Short-acting insulin analogues versus regular human insulin on postprandial glucose and hypoglycemia in type 1 diabetes mellitus: a systematic review and meta-analysis

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Abstract

Introduction: Strict glucose control using multiple doses of insulin is the standard treatment for type 1 diabetes mellitus (T1DM), but increased risk of hypoglycemia is a frequent drawback. Regular insulin in multiple doses is important for achieving strict glycemic control for T1DM, but short-acting insulin analogues may be better in reducing hypoglycemia and postprandial glucose levels.

Objective: We conducted a systematic review and meta-analysis of randomized controlled trials (RCTs) to assess the effects of short-acting insulin analogues vs regular human insulin on hypoglycemia and postprandial glucose in patients with T1DM.

Methods: Searches were run on the electronic databases MEDLINE, Cochrane-CENTRAL, EMBASE, ClinicalTrials.gov, LILACS, and DARE for RCTs published until August 2017. To be included in the study, the RCTs had to cover a minimum period of 4 weeks and had to assess the effects of short-acting insulin analogues vs regular human insulin on hypoglycemia and postprandial glucose levels in patients with T1DM. Two independent reviewers extracted the data and assessed the quality of the selected studies. The primary outcomes analyzed were hypoglycemia (total episodes, nocturnal hypoglycemia, and severe hypoglycemia) and postprandial glucose (at all times, after breakfast, after lunch, and after dinner). Glycated hemoglobin (HbA1c) levels and quality of life were considered secondary outcomes. The risk of bias of each RCT was assessed using the Cochrane Collaboration Risk of Bias table, while the quality of evidence for each outcome was assessed using the GRADEpro software. The pooled mean difference in the number of hypoglycemic episodes and postprandial glucose between short-acting insulin analogues vs. regular human insulin was calculated using the random-effects model.

Results: Of the 2897 articles retrieved, 22 (6235 patients) were included. Short-acting insulin analogues were associated with a decrease in total hypoglycemic episodes (risk rate 0.93, 95% CI 0.87–0.99; 6235 patients; I² = 81%), nocturnal hypoglycemia (risk rate 0.55, 95% CI 0.40–0.76, 1995 patients, I² = 84%), and severe hypoglycemia (risk rate 0.68, 95% CI 0.60–0.77; 5945 patients, I² = 0%); and with lower postprandial glucose levels (mean difference/MD −19.44 mg/dL; 95% CI −21.49 to −17.39; 5031 patients, I² = 69%) and lower HbA1c (MD −0.13%; IC 95% −0.16 to −0.10; 5204 patients; I² = 73%) levels.

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Introduction

Hyperglycemia caused by diabetes mellitus is associated with long-term diabetes-related complications, resulting in reduced life expectancy when compared to the general population without diabetes [1]. In type 1 diabetes mellitus (T1DM), increased mortality is explained by diabetic ketoacidosis and hyperglycemia early in life and cardiovascular diseases later in life. Strict glucose control is associated with a lower risk of diabetes-related complications and cardiovascular mortality [2, 3]. Besides sustained chronic hyperglycemia, another particular atherogenic action of postprandial glucose has emerged as another target to be pursued in the clinical practice aimed at reducing mean blood glucose and glycated hemoglobin (HbA1c) [4]. Lower postprandial glucose levels may be associated with a lower risk of cardiovascular outcomes in diabetes [5].

However, strict glucose control is associated with weight gain and a higher incidence of hypoglycemia [2]. Hypoglycemia can lead to seizures, cognitive impairment, decreased quality of life, loss of work productivity, impaired functioning on the following day, and non-adherence to treatment [6–10]. Hypoglycemia may also cause cardiac ischemia or arrhythmia mediated by the catecholamine secretion [11], eventually leading to a higher risk of death [12].

The development of insulin analogues through molecular structure modifications of human insulin is based on pharmacokinetic profiles that try to mimic the physiological secretion of insulin [13]. Short-acting insulin analogues (aspart, glulisine, and lispro) are thought to be better than regular human insulin due to faster absorption and faster onset of action, mimicking better the physiological prandial insulin peak of people without diabetes [14, 15] and leading to lower postprandial glucose levels [16]. Potentially, this allows for better glucose control, reduces the number of hypoglycemic episodes, and helps improve the patient’s quality of life by allowing for less restrictive mealtimes. However, a number of meta-analyses on short-acting insulin analogues have found only modest benefits on glucose control and the frequency of hypoglycemic episodes compared to therapy with regular human insulin [17–19].

Methods

This systematic review was carried out based on the methodology described in the Cochrane Collaboration tool [21].

