Continuous subcutaneous insulin infusion with short-acting insulin analogues or human regular insulin: efficacy, safety, quality of life, and cost-effectiveness

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Summary

Portable insulin infusion devices are effective and safe insulin delivery systems for managing diabetes mellitus, especially type 1 diabetes. Rapidly absorbed insulin analogues, such as insulin lispro or insulin aspart, may offer an advantage over regular human insulin for insulin pumps. Several open-label randomised crossover trials demonstrated that continuous subcutaneous insulin infusion (CSII) with insulin lispro provided a better control of postprandial hyperglycaemia and a slightly but significantly lower glycated haemoglobin level, with lower daily insulin requirement and similar or even less hypoglycaemic episodes. A CSII study comparing insulin lispro and insulin aspart demonstrated similar results with the two analogues, and better results than those with regular insulin. Because these analogues have a quicker onset and a shorter duration of action than regular insulin, one might expect an earlier and greater metabolic deterioration in case of CSII interruption, but a more rapid correction of metabolic abnormalities after insulin boluses when reactivating the pump. These expectations were confirmed in randomised protocols comparing the metabolic changes occurring during and after CSII interruption of various durations when the pump infused either insulin lispro or regular insulin. The extra cost resulting from the use of CSII and insulin analogues in diabetes management should be compensated for by better metabolic control and quality of life. In conclusion, CSII delivering fast-acting insulin analogues may be considered as one of the best methods to replace insulin in a physiological manner by mimicking meal and basal insulin requirements, without higher risk of hypoglycaemia or ketoacidosis in well-educated diabetic patients. Copyright © 2004 John Wiley & Sons, Ltd.

Keywords CSII; insulin analogue; insulin aspart; insulin lispro; insulin pump; ketoacidosis

Introduction

Continuous subcutaneous insulin infusion (CSII), often called insulin-pump therapy, was introduced in the 1970s as a way of achieving and maintaining strict control of blood glucose concentrations in people with type 1 diabetes [1]. Administration by CSII has provided additional flexibility in meal timing and modifying basal insulin replacement in response to circadian rhythms [2]. In the Diabetes Control and Complications Trial (DCCT), although this trial was not specifically designed to compare CSII and multiple daily injection therapies, lower glycated haemoglobin (HbA1c) levels could be achieved with
CSII because of a greater reproducibility and flexibility of insulin administration [3]. The superiority of CSII on multiple daily insulin injections, that is, better glycaemic control and stability with lower daily doses of insulin and reduction of hypoglycaemic episodes, has been confirmed in several studies [4,5] and in a recent meta-analysis of 12 randomized controlled trials comparing the two treatment modalities [6]. These advantages of CSII, as well as improvements in pump technology and new reimbursement modalities, have led to increasing acceptance and use of insulin-pump therapy [7–11].

Fast-acting insulin analogues offer the advantages of a faster and shorter hypoglycaemic action [12–14]. Numerous studies demonstrated that both insulin lispro [15–18] and insulin aspart [19–23] are able to better control postprandial hyperglycaemia, even when they are injected just before meals. Another advantage is a significant reduction in the risk of hypoglycaemic episodes and severe hypoglycaemia [16,24], as well as nocturnal hypoglycaemia [25], a favourable effect that has been confirmed in a meta-analysis of eight studies [26]. However, the positive impact on average blood glucose control, as assessed by HbA1C, was less obvious and could only be demonstrated in clinical trials in which adjustment of basal insulin supplements [27] and/or reduction of snacks [28] were recommended. The interpretation of such results was that while early postprandial hyperglycaemia was reduced by fast-acting insulin analogue, late plasma glucose levels were higher because of the extremely short action of the insulin analogue and the imperfect basal supplementation using intermediate-acting insulin such as NPH insulin once daily [29]. Delayed hypoglycaemia could theoretically be avoided when using the portable pump, as CSII allows adequate insulin delivery throughout the day and night, using variable insulin delivery rates when necessary. In addition, because the concept of CSII is predicated on an immediacy of insulin action [30,31], a rapid-insulin analogue should be ideal. One study demonstrated that potency and purity of insulin lispro were practically unchanged after 2 days of pumping in two different insulin infusion systems when syringes and catheters are replaced at 48-h intervals [32]. Thus, fast-acting insulin analogues are considered as the gold standard insulin for pump therapy, allowing better postprandial and overall glucose control without increasing the risk of hypoglycaemia [33,34]. However, in the most recent meta-analysis of the metabolic and psychosocial impact of CSII therapy in adults, adolescents, and children, including a total of 52 studies, consisting of 1547 patients, no single consideration was mentioned about the possible influence of the type of insulin delivered by the pump, short-acting insulin analogue versus regular human insulin, on the various outcomes considered [11].

