). With faster aspart, the time to onset of appearance and t50%Cmax(ins) were reduced by 57% and 35%, respectively, compared with insulin aspart. The tmax(ins) for faster aspart was 62.9 min and, for insulin aspart, it was 69.7 min (Fig. 2) [25].

Andersen et al. [26] looked at the PK properties of ultra-rapid BC lispro vs insulin lispro (0.4 U/kg vs 0.2 U/kg). Onset of action was significantly earlier with BC lispro when compared with insulin lispro, with a median (min;max) t50% Cmax(ins) of 15 (6;32) vs 27 (12;43) min, respectively. The median (min;max) tmax(ins) for BC lispro was 45 (25;120) min and for insulin lispro it was 60 (25;105) min (Fig. 2) [26].

In clinical practice, PD assessment holds more relevance than PK assessment. The glucose-clamp technique is the ‘gold standard’ for assessing insulin PD characteristics, providing data on onset, peak and duration of action, which are key in determining optimal bolus timing [27]. Euglycaemia is maintained after insulin administration via a concomitant intravenous glucose infusion at a variable rate. This variable glucose infusion rate (GIR) is indicative of whole-body insulin action. Clamp studies have demonstrated discordance between rapid-acting insulin analogue PK properties (absorption time) and PD properties (action time), which creates an obstacle in successfully replicating prandial, physiological insulin action (we have not found any clinical data that explain the differences seen between the PK and PD characteristics of these insulin analogues). For example, a study comparing PD profiles of insulin lispro 100 U/ml and insulin lispro 200 U/ml found that, although the times to maximum insulin concentration were 45 and 60 min, respectively, the times to maximum GIR were similar (120 vs 126.6 min for insulin lispro 100 vs 200 U/ml) [28]. Another study highlights the potential difference in timing of important outcomes. Directly comparing insulin aspart with insulin lispro, this study demonstrated a maximum insulin concentration of 30 min for both (P = 0.24), but a maximum GIR at 120 min, also for both (P = 0.61; Fig. 3) [29]. Swan et al. [30] investigated the effect of puberty on the PD and PK properties.

### Table 1 Pharmacokinetic and pharmacodynamic studies

<table>
<thead>
<tr>
<th>Study</th>
<th>Rapid-acting insulin analogues</th>
<th>PK characteristics</th>
<th>PD characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>INS-Tmax, min</td>
<td>GIR-Tmax, min</td>
</tr>
<tr>
<td>Plank et al. [22]</td>
<td>Insulin aspart</td>
<td>43.8 ± 3.9</td>
<td>n/a</td>
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<tr>
<td></td>
<td>Insulin lispro</td>
<td>46.7 ± 4.7</td>
<td></td>
</tr>
<tr>
<td></td>
<td>t50%Cmax(ins) 20.7 vs 31.6 min</td>
<td>196 ± 73</td>
<td>198 ± 65</td>
</tr>
<tr>
<td>Heise et al. [23]</td>
<td>Insulin glulisine</td>
<td>90 (40–120)</td>
<td>186 (155–263)</td>
</tr>
<tr>
<td></td>
<td>Insulin aspart</td>
<td>90 (50–150)</td>
<td>156 (83–245)</td>
</tr>
<tr>
<td></td>
<td>t50%Cmax(ins) 92 ± 38</td>
<td>132.5 (5.87)</td>
<td></td>
</tr>
<tr>
<td>Arnold et al. [24]*</td>
<td>Insulin aspart</td>
<td>62.9 (3.73)</td>
<td>124.3 (5.87)</td>
</tr>
<tr>
<td></td>
<td>Insulin aspart</td>
<td>69.7 (3.73)</td>
<td>132.5 (5.87)</td>
</tr>
<tr>
<td></td>
<td>t50%Cmax(ins) 45 (30–180)</td>
<td>120 (56)</td>
<td></td>
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<tr>
<td></td>
<td>Lispro 100</td>
<td>45 (30–180)</td>
<td></td>
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<tr>
<td></td>
<td>t50%Cmax(ins) 60 (30–180)</td>
<td>126.6 (49)</td>
<td></td>
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<tr>
<td></td>
<td>Lispro 200</td>
<td>60 (30–180)</td>
<td></td>
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<tr>
<td></td>
<td>tmax(ins) 62.9</td>
<td>120</td>
<td></td>
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<tr>
<td></td>
<td>Aspart 30</td>
<td>120</td>
<td></td>
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<tr>
<td></td>
<td>Insulin lispro</td>
<td>30</td>
<td></td>
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<td></td>
<td>tmax(ins) 45 (25–120)</td>
<td>109 (65–221)</td>
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<tr>
<td></td>
<td>Insulin aspart</td>
<td>60</td>
<td></td>
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<td></td>
<td>tmax(ins) 60 (25–105)</td>
<td>117 (71–225)</td>
<td></td>
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<tr>
<td></td>
<td>BC lispro</td>
<td>60</td>
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<tr>
<td></td>
<td>tmax(ins) 45 (25–120)</td>
<td>109 (65–221)</td>
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<td></td>
<td>Insulin lispro</td>
<td>60</td>
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<tr>
<td></td>
<td>tmax(ins) 60 (25–105)</td>
<td>117 (71–225)</td>
<td></td>
</tr>
</tbody>
</table>