Eligibility criteria

Studies were eligible for inclusion if they were randomized controlled trials (RCTs), included children and adults with a diagnosis of T1DM for at least 1 year, with or without chronic complications, and compared the use of subcutaneous short-acting insulin analogues (aspart, glulisine, and lispro) with regular human insulin for at least 4 weeks. The primary outcomes were hypoglycemia (all hypoglycemic episodes, nocturnal hypoglycemia, and severe hypoglycemia) and postprandial glucose (all meals and after breakfast, lunch, and dinner). Secondary outcomes included long-term glucose control assessed by HbA1c and changes in quality of life. Studies with pregnant women, patients with acute diabetic decompensation, or patients with type 2 diabetes, studies that used insulin pumps, experimental studies or retrospective studies, narrative reviews, letters, and congress abstracts were excluded.

Information sources

We searched the following electronic databases for studies published until August 2017: MEDLINE (via PubMed), EMBASE (via Elsevier), CENTRAL (the Cochrane Central Register of Controlled Trials), LILACS (Literatura Latino-Americana e do Caribe em Ciências da Saúde, via BVS), and DARE (Database of Abstracts of Reviews of Effects). The references cited by all the
relevant studies were hand searched. We performed an additional search for ongoing and/or unpublished studies in the US National Institute of Health Ongoing Trials Register (http://www.clinicaltrials.gov) and the International Clinical Trials Registry Platform (ICTRP - WHO). No language restrictions were applied.

Search strategy
We searched for the terms ‘diabetes mellitus, type 1’, ‘aspart’, ‘glulisine’, ‘lispro’, and related terms to obtain as many results as possible. The complete search strategies used for each database are provided as Additional file 1: Table S1.

Study selection
Duplicates were manually identified and excluded. The articles were then analyzed in two steps: firstly, two reviewers independently screened the titles and abstracts yielded by the search strategy against the inclusion and exclusion criteria; secondly, the same reviewers independently screened the full text reports and selected the articles that met the inclusion criteria. Disagreements were resolved by consensus. If no agreement could be reached, a third reviewer was consulted for arbitration. Agreement between reviewers was assessed using Cohen’s kappa coefficient. The Rayyan software (Rayyan Platform) was used for this selection process.

Data extraction and quality assessment
Two reviewers independently extracted data from each study using an extract table template, which provided the following information: title of the study; demographic characteristics; study design; intervention details; and outcomes. A third reviewer further assessed all RCTs to check for completeness of data.

In case of missing data, the authors of the studies were contacted for additional information. If the missing data could not be retrieved, the study was not included. Retrieved missing data were presented in a narrative form.

To assess the internal quality of the studies, we ran each RCT through the Cochrane Collaboration tool for assessing the risk of bias [21]. The following potential issues were assessed: random sequence generation; allocation concealment; blinding of participants and outcome assessors; blinding of outcomes; incomplete outcome data; selective reporting; and other sources of bias. For each domain, the risk of bias was rated as low, high, or unclear. The quality of evidence was assessed using the GRADEpro GDT software (GRADEpro 2014). The results were presented in a “Summary of Findings” table.

Data synthesis and analysis
Analyses were conducted using the RevMan 5.3 software. Relative risk was used as a summary measure of effect size for dichotomous outcomes, the mean difference was used for continuous outcomes, and the risk rate was used for outcomes related to the number of events. The meta-analysis was performed using a random-effects model based on the DerSimonian–Laird method, 95% confidence interval (95% CI). A p-value < 0.1 was considered statistically significant. Heterogeneity between the studies was assessed using I² statistic, in which values above 50% were indicative of high heterogeneity [21]. Heterogeneity as determined by the Chi square test was considered non-significant for I² values between 0 and 50%, moderate for values between 51 and 79%, and significant for values between 80 and 100%. Where possible, study data were pooled and summarized in meta-analysis charts (quantitative synthesis) using the RevMan 5.3 software; otherwise, the results of each study were presented individually (qualitative synthesis).

Subgroup and sensitivity analysis
Subgroup analyses were designed based on the effects by neutral protamine Hagedorn (NPH) human insulin vs. long-acting insulin analogues. We also carried out a sensitivity analysis for the primary outcome ‘all hypoglycemic episodes’ considering the risk of bias in the studies. A second meta-analysis was then performed for this outcome excluding studies with three or more domains classified as “high risk of bias”.

Results
Literature search
The electronic search yielded a total of 2897 references. We found 1761 references in MEDLINE, 670 in EMBASE, 216 in CENTRAL, 110 in ClinicalTrials.gov, and none in the International Clinical Trials Registry Platform (ICTRP-WHO). We then removed duplicates (328) and selected the studies as shown in Additional file 2: Figure S1.