One of the problems associated with portable pump therapy may be a higher risk of ketoadicotic episodes [35–37]. Indeed, technical problems (pump failure, catheter occlusion or disconnection, skin infection) may impair insulin delivery or insulin absorption and cause acute metabolic disorders. Properties of insulin analogues, that is, soluble, rapid-acting, and uniform absorption, should reduce the size of the subcutaneous insulin depot and, therefore, reduce the time interval between stopping of insulin delivery and occurrence of acute insulin deficiency, on the one hand, and reduce the time interval between administering recovery insulin boluses and reaching plasma-free insulin peaks, on the other hand. Thus, because of the peculiar pharmacokinetic profile of these two short-acting insulin analogues [15,18,22,23], an earlier insulin deprivation might occur in case of CSII interruption, while a more rapid metabolic correction of hyperglycaemia and ketosis might be attained after administration of insulin lispro or aspart than after that of human regular insulin.

This concise review aims at comparing both the efficacy on metabolic control, assessed by HbA1C levels, and safety, assessed by hypoglycaemia and ketoacidosis incidence, in CSII-treated type 1 diabetic patients using either short-acting insulin analogues (lispro or aspart) or human regular insulin. Moreover, we will compare the consequences of CSII interruption in diabetic patients treated either with regular insulin or with insulin lispro, as well as the efficacy of a rescue insulin replacement scheme after restarting the pump following several hours of insulin delivery interruption. Safety concern will also be discussed, especially when using CSII and insulin analogues during pregnancy. Finally, possible advantages on quality of life and cost-effectiveness analysis of these modalities of insulin replacement therapy will be briefly presented.

CSII and metabolic control: insulin lispro versus human regular insulin

At least six controlled trials compared the efficacy and safety of insulin lispro versus human regular insulin in CSII-treated type 1 diabetic patients [38–45]. The first pilot, double blind, crossover, comparative study treated 30 diabetic patients for 3 months with insulin lispro and for 3 months with human regular insulin in a randomized order [38]. All boluses were given immediately before the three main meals. At the end of the three-month treatment period, HbA1C levels were significantly lower with insulin lispro compared to those with human regular insulin (7.66 ± 0.13 vs 8.0 ± 0.16%; p < 0.005). One-hour postprandial blood glucose concentrations were significantly improved after breakfast, lunch, and dinner with insulin lispro, compared to those with regular insulin. The incidence of hypoglycaemia tended to be lower (NS) with insulin lispro than with regular insulin. The authors concluded that insulin lispro improves glycaemic control in CSII without increasing the risk of hypoglycaemia.

Five open-label, randomized, crossover trials compared insulin lispro with human regular insulin (two successive experimental periods of 2 to 4 months) in type 1 diabetic patients treated with a portable pump [39–43]. In all

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these open studies, except one [43], and in contrast to the previous double-blind trial [38], insulin boluses were given 0 to 5 min before meals for insulin lispro and 20 to 30 min before meals for regular insulin. Nevertheless, all trials reported reduced postprandial hyperglycaemia and lower HbA1c levels with insulin lispro as compared to those with regular insulin (Table 1). However, the differences were rather small (from −0.10 to 0.53% for HbA1c), although statistically significant, except in the study with the lower number of subjects [39]. The differences in the changes in HbA1c across the studies may be related to duration of study, intensity of glycose control, or timing of insulin bolus doses relative to meals. Hypoglycaemic episodes (usually defined as blood glucose levels <3 mmol/L) were not significantly different between the two treatment modalities in every study (Table 1).

A study demonstrated a positive impact on clinical status and quality of life of switching from regular human insulin to insulin lispro among patients using insulin pumps [44]. These results were confirmed in a long-term study comparing the therapeutic efficacy of insulin lispro with that of buffered regular human insulin in 62 patients on insulin-pump therapy [45]. The patients, initially treated with regular human insulin for 20 months, switched to using insulin lispro for another mean of 20 months. HbA1c level was significantly lower with insulin lispro than with regular insulin (7.4 vs 7.9%; \( p < 0.001 \)). Basal insulin requirements were higher, while pre-meal insulin boluses were lower with insulin lispro than with those with regular insulin, the total daily units of insulin being slightly but significantly lower during therapy with the fast-acting insulin analogue. The numbers of mild/moderate and severe hypoglycaemic episodes were similar in the two treatment periods. This first long-term study demonstrates the sustained efficacy of insulin lispro as compared to that of buffered regular insulin in decreasing HbA1c, without increasing the incidence of hypoglycaemia, among insulin-pump users.

One case report compared glucose profiles, HbA1c levels, and the risk of severe hypoglycaemia in one patient who successively moved from CSII using regular acting insulin to CSII using insulin lispro, and finally implantable pump with intraperitoneal insulin delivery [46]. Longitudinal data in this patient suggested that insulin lispro only tended to reduce average glycaemia and glycaemic fluctuations as compared to regular insulin, whereas a much greater improvement could be achieved with the implantable pump delivering insulin intraperitoneally, a finding recently confirmed in a pilot controlled study [47].