Values are means (SD) unless stated otherwise. *Values are medians (range). †Values are means (SEM). ‡Values are means [CV[%]]. BC lispro, BioChaparone insulin lispro; CV[%], coefficient of variance; GIR-Tmax, time to maximum glucose infusion rate; INS-Tmax, time to maximum serum insulin concentration; n/a, not available; PD, pharmacodynamic; PK, pharmacokinetic.
of insulin pump therapy in adolescents, and found that the peak action of insulin aspart was not observed until 90 min, 40 min after peak insulin concentration was reached.

The PD characteristics of faster aspart and BC lispro may be superior when compared with their respective rapid-acting analogue counterparts. When comparing faster aspart with...
insulin aspart, the onset of glucose-lowering effect was earlier with faster aspart, with a significantly earlier t50% GIRmax (31 vs 42 min, respectively; P < 0.001) and significantly earlier tGIRmax (109 vs 117 min, respectively; P = 0.0005) [26].

Continuous glucose monitoring (CGM) studies have shown that postprandial glucose levels peak at a mean of 70–80 min after eating in people with diabetes [31]. CGM measures interstitial glucose, with a lag of 4–10 min in relation to BG levels [32]. Although peak insulin levels are seen 40–60 min post-injection, peak insulin action occurs around 100–120 min after injection. Given this, it is reasonable to expect that the optimal time to administer rapid-acting insulin analogues is 15–20 min prior to eating, to synchronize insulin action peaks with postprandial glucose excursions, thus minimizing postprandial hyperglycaemia.

**Evidence from clinical studies**

Conflicting literature exists on optimal prandial bolus timing in clinical practice. Two studies in particular favour injection of prandial insulin 15–20 min before eating. Cobry et al. [33] carried out a crossover study in 23 young people with Type 1 diabetes (mean age 18.3 ± 4.4 years) on insulin pump therapy. The trial had three treatment arms: delivering an insulin glulisine bolus by insulin pump 20 min prior to a meal (−20 min), immediately before the meal (0 min) or 20 min after meal initiation (+20 min). At 60 min, the −20 min arm showed significantly lower glycaemic excursions than both the 0 min arm and the +20 min arm (−20 min = 10.0 ± 3.70 mmol/l vs 12.33 ± 3.27 mmol/l and 13.1 ± 2.59 mmol/l, respectively). At 120 min after meal initiation, the −20 min arm likewise showed significantly lower BG values than both the 0 min and +20 min arms (−20 min = 9.79 ± 3.9 mmol/l vs 11.5 ± 2.7 mmol/l and 11.4 ± 2.8 mmol/l, respectively; Table 2). Peak BG levels were also significantly lower in the −20 min arm compared with the 0 min arm and in the +20 min arm (−20 min = 11.2 ± 0.44 mmol/l; P = 0.0001) and the +20 min arm (13.7 ± 0.47 mmol/l; P < 0.0001; Fig. 4). No difference in BG readings was observed when insulin was administered immediately prior to the meal compared with 20 min post-meal. Hypoglycaemic episodes recorded were highest in the +20 min arm compared with the 0 min and −20 min arms (five vs one vs four), respectively [33].

Luijf et al. [34] studied 10 people with Type 1 diabetes on insulin pump therapy with a mean age of 45.5 ± 12.1 years, in a three-way, randomized, crossover trial. Insulin aspart was administered at 30, 15 or 0 min before mealtime. Each participant was provided with a breakfast similar to their usual breakfast. Area under the glucose curve was lower in the −15 min arm (0.41 ± 0.51 mmol/l/min) than in the −30 min arm (1.89 ± 0.72 mmol/l/min; P = 0.029) and 0 min arm (2.11 ± 0.66 mmol/l/min; P = 0.030). Maximum glucose excursion was almost 30% lower in the −15 min arm.
<20 g of fat. The protein content was not revealed. In the latter study, the nutritional content of the meal was not mentioned, which is an important missing variable. The infusion sites of the pumps were also not mentioned.