Description of studies
In the initial search, 2569 potentially relevant citations were retrieved, of which 22 articles met the inclusion criteria. Eight RCTs analyzed the effects of aspart vs. regular human insulin [35–42], one analyzed the effects of glulisine vs. regular human insulin [43], and 13 analyzed the effects of lispro vs. regular human insulin [22–34]. The characteristics of each study are described in Table 1.

Most studies (77.2%) were multicenter trials, and the countries with the largest participation were the United
Table 1 Characteristics of the included studies

| Study/year                  | Design                      | N  | Age (years) | Time with DM (mean, years) | Comparison                                                                 | Basal insulin                                                                 | Treatment time | Outcomes                                                                 |
|-----------------------------|-----------------------------|----|-------------|-----------------------------|-----------------------------------------------------------------------------|-----------------------------------------------------------------------------|----------------|--------------------------------------------------------------------------|
| Anderson et al. (1997)      | Open, multicenter, crossover| 336| 32.1 (mean) | 12                          | LISPRO (right before meal) × REGULAR (30–45 min before meal)               | NPH and Long-acting insulin analogue (Humulin U)                          | 12 months      | Total hypoglycemia, Postprandial glucose, Glycated hemoglobin           |
| Annuzzi et al. (2001)*      | Multicenter, crossover      | 85 | 31 (mean)   | 12                          | LISPRO (right before meal) × REGULAR (30–45 min before meal)               | NPH                                                                         | 3 months       | Total hypoglycemia, Severe hypoglycemia, Glycated hemoglobin, Quality of life |
| Brock Jacobsen et al. (2011)| Double blind, single center, crossover | 16 | From 18 to 60 | 1                           | ASPART × REGULAR (both right before meal)                                  | NPH                                                                         | 2 months       | Total hypoglycemia, Nocturnal hypoglycemia, Postprandial glucose, Glycated hemoglobin, Quality of life |
| Cherubini et al. (2006)     | Open, single center, parallel| 30 | 8.1 (mean)  | 5.2                         | ASPART (2 min before meal) × REGULAR (30 min before meal). Counting of carbohydrates | Glargine                                                                     | 4.5 months     | Total hypoglycemia, Postprandial glucose, Glycated hemoglobin            |
| Danne et al. (2007)         | Open, multicenter, crossover| 26 | 5 (mean)    | 1.8                         | ASPART (right before meal) × REGULAR (30 min before meal)                  | NPH                                                                         | 3 months       | Total hypoglycemia, Severe hypoglycemia, Patient satisfaction            |
| Fairchild et al. (2000)     | Open, single center, crossover| 35 | From 5 to 10 | 3.1                        | LISPRO (right before meal) × REGULAR (30 min before meal)                  | NPH                                                                         | 3 months       | Total hypoglycemia, Nocturnal hypoglycemia, Severe hypoglycemia, Postprandial glucose, Glycated hemoglobin |
| Ferguson et al. (2001)      | Open, single center, crossover| 33 | 46 (mean)   | 26                          | LISPRO (right before meal) × REGULAR (30 min before meal)                  | NPH                                                                         | 4.5 months     | Total hypoglycemia, Nocturnal hypoglycemia, Severe hypoglycemia, Postprandial glucose, Glycated hemoglobin, Quality of life |
| Study/year | Design | N   | Age (years) | Time with DM (mean, years) | Comparison | Basal insulin | Treatment time | Outcomes                                      |
|-----------|--------|-----|-------------|---------------------------|------------|---------------|----------------|----------------------------------------------|
| Ford-Adams et al. (2003) | Multicenter, crossover | 23  | From 7 to 11 | Not provided | LISPRO x REGULAR | NPH | 4 months | Total hypoglycemia, Nocturnal hypoglycemia, Glycated hemoglobin |
| Gale et al. (2000) | Double blind, multicenter, crossover | 93  | 35 (median) | 13 (median) | LISPRO x REGULAR (both right before meal) | NPH | 3 months | Total hypoglycemia, Nocturnal hypoglycemia, Severe hypoglycemia, Postprandial glucose, Glycated hemoglobin, Quality of life |
| Garg et al. (2005) | Open, multicenter, parallel | 860 | 40.3 (mean) | 20 | GLULISINE (0-15 min before meal) x REGULAR (30–45 min before meal) | Glargine | 3 months | Total hypoglycemia, Severe hypoglycemia, Postprandial glucose, Glycated hemoglobin |
| Heller et al. (1999) | Open, multicenter, crossover | 165 | 38 (mean) | 16 | LISPRO (right before meal) x REGULAR (30 min before meal) | NPH | 4 months | Total hypoglycemia, Nocturnal hypoglycemia, Severe hypoglycemia, Postprandial glucose, Glycated hemoglobin |
| Heller et al. (2004) | Double blind, multicenter, crossover | 155 | 35.7 (mean) | 2 | ASPART x REGULAR (both right before meal) | NPH | 4 months | Total hypoglycemia, Severe hypoglycemia, Nocturnal hypoglycemia, Glycated hemoglobin |
| Holcombe et al. (2002) | Open, multicenter, crossover | 463 | 14.9 (mean) | 6 | LISPRO (right before meal) x REGULAR (30–45 min before meal) | NPH | 4 months | Total hypoglycemia, Nocturnal hypoglycemia, Severe hypoglycemia, Postprandial glucose, Glycated hemoglobin, Adverse event |
| Study/year                                      | Design               | N    | Age (years) | Time with DM (mean, years) | Comparison                                                                 | Basal insulin | Treatment time | Outcomes                                      |
|------------------------------------------------|----------------------|------|-------------|-----------------------------|---------------------------------------------------------------------------|---------------|----------------|-----------------------------------------------|
| Hollenman et al. (1997)                        | Open, multicenter, crossover | 199  | 35.4 (mean) | 13                          | LISPRO (right before meal) x REGULAR (30 min before meal)                    | NPH           | 3 months       | Total hypoglycemia Nocturnal hypoglycemia    |
| Home et al. (1998)                             | Multicenter, crossover | 104  | 34 (mean)   | 15                          | ASPART x REGULAR (both right before meal)                                  | NPH           | 1 month        | Total hypoglycemia Severe hypoglycemia       |
| Home et al. (2000), Bott et al. (2003), Home et al. (2006) | Open, multicenter, parallel | 1070 | 38 (mean)   | 15                          | ASPART (right before meal) x REGULAR (30 min before meal)                  | NPH           | 6 months       | Total hypoglycemia Severe hypoglycemia       |
| Jacobs et al. (1997)                           | Open, multicenter, crossover | 12   | 18 (mean)   | Not provided                | LISPRO (right before meal) x REGULAR (30 min before meal)                  | NPH           | 1 month        | Total hypoglycemia Nocturnal hypoglycemia    |
| Provenzano et al. (2001)                       | Open, multicenter, crossover | 12   | 28 (mean)   | 12                          | LISPRO x REGULAR (both right before meal)                                  | Long-acting insulin | 168 days   | Total hypoglycemia Nocturnal hypoglycemia    |
| Raskin et al. (2000)                           | Open, multicenter, parallel | 882  | 39.2 (mean) | 1.5                         | ASPART (right before meal) x REGULAR (30 min before meal)                  | NPH           | 6 months       | Total hypoglycemia Severe hypoglycemia       |
| Tamas et al. (2001)                            | Open, multicenter, parallel | 423  | From 18 to 70 | 14                          | ASPART (0 to 5 min before meal) x REGULAR (30 min before meal)              | NPH           | 16 months      | Total hypoglycemia Severe hypoglycemia       |
States, United Kingdom, Australia, Italy, Germany, Canada, Denmark, and Finland. All studies were published between 1996 and 2011.