Thus, one key advantage of insulin lispro in CSII is the similar or even reduced incidence of hypoglycaemic episodes despite a better overall metabolic control, as compared to that of regular human insulin [33]. One specific experimental trial demonstrated that the counter-regulatory hormone responses are maintained with the use of insulin lispro in CSII, compared to that of regular human insulin, resulting in improved hepatic glycogen output in response to glucagon [48].

### Table 1. Changes in postprandial hyperglycaemia, HbA1c levels, and incidence of hypoglycaemic episodes in clinical trials comparing insulin lispro with human regular insulin in type 1 diabetic patients treated with continuous subcutaneous insulin infusion

<table>
<thead>
<tr>
<th>References</th>
<th>( n )</th>
<th>Period (months)</th>
<th>Design</th>
<th>Postprandial hyperglycaemia ( \Delta ) (mmol/L)</th>
<th>HbA1c ( \Delta ) (%)</th>
<th>Hypoglycaemia ( \Delta ) (30 days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zinman et al. 1997 [38]</td>
<td>30</td>
<td>2 x 3</td>
<td>D8 CO</td>
<td>−1.44 ( p = 0.006 )</td>
<td>−0.34 ( p &lt; 0.001 )</td>
<td>−1.6 ( NS )</td>
</tr>
<tr>
<td>Schmauss et al. 1998 [39]</td>
<td>11</td>
<td>2 x 3</td>
<td>Open CO</td>
<td>−1.50 ( p = 0.030 )</td>
<td>−0.35 ( NS )</td>
<td>+0.8 ( NS )</td>
</tr>
<tr>
<td>Melki et al. 1999 [40]</td>
<td>39</td>
<td>2 x 3</td>
<td>Open CO</td>
<td>−1.64 ( p = 0.001 )</td>
<td>−0.53 ( p &lt; 0.01 )</td>
<td>−0.9 ( NS )</td>
</tr>
<tr>
<td>Renner et al. 1999 [41]</td>
<td>113</td>
<td>2 x 4</td>
<td>Open CO</td>
<td>−1.60 ( p = 0.001 )</td>
<td>−0.1 ( p &lt; 0.01 )</td>
<td>−0.4 ( NS )</td>
</tr>
<tr>
<td>Johansson et al. 2000 [42]</td>
<td>41</td>
<td>2 x 2</td>
<td>Open CO</td>
<td>−1.50 ( p = 0.001 )</td>
<td>−0.2 ( p &lt; 0.01 )</td>
<td>−0.4 ( NS )</td>
</tr>
<tr>
<td>Raskin et al. 2001 [43]</td>
<td>58</td>
<td>2 x 3</td>
<td>Open CO</td>
<td>−2.04 ( p = 0.012 )</td>
<td>−0.23 ( p &lt; 0.005 )</td>
<td>0 ( NS )</td>
</tr>
<tr>
<td>Garg et al. 2000 [45]</td>
<td>52</td>
<td>2 x 20</td>
<td>Switch</td>
<td>NA ( p &lt; 0.001 )</td>
<td>−0.50 ( NS )</td>
<td>−0.4 ( NS )</td>
</tr>
<tr>
<td>Bode et al. 2002 [34]</td>
<td>28(+)</td>
<td>4</td>
<td>Open parallel</td>
<td>0 ( NS )</td>
<td>+0.03 ( NS )</td>
<td>−0.4 ( NS )</td>
</tr>
</tbody>
</table>

\( \Delta \), Value with insulin lispro minus value with human regular insulin; CO, crossover; DB, double blind; Switch, from regular insulin to insulin lispro; NA, not available; NS, not significant; (+) 59 patients received human regular insulin.
insulin, NS). A similar number of patients experienced hypoglycaemia (blood glucose <2.5 mmol/L) during the study (74% with insulin aspart vs 60% with regular insulin; NS). The authors concluded that insulin aspart and regular human insulin were effective and well tolerated when used in CSII therapy.

Only one randomized study compared the three types of insulin, that is, insulin aspart (n = 59) versus buffered human insulin (n = 59) versus insulin lispro (n = 28) [34]. In this recent multicentre, open-label, randomized, parallel-group study, bolus insulin doses were administered 30 min before meals for regular insulin or immediately before meals for the two insulin analogues. After 16 weeks of treatment, mean changes in baseline HbA1c values were not significantly different between the three groups (0.00% for aspart, 0.15% for regular, and 0.18% for lispro; NS). The rates of hypoglycaemic episodes per patient per month were similar in the three groups. The conclusion was that insulin aspart in CSII was as efficacious and well tolerated as regular insulin and insulin lispro and is a suitable insulin for external pump therapy.

Thus, the use of a short-acting insulin analogue in the pump improves HbA1c and blood glucose stability, without increasing the risk of hypoglycaemia. However, frequent blood glucose self-monitoring is required in order to optimize insulin adjustments [51].