Several studies have compared prandial (immediately before eating) and postprandial administration of rapid-acting insulin analogues (Table 2). Brunner et al. [35] compared insulin aspart administered immediately before (0 min) and 15 min after the start of the meal, along with RHI 15 min before and immediately before the meal [35]. This was a well-designed study in which participants’ glucose levels were kept within a range of 100.8–140.4 mg/dl (5.6–7.8 mmol/l) prior to commencement, with a variable insulin infusion. A standardized breakfast was used (543 kcal, 55% carbohydrate, 35 g of protein and 28 g of fat). That study showed that insulin aspart at 0 min was superior to insulin aspart at +15 min and was similar to RHI at −15 min. The lowest postprandial glucose level achieved was in insulin aspart at 0 min but was higher than most target values at 11.2 mmol/l, compared with 13.2 mmol/l with insulin aspart at +15 min. Insulin aspart injected 15 min before mealtime was not investigated in that study. Importantly, late hypoglycaemia occurred in 21% of the experiments (0 min, n = 6; +15 min, n = 6) [35].

One of the most comprehensive studies was performed by Schernthaner et al. [36] comparing RHI at −40, −20 and 0 min and insulin lispro at −20, 0 and +15 min on postprandial glucose levels. Participants in that study had a standardized meal consisting of 584.5 kcal, 45.5 g of carbohydrate, 35 g of protein and 28 g of fat. BG excursions at 60 min after injection were significantly lower with insulin lispro at −20 min when compared with all other treatments, particularly insulin lispro 0 min and +15 min (−1.12 ± 2.13 vs 0.19 ± 1.72 vs 2.20 ± 1.49 mmol/l, respectively). At 90 and 120 min, insulin lispro −20 min and 0 min were superior to all other treatments, with insulin lispro 0 min performing much better at 90 and 120 min than at 60 min when compared with insulin lispro −20 min (−1.44 ± 1.60

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**Table 2 Clinical studies**

<table>
<thead>
<tr>
<th>Study</th>
<th>Rapid-acting insulin analogue</th>
<th>CSII or MDI</th>
<th>Time of insulin administration in relation to mealtime, min</th>
<th>Most effective time at lowering postprandial hyperglycaemia, min</th>
<th>Postprandial glucose levels, mmol/l</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cobry et al. [33]</td>
<td>Insulin glulisine</td>
<td>CSII</td>
<td>−20, 0, +20*</td>
<td>−20</td>
<td>11.0 ± 3.8 vs 13.7 ± 3.0 vs 13.8 ± 2.3 (max)</td>
</tr>
<tr>
<td>Luijé et al. [34]</td>
<td>Insulin aspart</td>
<td>CSII</td>
<td>−30, −15, 0</td>
<td>−15</td>
<td>11.74 ± 0.8 vs 9.26 ± 0.72 vs 12.29 ± 0.93 (max)</td>
</tr>
<tr>
<td>Brunner et al. [35]</td>
<td>Insulin aspart</td>
<td>MDI</td>
<td>0, +15</td>
<td>0</td>
<td>11.2 (10.4–12.0) vs 13.2 (12.3–14.2) (max)</td>
</tr>
<tr>
<td>Schernthaner et al. [36]</td>
<td>Insulin lispro</td>
<td>MDI</td>
<td>−20, 0, +15*</td>
<td>−20</td>
<td>n/a</td>
</tr>
<tr>
<td>Jovanovic et al. [38]</td>
<td>Insulin aspart</td>
<td>MDI</td>
<td>−5 to 0, +30*</td>
<td>−5 to 0</td>
<td>5.7 ± 0.5 vs 8.3 ± 0.55 (max)</td>
</tr>
<tr>
<td>Schernthaner et al. [37]</td>
<td>Insulin lispro</td>
<td>MDI</td>
<td>0, +30</td>
<td>0</td>
<td>7.71 ± 1.83 vs 8.66 ± 2.13 (mean)</td>
</tr>
</tbody>
</table>

*Up to 15, 20 and 30 min after commencement of eating. CSII, continuous subcutaneous insulin infusion; max, maximum postprandial glucose level; mean, mean postprandial glucose level; MDI, multiple daily injections; n/a, not available.