The selected studies contributed to a combined sample of 6235 patients for the meta-analysis. Sample sizes varied between the studies, with a minimum of 12 [23, 30] and a maximum of 1184 patients [32]. The mean age of the participants ranged from 5 to 60 years, with five studies including only children and adolescents [26, 31, 33, 34, 39]. The time since the diagnosis of T1DM ranged from 1 to 20 years.

NPH insulin was the most widely used type of basal insulin. Duration of treatment ranged between 1 and 16 months. RCTs aimed at assessing metabolic stabilization and drug adaptation carried out a run-in period that lasted up to 2 months. In most articles, short-acting insulin analogues were administered immediately before meals, and regular human insulin was administered 30–45 min before meals.

Assessment of the risk of bias
The risk of bias was assessed using the Cochrane Collaboration tool [21]. The domains with the highest risk of bias were the lack of patient and research team blinding for the treatments and the assessment of subjective outcomes. Results of the assessment and the percentage distribution of the risk of bias by domain are shown in Additional file 3: Figure S2, Additional file 4: Table S2 and Additional file 6: Table S3.

Hypoglycemia
All hypoglycemic episodes
All 22 studies included in this meta-analysis had information on the number of all hypoglycemic episodes per month and were thus included in our count [22–43]. Short-acting insulin analogues (aspart, glulisine, and lispro) were not associated with a lower number of all hypoglycemic episodes per month when compared to regular human insulin (risk rate 0.94, 95% CI 0.89–1.00; 6235 patients; I² = 80%).