**Interruption of CSII with regular insulin: comparison with normal functioning of the pump**

CSII use was initially associated with an overall increased frequency of diabetic ketoacidosis [35,36]. However, as pointed out in a recent meta-analysis of 52 studies dealing with insulin-pump therapy [11], this increased risk of diabetic ketoacidosis was not evident in studies published after 1993, suggesting that CSII may no longer be associated with a greater risk of ketoacidotic episodes compared to other forms of insulin administration. It may be that earlier data on the increased risk of diabetic ketoacidosis led clinicians to emphasize ketoacidosis prevention with their patients and manufacturers to improve the security of insulin pumps by inserting several alarms in case of malfunction of the device. Thus, with proper education and pump practice, the frequency of ketoacidosis appears to be the same on CSII and insulin injection therapy [10].

In an original pilot study, our group investigated the changes in blood glucose, plasma non-esterified fatty acids (NEFA), 3-hydroxybutyrate (3-OHB), glucagon, and free insulin in eight C-peptide-negative type 1 diabetic patients whose pumps were deliberately stopped between 23.00 h to 05.00 h [52]. A control test with the pump functioning normally was carried out in each patient and the experimental protocols were carried out in a random order. Considering the values at 23.00 h as reference, interruption of insulin infusion resulted in (1) a rapid decrease in plasma-free insulin levels, significant after 1 h and reaching a nadir after 6 h; (2) a rise in blood glucose that was significant at hour 3 and reached 17.4 ± 1.9 mmol/L at hour 6; (3) a moderate increase in plasma NEFA concentrations that remained in the range of 700 to 800 μmol/L; (4) an early and linear rise in plasma 3-OHB, significant after 1 h and averaging 1290 ± 140 μmol/L after 6 h, with presence of ketonuria; and (5) a late increase (hour 5) in plasma glucagon.

Besides insulin deprivation due to pump arrest, counter-regulatory hormones play a significant role in the increase of blood glucose and 3-OHB levels, as such increments were reduced almost by half after either continuous intravenous infusion of somatostatin or a single subcutaneous injection of octreotide (a long-acting somatostatin analogue) inhibiting both growth hormone and glucagon secretion [review in 37]. Our group demonstrated that several other factors significantly influence the amplitude and kinetics of metabolic deterioration: the residual insulin secretion (less deterioration in C-peptide positive diabetic patients), the quality of previous metabolic control (less deterioration in subjects with prevailing normoglycaemia), insulin concentration in the pump cartridge (less deterioration when using concentrated insulin), and the presence of insulin antibodies (less deterioration in patients with high circulating IgG antibodies) [37]. Finally, as expected, the duration of CSII interruption plays a crucial role. We demonstrated that 1-h pump arrest does not significantly alter the metabolic profile in well-controlled diabetic patients, while a 2-h period of CSII interruption is enough to induce a significant rise in plasma glucose and 3-OHB levels; in this condition, additional insulin boluses are required to prevent or correct the metabolic disorders [53].

Thus, the prolonged, either intentional or accidental, interruption of CSII delivering regular human insulin has metabolic consequences that should be carefully borne in mind. These consequences reflect the increased risk of ketoacidotic episodes reported in diabetic patients treated with CSII [37]. This risk can be markedly reduced if regular home blood glucose monitoring is performed and if the well-educated diabetic patient reacts rapidly to incipient metabolic deterioration with the administration of insulin supplements based on a regimen determined by blood glucose and the presence or absence of ketonuria [37,51]. From a practical point of view, in well-controlled diabetic patients, it seems reasonable to permit short-term interruption of CSII for up to 1 h, while longer interruption must be followed by insulin supplements [54].

**Interruption of CSII with insulin lispro: comparison with regular insulin**

Three controlled studies compared the metabolic consequences of CSII interruption when using insulin lispro versus regular insulin [55–57]. All trials studied C-peptide-negative type 1 diabetic patients, and measured
at regular intervals various metabolic (plasma glucose, NEFA, 3-OHB, ketonuria) and hormonal (plasma-free insulin, glucagon) parameters. However, the protocols differed as far as the study design (parallel vs crossover), the duration of CSII interruption (5 vs 6 vs 9 h), the time period of pump arrest (nocturnal vs diurnal), and the delay since the last insulin bolus administration were concerned (Table 2).

A total of 18 well-controlled type 1 diabetic patients were studied: 9 received human regular and 9 received insulin lispro. Basal insulin infusion was stopped from 03.00 to 09.00 h [55]. Plasma glucose concentrations rose to 13.8 ± 1.9 and 16.0 ± 1.7 mmol/L in the regular insulin- and insulin lispro–treated groups respectively (NS). No significant differences were seen between the therapy groups at any time in the insulin levels or in the concentrations of plasma glucose or 3-OHB. The authors concluded that interruption of the basal insulin infusion in the middle of the night does not result in more rapid metabolic deterioration in patients treated with lispro compared to those treated with regular insulin. However, this trial used a parallel-group study design rather than a crossover protocol, which may hinder the demonstration of differences between the two insulins as different groups of patients were used for the comparison.