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(4.77 ± 0.52 mmol/l) than in the −30 min arm (6.48 ± 0.76 mmol/l; P = 0.025) and 0 min arm (6.93 ± 0.76 mmol/l; P = 0.022). Time spent in the +3.5 to +10 mmol/l range was higher in the −15 min arm (224.5 ± 25.0 min) than in the 0 min arm (90.5 ± 23.2 min; P = 0.001). There was no significant difference in occurrence of hypoglycaemia between arms (P = 0.901) [34].

While both these studies were performed exclusively in people using insulin pumps, the results should be applicable to people using MDI regimens, as the bolus aspect of these therapies is very similar. Importantly, participants in both of these studies had quite tight glucose control immediately prior to commencement of the study: 5.5–10 mmol/l in the Cobry et al. [33] study and 3.5–7.8 mmol/l in the Luijé et al. study [34]. In the former, test meals consisted of a known, fixed amount of carbohydrate that was not specified and
Dose timing may be less critical in people with Type 2 diabetes, at least while they retain useful amounts of endogenous insulin. A study by Gredal et al. [42] assessed the optimal dose and timing of aspart in people with Type 2 diabetes. No difference in postprandial glucose profile was demonstrated whether insulin aspart 0.04 IU/kg was administered 15 or 30 min before mealtime. Doubling the dose increased the risk of hypoglycaemia [42].

Ratner et al. [43] investigated the effect of insulin glulisine injected either preprandially (0–15 min) or postprandially (+20 min) on glycaemic control and weight gain in people with Type 2 diabetes. Participants were also taking insulin glargine once daily ± metformin. This study lasted for 52 weeks, with 322 people completing the study. At study end, insulin glulisine achieved similar glycaemic control whether it was administered before or after meals (HbA1c: 7.04% pre-meal vs 7.16% post-meal; P < non-significant). Overall hypoglycaemia incidence and severe hypoglycaemia rates were not significantly different between pre-meal and post-meal groups; however, symptomatic and nocturnal hypoglycaemia rates were higher in the postprandial group. There was no significant difference in weight gain [43].

Many people with Type 2 diabetes requiring insulin therapy use biphasic or mixed insulin. Warren et al. [44] compared biphasic aspart insulin (BIAsp 30, a biphasic formulation of insulin aspart, 30% soluble and 70% protamine-crystallized) injected 5 min before or 15–20 min
after eating in an elderly population (aged >65 years) with Type 2 diabetes. Mean plasma glucose values during a 4-h meal test at the end of each treatment were similar for pre- and postprandial BAAsp 30 (8.5 ± 3.2 mmol/l and 8.9 ± 3.3 mmol/l, respectively; difference not significant). The mean BG increment from self-measured BG values, however, was slightly but significantly greater after postprandial injection than after preprandial injection (treatment difference: 0.9 mmol/l, 95% CI 0.03;1.63). No increased risk of hypoglycaemia was seen with postprandial injection [44].

**Effect of other factors on postprandial glucose control**

When interpreting data from the described studies, it is important to consider other factors that can adversely affect postprandial glycaemia and potentially skew results.

The nutritional content of food, particularly the protein and fat content, has been shown to affect postprandial glycaemia. Some studies have shown that meals containing carbohydrates that are high in dietary fat cause sustained late postprandial hyperglycaemia. One study showed that the addition of 35 g dietary fat increased postprandial glucose concentrations by 2.3 mmol/l at 5 h and another demonstrated that the addition of 50 g fat caused significant hyperglycaemia over 5 h [45,46]. Protein has also been shown to increase postprandial glucose levels, with one study reporting that the addition of 35 g of protein to a 30-g carbohydrate meal resulted in a 2.6-mmol/l increase in BG at 5 h [47]. There are also data to suggest that food order has a significant role to play. A study by Shukla et al. [48] showed that when protein and fat were consumed 15 min before carbohydrate, the mean post-meal glucose levels were lower by 28.6% (P = 0.001), 36.7% (P = 0.001) and 16.8% (P = 0.03) at 30, 60 and 120 min, respectively, and the incremental AUC was 73% lower. The glycaemic index of food may also affect postprandial BG levels, as foods with a high glycaemic index cause a large and rapid rise in BG, whereas those with a low glycaemic index produce small fluctuations in BG [49]. It has also been shown that large carbohydrate meals may contribute to late postprandial hyperglycaemia [50]. These studies highlight the importance of knowing and understanding the nutritional content of meals, as this can have a bearing on prandial glucose levels and insulin requirements. We may speculate that bolusing 15–20 min before eating is of most importance with high-glycaemic index foods, and that dividing administration of doses may allow optimum postprandial glucose control for meals of high fat and/or protein content. One study found that an 8-h dual-wave bolus given pre-meal using an insulin pump provided the best postprandial glucose control after a high-fat meal [51].