A sensitivity analysis was performed for this primary outcome (total episodes of hypoglycemia) by excluding studies with a high risk of bias [31, 41]. The results showed that short-acting insulin analogues were associated with a lower number of total hypoglycemic episodes per month when compared with regular human insulin (risk rate 0.93, 95% CI 0.87–0.99; 6180 patients, 20 studies; I² = 81%) (Fig. 1). The monthly rate of total hypoglycemic episodes after the use of short-acting insulin analogues was 7% lower than in the group who used regular human insulin.

A subgroup analysis comparing the use of NPH and long-acting insulin analogues as basal insulin did not show difference between them regarding the number of total hypoglycemic episodes per month when compared with regular human insulin (risk rate 0.93, 95% CI 0.87–0.99; 6180 patients, 20 studies; I² = 81%) (Fig. 1). The monthly rate of total hypoglycemic episodes after the use of short-acting insulin analogues was 7% lower than in the group who used regular human insulin.

Nocturnal hypoglycemia
Of the 22 studies, eight assessed episodes of nocturnal hypoglycemia, one with glulisine and seven with lispro. Their results were combined for this meta-analysis [24–28, 33, 34, 43]. In the only RCT that compared glulisine with regular human insulin, no difference was found between them (risk rate 0.93, 95% CI 0.76–1.13; 564 patients). The short-acting insulin analogues (lispro

Table 1 (continued)

| Study/year            | Design          | N   | Age (years) | Time with DM (mean, years) | Comparison  | Basal insulin | Treatment time | Outcomes                                      |
|----------------------|-----------------|-----|-------------|---------------------------|-------------|----------------|----------------|-----------------------------------------------|
| Tupola et al. (2001) | Open, multicenter, crossover | 29  | 6 (mean)    | 3                         | LISPRO (30 min after the patient started eating) × REGULAR (20 to 39 min before meal) | NPH          | 3 months       | Total hypoglycemia Glycated hemoglobin      |
| Vale et al. (2001)   | Open, multicenter, parallel | 1184 | 38.7 (mean) | 19.4                      | LISPRO (right before meal) × REGULAR (30 min before meal) | NPH          | 3 months       | Total hypoglycemia Severe hypoglycemia Postprandial glucose Glycated hemoglobin |

* Data from this study were taken from Siebenhofer et al. [51]
and glulisine) were associated with a 45% lower risk rate of nocturnal hypoglycemia when compared with regular human insulin (risk rate 0.55, 95% CI 0.40–0.76, 1995 patients, $I^2=84\%$) (Fig. 2).

A subgroup analysis comparing the use of NPH and long-acting insulin analogues as basal insulin showed that NPH insulin was associated with a lower number of nocturnal hypoglycemic episodes per month when compared with long-acting insulin analogues (NPH as basal insulin: risk rate 0.48, 95% CI 0.36–0.64, 1011 patients, 7 studies, $I^2=71\%$; glargine as basal insulin: risk rate 0.93, 95% CI 0.76–1.13, 860 patients, one study).

**Severe hypoglycemia**

Of the 22 studies, 15 analyzed the episodes of severe hypoglycemia. Their results were combined for this meta-analysis [24–28, 32, 33, 35–38, 40–43]. Short-acting insulin analogues (aspart, glulisine, and lispro) were associated with a 32% lower risk rate of severe hypoglycemia when compared with regular human insulin (risk