An open-label randomized crossover trial investigated seven type 1 diabetic patients treated with portable pump. CSII was interrupted from 22.00 to 07.00 h the next day in two experimental conditions: either with regular insulin or with insulin lispro [56]. With insulin lispro, the metabolic changes developed 1.5 to 2 h earlier than with regular human insulin. After 3 h, blood glucose averaged 4.93 ± 2.87 mmol/L with regular insulin and 8.97 ± 3.48 mmol/L with insulin lispro (p < 0.05). Decreases in base excess were significantly greater with insulin lispro (−1.69 ± 0.83 mmol/L) than with regular insulin (−0.41 ± 1.04 mmol/L; p < 0.05).

Another randomized, crossover, open-label trial comparing insulin lispro and regular insulin was performed in 10 type 1 diabetic patients [57]. CSII was interrupted from 07.00 to 12.00 h. The plasma-free insulin level decreased significantly with the two treatments, but was significantly lower with lispro than with regular insulin (p < 0.05–0.01). The plasma glucose level was significantly higher in the insulin lispro group than in the regular insulin group from 120 until 300 min after stopping CSII (13.93 ± 3.77 vs 10.77 ± 4.38 mmol/L, p < 0.05 at 180 min and 17.04 ± 3.27 vs 12.98 ± 4.33 mmol/L, p < 0.01 at 300 min). Plasma NEFA concentrations increased more rapidly and were significantly higher in the lispro group than in the regular group (p < 0.05–0.01). Plasma 3-OHB increased earlier with lispro (60 vs 120 min), but was not statistically different between the two treatments.

Thus, the two trials using a randomized crossover study protocol, allowing a direct comparison in the same diabetic patients, reported similar results and concluded that metabolic deterioration occurred earlier and was of greater amplitude with insulin lispro than with human regular insulin [56,57].

Metabolic correction with regular Insulin versus insulin lispro replacement scheme

After a 6-h CSII nocturnal (23.00–05.00 h) interruption using regular insulin, a scheme for a prompt return to adequate control was tested in eight C-peptide-negative type 1 diabetic patients [52]. This scheme consisted of insulin supplements administered via the pump and based on blood glucose monitoring and semi-quantitative evaluation of ketonuria. Resetting the pump at its basal rate at 05.00 h and giving insulin supplements (2–8 U) at 06.45 h (with the usual breakfast dose) and again at 10.00 h have proved efficacious in restoring rapidly satisfactory metabolic control. Indeed, plasma glucose and 3-OHB levels were similar at noon, both after the CSII interruption test and in the test with normal functioning of the pump throughout the night. This study allowed establishing precise guidelines that permit the patient to restore adequate metabolic control within a few hours on the basis of simple blood and urine determinations performed by the patient himself/herself [54].

Table 2. Changes in plasma-free insulin, glucose, and 3-OHB levels during CSII interruption in three trials comparing regular insulin with fast-acting insulin lispro. When precise data were not directly available in the original paper, they were estimated from the curves of the corresponding figures

<table>
<thead>
<tr>
<th>References</th>
<th>Δ Free insulin (pmol/L)</th>
<th>Δ Glucose (mmol/L)</th>
<th>Δ 3-OHB (µmol/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Attia et al. 1998 [55]</td>
<td>Off/Regular (n = 9)</td>
<td>−47</td>
<td>+13.0</td>
</tr>
<tr>
<td>(8h-stop of CSII: parallel)</td>
<td>Off/Lispro (n = 9)</td>
<td>−60</td>
<td>+12.2</td>
</tr>
<tr>
<td>Reichel et al. 1998 [56]</td>
<td>Off/Regular (n = 7)</td>
<td>−10.0</td>
<td>+12.6</td>
</tr>
<tr>
<td>(9h-stop of CSII: crossover)</td>
<td>Off/Regular (n = 7)</td>
<td>−36</td>
<td>−4.9</td>
</tr>
<tr>
<td>Guerdi et al. 1999 [57]</td>
<td>Off/Regular (n = 10)</td>
<td>NA</td>
<td>p &lt; 0.05</td>
</tr>
</tbody>
</table>

Δ, Value at the end of CSII interruption minus baseline value; (*), base excess (instead of 3-OHB not measured in this study); NA, not available; NS, not significant.