Gastric emptying rate is also an important variable that can influence postprandial glycaemia in both people with and without diabetes, with significant inter-individual variability. Pre-meal glucose affects gastric emptying, with hyperglycaemia causing a ‘physiological’ slowing, as may meal composition and other concomitant medication such as glucagon-like peptide-1 receptor agonists [52–54]. People with gastroparesis may also need a different bolus profile, such as a dual-wave or a square-wave, to mimic the delayed gastric absorption of carbohydrate [55]. One study examined the intra-individual variability in postprandial glucose excursions in a small cohort of people with Type 1 diabetes on MDI, using standardized test meals with either insulin lispro (15 min pre-meal) or regular human insulin (30 min pre-meal). The intra-individual coefficients of variance of the mean glucose excursions after the meals were significant, and also lower with insulin lispro, at most time points: 1 h, 66% vs 71%; 2 h, 49% vs 69%; 4 h, 66% vs 75% and 5 h, 49% vs 72% [56].

There are, of course, some circumstances in which safety or practicality governs the timing of insulin. Examples may be people working in critical environments or where they cannot guarantee eating of food 15–20 min after a bolus, when eating out at social events or when predicting the exact carbohydrate content of the meal ahead is not possible. In these situations, pre- rather than post-meal administration remains optimal but, if safety or convenience leads to occasional post-meal administration of prandial insulin, rapid-acting analogues are the safer option.

Administration of fast-acting analogues by a parent uncertain of a child’s appetite within 15 min after the child starts to eat may be an acceptable compromise between parental anxiety and normoglycaemia, but it should not be allowed to become a habit as the child’s behaviour becomes more predictable, and adults with diabetes should be discouraged from postprandial injections. Another special circumstance, in the opposite direction, is pregnancy. A study by Murphy et al. [57] showed that postprandial BG levels are impaired by significantly slower glucose disposal in late gestation, with the authors suggesting that optimal bolus timing in late pregnancy may be 30–40 min pre-meal compared to 15–20 min in early pregnancy.

The site at which s.c. insulin is injected can also affect the PK characteristics of insulin. Abdominal injecting of rapid-acting insulin analogues results in the highest concentration of insulin at the earliest time when compared with insulin administration in the arm, thigh or buttocks [58].

**Conclusions**

The data from the present review of the literature provide clear clinical evidence for the superiority and safety of injecting 15–20 min pre-food, with almost 30% lower post-meal glucose levels, a lower AUC for hyperglycaemia and less post-meal hypoglycaemia when the pre-meal glucose levels are in range. We therefore recommend that people with
diabetes should aim to do this whenever possible, accepting that there may be individual circumstances when this is not practical. Postprandial administration of rapid-acting insulin analogues is a less effective method of controlling BG levels in the postprandial phase, and carries a significant risk of hypoglycaemia.

The PK/PD studies of rapid-acting insulin analogues show that the time to maximum insulin levels is between 40 and 60 min, but time to peak insulin effect is up to 120 min after injection. Given that BG levels peak before the maximum peak effect of insulin has been reached, it makes sense to administer rapid-acting insulin analogues 15–20 min before mealtimes to try to synchronize BG and insulin peaks in an attempt to avoid postprandial hyperglycaemia. Fear of hypoglycaemia by adopting this approach may prevent patients from following this advice but may be allayed by discussion of the lack of insulin action during that time. Indeed, the data show that risk of post-meal hypoglycaemia is highest with analogue insulins when administered 20 min after the start of eating. The delay in injecting after a meal in adults with Type 1 diabetes using post-meal insulin administration may often be greater than that, which would be expected to exacerbate the problems further.

As our understanding of post-meal glucose control increases, we may need to develop better strategies to cope with complex meals that may require different time–action profiles. With the advent of newer faster-acting analogues, we will need further clinical studies to understand the optimum timing of these insulins; however, given the profile of post-meal glucose excursions, the time to peak insulin action would need to be <45 min if these insulins were to be effectively used after eating.

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Competing interests
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