| Study or Subgroup | log(Rate Ratio) | SE | Weight | Rate Ratio | Rate Ratio |
|-------------------|----------------|----|--------|------------|------------|
| Anderson et al, 1997 | -0.094          | 0.0518 | 6.2%  | 0.91 [0.82, 1.01] |           |
| Annuzzi et al, 2001 | 0.2231          | 0.0893 | 4.5%  | 1.25 [1.04, 1.50] |           |
| Brock-Jacobsen et al, 2011 | -0.298          | 0.0846 | 4.9%  | 0.74 [0.63, 0.86] |           |
| Cherubini et al, 2006 | -0.173          | 0.3409 | 0.8%  | 0.84 [0.43, 1.64] |           |
| Danne et al, 2007 | 0.0488          | 0.088 | Not estimable |                   |           |
| Fairchild et al, 2000 | 0.2808          | 0.1695 | 2.4%  | 1.32 [0.95, 1.85] |           |
| Ferguson et al, 2001 | 0.0361          | 0.0419 | 6.6%  | 1.04 [0.96, 1.13] |           |
| Ford-Adams et al, 2003 | -0.0828         | 0.0586 | 5.9%  | 0.92 [0.62, 1.03] |           |
| Gale et al, 2000 | -0.1759         | 0.1535 | 2.7%  | 0.84 [0.62, 1.13] |           |
| Garg et al, 2005 | 0.0197          | 0.0451 | 6.5%  | 1.02 [0.93, 1.11] |           |
| Heller et al, 1999 | -0.3999         | 0.0464 | 6.4%  | 0.67 [0.61, 0.73] |           |
| Heller et al, 2004 | -0.0971         | 0.0346 | 6.8%  | 0.91 [0.85, 0.97] |           |
| Holcombe et al, 2002 | -0.0835         | 0.0454 | 6.4%  | 0.92 [0.84, 1.01] |           |
| Holleman et al, 1997 | -0.0414         | 0.0295 | 7.0%  | 0.96 [0.91, 1.02] |           |
| Home et al, 1998 | -0.0813          | 0.0582 | 6.6%  | 0.92 [0.82, 1.03] |           |
| Home et al, 2000 | -0.0444          | 0.0442 | 6.5%  | 1.05 [0.96, 1.14] |           |
| Jacobs et al, 1997 | -0.0305         | 0.1123 | 3.9%  | 0.97 [0.78, 1.12] |           |
| Provenzano et al, 2001 | -0.5548       | 0.1647 | 2.5%  | 0.57 [0.42, 0.78] |           |
| Raskin et al, 2000 | -0.0497         | 0.1326 | 3.3%  | 0.95 [0.73, 1.23] |           |
| Tamas et al, 2001 | 0.0156          | 0.1022 | 4.2%  | 1.02 [0.83, 1.24] |           |
| Tupola et al, 2001 | 0.1076          | 0.14 | Not estimable |                   |           |
| Valie et al, 2001 | -0.0203         | 0.0433 | 6.5%  | 0.98 [0.90, 1.07] |           |

Total (95% CI) | 100.0% | 0.93 [0.87, 0.99] | 100.0% | 0.55 [0.40, 0.76] | 100.0% | 1.00 [0.80, 1.23] |

Heterogeneity: Tau² = 0.01; Chi² = 102.10, df = 19 (P < 0.0001); P = 81% | Heterogeneity: Tau² = 0.12; Chi² = 44.88, df = 7 (P < 0.0001); P = 84% | Heterogeneity: Tau² = 0.39; Chi² = 23.60, df = 5 (P < 0.0001); P = 71% |

Test for overall effect: Z = 2.21 (P = 0.03) | Test for overall effect: Z = 7.21 (P = 0.0001) | Test for overall effect: Z = 4.03 (P = 0.0001) |

**Fig. 1** Forest plot representing all hypoglycemic episodes (for aspart, glulisine and lispro). SAI Short-acting insulin, RHI Regular human insulin

**Fig. 2** Forest plot representing nocturnal hypoglycemia (for aspart, glulisine and lispro). SAI Short-acting insulin, RHI Regular human insulin
rate 0.68, 95% CI 0.60–0.77; 5945 patients, 15 studies; \(I^2 = 0\%\) (Fig. 3).

A subgroup analysis comparing the use of NPH or long-acting insulin analogues as basal insulin showed that NPH insulin was not associated with a lower number of total hypoglycemic episodes per month when compared with long-acting insulin analogues (NPH as basal insulin: risk rate 0.70, 95% CI = 0.61–0.79, 4848 patients, 14 studies, \(I^2 = 0\%\); glargine as basal insulin: risk rate 0.40, 95% CI 0.21–0.73, 860 patients, one study).

### Postprandial glucose

Of the 22 studies, 15 analyzed postprandial glucose (any meal). Their results were combined for this meta-analysis [22–26, 28, 29, 32, 33, 35, 36, 39, 40, 42, 43]. Short-acting insulin analogues (aspart, glulisine, and lispro) were associated with lower postprandial glucose levels when compared with regular human insulin (mean difference/MD = -19.44 mg/dL; 95% CI = -21.49 to -17.39; 5031 patients, \(I^2 = 69\%\)) (Fig. 4).

Thirteen studies assessed postprandial glucose levels 2 h after breakfast [22–26, 28, 29, 32, 33, 36, 40, 42,
43]. We were able to pool the data from 12 studies in this meta-analysis, and the results showed that short-acting insulin analogues (aspart, glulisine, and lispro) were associated with lower postprandial glucose levels after breakfast when compared with regular human insulin (MD \(-22.35\) mg/dL; 95% CI \(-23.52\) to \(-21.17\); 4623 patients; I\(^2\) = 50%) (Additional file 5: Figure S3A).