In the study of Attia et al. [55], patients with plasma glucose levels >13.9 mmol/L or moderate ketonuria at the end of a 5-h pump arrest received a single subcutaneous injection of either human regular or insulin lispro. In this insulin-replacement phase, plasma-free insulin levels rose more rapidly in those treated with insulin lispro, reaching a greater peak value (150 ± 20 vs 94 ± 16 pmol/L, p < 0.05, at 60 min), while plasma glucose concentrations decreased to a lower nadir after insulin lispro (9.7 ± 0.4 vs 13.7 ± 0.7 mmol/L, p < 0.01, at 120 min after insulin administration). Plasma 3-OHBD levels decreased rapidly in both groups, the slope of the reduction being more abrupt with insulin lispro, allowing finishing the test with roughly similar values.

In the study performed by Guerci et al. [57], the pump of each patient was reactivated at its usual basal rate at 12.00 h (after 5 h of CSII interruption), at which time the patients ate a similar calibrated lunch and activated their usual pre-lunch insulin boluses. Additionally, a correcting insulin scheme (0-4 U) was given each hour afterwards according to plasma glucose levels and ketonuria. After restarting the pump, the plasma-free insulin peak occurred earlier and was greater with lispro than with regular insulin (224 ± 121 vs 97 ± 31 pmol/L; p < 0.05). As a consequence, plasma glucose concentrations decreased with insulin lispro, but continued to increase with regular insulin during the first 2 h after restarting the pump (the difference being statistically significant 60 min after insulin replacement). Plasma NEFA and 3-OHBD levels decreased significantly with the two treatments, but more dramatically with lispro treatment.

Thus, the two studies comparing correction schemes with insulin lispro and regular insulin concluded that the fast-acting insulin analogue is more effective in treating mild ketosis and hyperglycaemia following CSII interruption [55,57]. In well-educated patients, this may represent an advantage provided that early detection of the metabolic deterioration is made and prompt adequate response is brought.

**Immunological considerations with short-acting insulin analogues during pump therapy**

It has been suggested that CSII [58] as well as long-term intraperitoneal insulin administration via implantable programmable insulin delivery systems [59] may favour the development of insulin antibodies. However, the role of these insulin antibodies on metabolic control remains controversial [60,61]. These antibodies might play the role of reservoir in certain circumstances and, for instance, dampen the metabolic deterioration occurring after CSII interruption [62,63]. To our knowledge, no study specifically investigated the development of antibodies during CSII when delivering short-acting insulin analogues.

Insulin lispro and recombinant human insulin have similar immunogenicity [64]. Recently, a multinational, multicentre combination of controlled and non-controlled, open-label studies of 4.5 years’ duration evaluated the long-term immunologic profile of subcutaneously administered insulin lispro [65]. By measuring lispro-specific, insulin-specific, and cross-reactive antibodies, it was concluded that the immunogenic profile of patients treated with insulin lispro was comparable to that of patients treated with recombinant human insulin. In addition, the incidence of insulin allergy was not different from that in patients treated with recombinant regular human insulin. Anecdotal reports suggested that both antibody-mediated insulin resistance [66] and generalised allergy to human insulin [67] could be successfully treated with insulin lispro. Insulin desensitization with CSII delivering insulin lispro has been reported in several type 1 diabetic patients [68–70], suggesting that the insulin pump and short-acting insulin may be useful as alternative treatments in insulin allergy.

Lipoatrophy as a cutaneous complication of insulin therapy has been extremely rare since the introduction of recombinant human insulin. Two cases of lipoatrophy associated with lispro insulin have been reported in two insulin-pump-treated diabetic patients [71]. Recently, a singular case in which lipoatrophy occurred in two different locations with both buffered regular human insulin and insulin lispro has been described in a patient treated by CSII [72]. Interestingly, the lispro-induced lipoatrophy area was smaller than that associated with human regular insulin, a difference that might be related to the reduced ability of insulin lispro to aggregate.

**CSII with insulin analogues during pregnancy**

CSII has been advocated as an alternative to multiple-dose insulin injections in diabetic pregnant women as it more closely mimics physiological insulin delivery and allows better metabolic control [73–75]. Only few studies have investigated the safety of insulin lispro in human pregnancy as far as foetal and maternal outcomes are concerned [76–78]. The risk of severe hypoglycaemic or ketoacidotic episodes may be particularly dramatic in pregnant women, two complications that could be positively or negatively influenced by the use of both CSII and short-acting insulin analogues. Apparently, the metabolic observations made in non-pregnant women were confirmed during pregnancy with a trend to better blood glucose control with lower hypoglycaemic episodes and no significant increase in severe ketoacidotic episodes [73–75]. Because the risk of developing ketoacidosis is higher during pregnancy, the problem of pump arrest should be a matter of concern in pregnant diabetic women. One of the most important criteria in selecting patients for CSII is their willingness to test their capillary glucose levels several times each day [73].
This prerequisite is especially important when fast-acting insulin analogue is used in the pump in order to detect early metabolic deterioration and administer insulin bolus to promptly restore adequate metabolic control in well-educated women. Despite better acceptability, less hypoglycaemia and possibly better glycaemic control, doubts were raised by a few reported cases of congenital anomalies using insulin lispro [77,78]. However, some controlled studies were unable to find any increase in the adverse outcome using insulin lispro in diabetic pregnancies, in either gestational or pre-gestational diabetes [77,79]. A recent report on pooled data from seven centres with experience in the use of insulin lispro during pregnancy mentioned comparable outcomes to other large studies using conventional insulin [80]. A recent study examined whether insulin lispro crosses the placenta using the technique of perfusing a human placental lobule in vitro [81]. It showed that insulin lispro is not likely to cross the placenta at a single standard dose and suggested that insulin lispro is unlikely to reach and harm the unborn baby. These results are in agreement with the absence of detectable insulin lispro in the cord blood of newborns from mothers who received continuous intravenous infusion of insulin lispro and dextrose during labour [76].

Only scarce data are available yet on the efficacy and safety of the use of insulin aspart in pregnant women. A recent study demonstrated that effective postprandial glycaemic control in women with gestational diabetes mellitus who required insulin was brought about by insulin aspart, as compared to regular human insulin, through higher insulin peak and lower demand on endogenous insulin secretion [82]. It has been suggested that large prospective randomized clinical trials should be performed for further evaluation of any possible association between the use of insulin lispro or insulin aspart during pregnancy and an increased rate of congenital malformations [79]. Up to now, rapid-acting insulin analogues are not approved for use in pregnancy [83].

In one report, rapid acceleration of proliferative retinopathy was seen during pregnancy in 3 of 10 women with diabetes treated with insulin lispro [84]. In other reports, insulin lispro was not associated with the progression of diabetic retinopathy during pregnancy [77,85]. A recent prospective open study of 69 pregnant women with diabetes (36 treated with insulin lispro and 33 treated with conventional regular human insulin) confirmed that insulin lispro improves glycaemic control during diabetic pregnancy compared to regular insulin, with no adverse impact on progression of diabetic retinopathy [86].

Education and patient’s selection for CSII

Only well-educated diabetic patients are those who benefit more from CSII. As emphasised by the American Diabetes Association recommendations [83], CSII therapy should ideally be prescribed, implemented, and followed by a skilled professional team familiar with CSII therapy and capable of supporting the patient. Experience with insulin-pump therapy indicates that candidates for CSII must be strongly motivated to improve glucose control and willing to work with their health-care provider in assuming substantial responsibility for their day-to-day care. They must also understand and demonstrate appropriate use of the insulin pump and self-monitoring of blood glucose [10,83]. Patients who are not compliant on insulin injection regimens will do badly on pump therapy and, in general, should not be selected [10]. All clinical trials demonstrating a significant advantage of short-acting insulin analogues, in comparison with regular human insulin, as far as metabolic control was concerned, were obtained in CSII-treated patients who satisfied all these prerequisites. We can also speculate that only those patients will benefit from an increased quality of life with an acceptable cost-effectiveness ratio when using such sophisticated methods of insulin delivery.

CSII with insulin analogues and quality of life

Quality of life is an important health outcome, particularly in persons with a chronic disease such as diabetes mellitus. It is therefore important that the ability of patients to comply and their satisfaction with treatment, together with the effect of therapy on overall quality of life, are taken into account by physicians. Both CSII and fast-acting insulin analogues may positively influence the quality of life of diabetic patients.

Among CSII-treated patients, the effectiveness and flexibility of CSII appeared to compensate for problems caused by both the external pump and the subcutaneous catheter, and for the interference with various activities like work, sex, and sport. The favourable disposition towards CSII appears to stem from the feeling that CSII is better than multiple daily injections in achieving good metabolic control and produces a sense of well-being, increased freedom, and greater autonomy [87]. Patient reactions to CSII have been largely enthusiastic and the discontinuation rates low [10,88].

Increased satisfaction with treatment and improved quality of life with insulin lispro relative to regular human insulin have been shown in a number of studies with quality of life as primary and secondary endpoints in diabetic patients [89]. A large majority of patients expressed a preference for insulin lispro over human soluble insulin on the grounds of increased freedom with meal timing and increased independence in everyday activities. However, it should be noted that most of these trials were conducted in an uncontrolled manner. Quality-of-life data were reported in great detail by Kotsanos et al. [90], who based their analysis on a large non-blind randomized comparative crossover trial.
in 1008 patients with type 1 diabetes, of whom 468 were available for quality-of-life assessment. An expanded and modified form of the 'Diabetes Quality of Life Clinical Trial Questionnaire' was used with two diabetes-specific domains (treatment satisfaction and treatment flexibility) being specified as primary outcomes. After 3 months' treatment, only mean treatment flexibility and treatment satisfaction scores were significantly increased relative to baseline to a greater extent with insulin lispro than with human soluble insulin. All other primary and secondary domains of health-related quality-of-life findings were comparable for insulin lispro and regular human insulin. Favourable results were also recently reported in a 6-month, multinational, randomized, large, open-label trial comparing insulin aspart with soluble human insulin in 424 adult type 1 diabetic patients on an insulin basal-bolus scheme, but not CSII [91]. Under the study conditions, insulin aspart improved treatment satisfaction and quality of life with regard to diet restriction when compared with human regular insulin, and this effect was judged as being 'not trivial'.