Fourteen studies assessed postprandial glucose levels two hours after lunch. We were able to pool the data from 11 studies in this meta-analysis [23–26, 28, 29, 32, 33, 35, 36, 42], and the results showed that short-acting insulin analogues (aspart and lispro) were associated with lower postprandial glucose levels after lunch when compared with regular human insulin (MD \(-10.86\) mg/dL, 95% CI \(-13.41\) to \(-8.31\); 3675 patients; I\(^2\) = 54%). The three remaining studies [27, 30, 43] did not provide sufficient data to be pooled in this meta-analysis. Individually, none of the articles showed any difference when comparing insulin lispro with regular human insulin for postprandial glucose after lunch (Additional file 5: Figure S3B).

Fourteen studies assessed postprandial glucose levels 2 h after dinner [23, 24, 26–29, 31–33, 35, 36, 40, 42, 43]. We were able to pool the data from 12 studies in the meta-analysis, and the results showed that short-acting insulin analogues (aspart, glulisine, and lispro) were associated with lower postprandial glucose levels after dinner when compared with regular human insulin (MD \(-19.52\) mg/dL, 95% CI \(-21.73\) to \(-17.31\); 4530 patients; I\(^2\) = 90%). In the two studies not included in this meta-analysis, no difference was found between lispro and regular human insulin [27, 31]. (Additional file 5: Figure S3C).

**Glycated hemoglobin**

Nineteen studies evaluated HbA1c at the end of treatment [22–28, 30, 32–40, 42, 43]. We were able to pool the data from 15 studies in this meta-analysis, and the results showed that short-acting insulin analogues (aspart or lispro) were associated with lower HbA1c when compared with regular human insulin (MD \(-0.13\), 95% CI \(-0.16\) to \(-0.10\); 5204 patients; I\(^2\) = 73%).

When short-acting insulin analogues were assessed separately, only aspart was associated with lower HbA1c when compared with regular human insulin (MD \(-0.14\), 95% CI \(-0.20\) to \(-0.02\); 2822 patients, I\(^2\) = 40%). Lispro was not associated with lower HbA1c levels when compared with regular human insulin (MD \(-0.09\), 95% CI \(-0.17\) to \(-0.02\); 2552 patients; I\(^2\) = 40%).

Four studies were not included in the meta-analysis because they did not have available data for input [23, 31, 42, 43]. The first three showed no statistically significant difference between the groups, and Jacobs et al. [23] showed a difference in favor of regular human insulin.

**Quality of life and patient satisfaction**

Five studies assessed patients’ quality of life [24, 27, 28, 36, 40]. Fergusson et al. and Gale et al. [27, 28] did not present the results, but reported no difference between the short-acting insulin analogue and regular human insulin groups. Holleman et al. [24] reported a greater flexibility in the short-acting insulin analogue group (p < 0.0001) and an even better adaptation of mealtimes (p < 0.0001), physical activity planning (p < 0.0001), and activities (p < 0.0001). Home et al. [36] used the Diabetes Treatment Satisfaction Questionnaire (DTSQ) and found a significant difference in favor of the short-acting insulin analogue group, with a score 2.3 points higher than that of the regular insulin group (95% CI 1.32 points to 3.28 points). Tamás et al. [40] also reported no difference in the overall DTSQ score, but the group that received short-acting insulin analogues reported greater flexibility of use (p = 0.022).

**Discussion**

In this systematic review and meta-analysis we originally report clinical evidence on therapeutical use of short-acting insulin analogues compared with regular insulin while focusing on the main benefits of these analogues, namely the reduction of hypoglycemia and postprandial glucose levels. The combined data of 22 RCTs showed that short-acting insulin analogues are associated with a decrease in total hypoglycemic episodes, nocturnal and severe hypoglycemia, and post-breakfast, post-lunch and post-dinner glucose levels.

Fullerton et al., in a systematic review that aimed to evaluate long-term safety of short-acting insulin analogues, also assessed the outcomes described here; however, since their research was focused on long-term studies, fewer RCTs were retrieved when compared to the present study [17]. The results of these reviews cannot be compared due to the high probability of inconsistencies. Another recently-published review analyzed only trials comparing aspart with regular human insulin, but also gathered data from a smaller set of studies [44]. Since the three short-acting insulin analogues are very pharmacologically similar regarding time of onset, peak activity, and duration of action [16], analyzing data from trials conducted with only one short-acting insulin analogue yields a lower number of studies, resulting in less statistical power. We saw no clear advantage in using this approach for the current study. A third systematic review and meta-analysis described only the results of hypoglycemia, and again included a smaller
Limited evidence analyzed in this systematic review suggests that, for patients with T1DM, the treatment with short-acting insulin analogues is more convenient than with regular human insulin. The higher satisfaction levels and greater flexibility attributed to short-acting insulin analogues could be explained by the fact that they can be administered immediately before meals, as opposed to the anticipated 30 to 45 min when administering regular human insulin. In a study involving 1184 patients with T1DM, adherence to the correct timing of regular human insulin was 7% for patients who took it more than 30 min before meals, 60% for those who took it 15-30 min before meals, and 33% for those who took it 15 min before meals. Regarding the administration of insulin lispro, 98% of the patients followed the orientation (0 to 15 min before meals) [32]. The possibility of administration of short-acting insulin analogues immediately after meals is another important benefit, as it may not always be possible to predict how much food (carbohydrates) the patient will have eaten at the end of the meal.