It should be emphasised, however, that none of these studies assessing quality of life included patients treated with CSII and that none of the above-mentioned studies comparing short-acting insulin versus human soluble insulin in CSII-treated patients with type 1 diabetes reported specific data on quality of life.

Pharmacoeconomical aspects of CSII and insulin analogues

It is inevitable that new treatments for diabetic patients will be considered critically by health-care planners and providers in the prevailing global environment of increasing costs of medical care and pressure for rational allocation of resources [89]. Both portable insulin pumps and short-acting insulin analogues represent an extra cost as compared to classical insulin therapy using conventional insulin injections of human regular insulin. Intensive insulin therapy reduces costs by decreasing complications [92]. However, neither CSII nor short-acting insulin analogues have proven their superiority in reducing the risk of microvascular or macrovascular complications in long-term studies. Thus, the only available data concern the possible reduction of acute metabolic complications, such as severe hypoglycaemic episodes.

A Markov model was recently constructed to estimate the costs and outcomes for patients with type 1 diabetes treated with CSII compared to those treated with multiple daily injections [93]. The primary outcome was quality-adjusted life years (QALYs). It was concluded that CSII is a worthwhile investment when targeted at those who might benefit most. Results were more sensitive to the number of hypoglycaemic events per patient. CSII was most cost-effective in patients who had more than two severe hypoglycaemic events per year and who required admission to hospital at least once every year. Cases in which CSII might not be economically viable are cases in which diabetes is well controlled with few severe hypoglycaemic events on conventional insulin therapy.

Pharmacoeconomic studies of insulin lispro used surrogate or short-term clinical endpoints rather than longer-term effects such as vascular complications avoided, and study details were often lacking [89]. Participants in well-designed studies have expressed a preference for lispro-based insulins and have been shown to be willing to pay for the advantages they offer [89]. The limited cost-effectiveness data currently available for insulin lispro suggest that the additional cost associated with the insulin analogue relative to regular human insulin is justified in terms of cost per episode of severe hypoglycaemia avoided or per 1% reduction in HbA1c level. However, details of methods and sources and determination of costs and robustness of data were often lacking in the available reports [89]. Thus, further research into the pharmacoeconomic implications of short-acting insulin analogue use in the long-term is needed, particularly with respect to effects of indirect costs and those associated with diabetic complications. It would be of much interest to perform such an analysis in a large cohort of type 1 diabetic patients treated in the long term by CSII delivering short-acting insulin analogues.

Conclusions

Approximately 80 years after the discovery and first human use of insulin, we are still striving to replace insulin in a physiological manner. Administration by CSII is best mimicking basal insulin secretion and meal insulin requirements. The fast-acting nature of insulin analogues provides patients with greater flexibility of their insulin requirements because the bolus can be administered immediately before meals instead of minus 20 to 30 min before meals as recommended for human regular insulin. This greater flexibility likely leads to improved compliance and a better quality of life for patients using CSII therapy. In addition, better control of postprandial hyperglycaemia leads to slightly improved HbA1c levels, without increasing the risk of hypoglycaemia, as confirmed in a recent meta-analysis of clinical trials comparing CSII with short-acting insulin analogues versus human regular insulin [94]. Most studies tested insulin lispro, although the only clinical trial comparing directly lispro versus aspart insulin reported similar results with the two insulin analogues.

A several-hour interruption of CSII containing insulin lispro is associated with an earlier and greater metabolic deterioration than when the pump delivers human regular insulin. Consequently, patients treated with short-acting insulin analogues (either lispro or aspart) in an insulin pump have to be well educated about the pharmacokinetic properties of their insulin and about the possibility that ketoacidotic deterioration after an interruption of the insulin delivery may occur somewhat
earlier in comparison to regular human insulin. Finally, the fast-acting insulin analogue, owing to its more rapid absorption from the subcutaneous site, is more effective in correcting this metabolic deterioration after restarting the pump, which may partially compensate for the higher risk of ketoadocis in case of CSII interruption.

Both CSII and short-acting insulin analogues may offer some improvements regarding quality of life relative to conventional administration of regular human insulin, especially as far as treatment flexibility is concerned. However, further research into the pharmaco-economic implications of CSII and short-acting insulin analogues in the long term is needed, particularly with respect to effects on indirect costs and those associated with complications of diabetes mellitus.

In conclusion, improved insulins, better methods of insulin delivery, and advances in glucose monitoring will allow to progress towards physiological insulin replacement and, hopefully, reduce the long-term complications of diabetes mellitus.

References