Due to the scarcity of studies assessing the impact of short-acting insulin analogues on the quality of life of patients with T1DM and the methodologies used, previously-published systematic reviews either did not analyze this outcome or did not reach a conclusion [50]. According to Fullerton et al. [17], with a more adequate methodology (the DTSQ) [51], three studies reported no improvement in treatment satisfaction, while four studies indicated an improvement in this outcome with short-acting insulin analogues when compared to regular human insulin.

As opposed to other meta-analyses [17], this review provided information regarding the use of insulin analogues in children. However, no association was found between the use of short-acting insulin analogues or regular human insulin and the number of hypoglycemic episodes, postprandial glucose reductions, and HbA1c, probably because of the low number of studies included.

The main methodological strengths of this review are as follows: the most adequate outcomes considering the pharmacokinetics of short-acting insulin analogues; the most comprehensive and systematic literature search among systematic reviews on this subject, with no language restriction; and the specific and reproducible eligibility criteria, study selection, and data extraction.

However, some limitations should be pointed out. The first is that most studies included in our systematic review may not represent current T1DM treatment practice. Most trials excluded patients with hypoglycemia unawareness or with a high risk of hypoglycemia, which in fact makes up the largest population group that could benefit from insulin analogues in the current...
clinical practice. Additionally, the low quality of most studies identified in this systematic review may limit the interpretation of the presented data. The differences in the definition of total and nocturnal hypoglycemic episodes, as well as the methods for recording hypoglycemic episodes based on the presence of symptoms or on the obligatory verification of blood glucose independently of symptoms, are real limitations frequently observed in clinical trials. Another limitation is the absence of masking, which could also result in a high risk of bias. However, it is unlikely that future studies will adequately mask the participants, as this would require a significant increase in the number of insulin applications. The analyses with NPH as basal insulin included 7, 18 and 14 studies (nocturnal hypoglycemia, total hypoglycemia and severe hypoglycemia, respectively), and those with long-acting insulin analogues included 1, 3 and 1 studies (nocturnal hypoglycemia, total hypoglycemia and severe hypoglycemia, respectively), and these analyses presented high heterogeneity, precluding their consideration as a definitive evidence of the possible superiority of NPH as compared to long-acting insulin analogues. This information should, thus, be interpreted with caution. A direct comparison between NPH insulin and long-acting analogues is beyond the scope of this review. Another important point is that, over the years, there has been a significant evolution in insulin therapy, which can be observed in the clinical heterogeneity between studies in the past 20 years.

In summary, short-acting insulin analogues were associated with fewer nocturnal and severe hypoglycemic events and better glucose control (slightly lower HbA1c and lower postprandial blood glucose levels) when compared with regular human insulin in subjects with type 1 diabetes.

Additional files

Additional file 1: Table S1. Search strategy applied to all databases.

Additional file 2: Figure S1. Flow diagram: identification and selection of articles included in the meta-analysis.

Additional file 3: Figure S2. Percentage distribution of risk of bias by domain.

Additional file 4: Table S3. Characteristics and Quality Of Studies.

Additional file 5: Figure S3. Forest plot representing postprandial glucose for breakfast (A), lunch (B), and dinner (C) (for aspart, glulisine and lispro). SAI: Short-Acting Insulin; RHI: Regular Human Insulin.

Additional file 6: Table S3. Analysis of risk of bias of the selected studies.

Authors’ contributions
Conception and design: KFSM, LRB, BDS; data search: KFSM, GJMP, ALM, RR; analysis and interpretation of data: KFSM, LRB, BP, BDS; drafting of the manuscript: KFSM, LRB, BDS; revising it critically for important intellectual content: LEPC, WJM, LAT, HCP; final approval of the manuscript submitted: KFSM, LRB, GJMP, ALM, RR. All authors read and approved the final manuscript.

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Competing interests
The authors declare that they have no competing interests.

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Not applicable.

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KFSM had full access to the data and takes full responsibility for its integrity.

Ethics approval and consent to participate
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