

This is the author's manuscript



AperTO - Archivio Istituzionale Open Access dell'Università di Torino

Efficacy and safety of dual SGLT 1/2 inhibitor sotagliflozin in type 1 diabetes: meta-analysis of randomised controlled trials

Original Citation:	
Availability:	
This version is available http://hdl.handle.net/2318/1705347	since 2019-07-03T11:53:09Z
Published version:	
DOI:10.1136/bmj.I1328	
Terms of use:	
Open Access Anyone can freely access the full text of works made available as under a Creative Commons license can be used according to the tof all other works requires consent of the right holder (author or protection by the applicable law.	erms and conditions of said license. Use

(Article begins on next page)



UNIVERSITÀ DEGLI STUDI DI TORINO

This is an author version of the contribution published on:

Questa è la versione dell'autore dell'opera:

[British Medical Journal, 365:I1328, 2019, doi: 10.1136/bmj.l1328]

The definitive version is available at:

La versione definitiva è disponibile alla URL:

https://www.bmj.com/content/365/bmj.l1328.long

http://creativecommons.org/licenses/by-nc-nd/4.0/

Efficacy and safety of dual SGLT 1/2 inhibitor sotagliflozin in type 1 diabetes: metaanalysis of randomised controlled trials

RUNNING TITLE: meta-analysis of sotagliflozin in type 1 diabetes:

Giovanni Musso¹MD, Roberto Gambino² PhD, Maurizio Cassader² PhD, Elena

Paschetta¹MD.

1 HUMANITAS Gradenigo, Turin

2 Laboratoty of Diabetes and Metabolic disorders, Department of Medical Sciences, University of Turin, Italy

Corresponding author:

Giovanni Musso

HUMANITAS Gradenigo

C. so Regina Margherita 8

10132 Turin, Italy

E-mail: giovanni_musso@yahoo.it

Word count: 4812

KEY POINTS

QUESTION

• What is the efficacy and safety of the novel dual dual sodium glucose co-transport ½(SGLT1/2 inhibitor sotagliflozin in patients with type 1 diabetes mellitus (T1DM)?

FINDINGS

- The dual SGLT1/2 inhibitor sotagliflozin improves glycemic and nonglycemic outcomes and reduces the incidence of hypoglycemia and of severe hypoglycemia in T1DM.
- Diabetic ketoacidosis (DKA) is the main adverse event associated with sotagliflozin treatment.
 The risk of DKA varies depending on initial HbA1c levels and basal insulin dose reduction during treatment. An increased risk of genital tract infections and diarrhea, but not of urinary tract infections, is also associated with sotagliflozin.

MEANING

- Sotagliflozin has incremental benefit over other adjunctive therapies, including incretin analogues and SGLT2 inhibitors, seeking an indication as an adjunct therapy to insulin in T1D.
- Careful patient selection and insulin dose adjustment may help minimize the risk of DKA associated with sotagliflozin treatment

Abstract

Background. Patients with type 1 diabetes mellitus (T1DM) achieve target glycemic control in 30% of cases and are encumbered with hypoglycemia, the main factor limiting optimal glucose control and a strong predictor of adverse outcomes and death. Hence, these patients urgently need adjunctive therapies to insulin.

Purpose. To assess efficacy and safety of the first-in-class dual sodium glucose co-transport 1/2 inhibitor sotagliflozin in T1DM.

Data sources. MEDLINE, Cochrane Library, EMBASE, International meeting abstracts, international and national clinical trial registries, websites of US, European and Japanese regulatory authorities, through Jan 10th, 2019.

Study Selection: Randomized controlled trial s(RCTs) evaluating the effect of sotagliflozin vs. active comparison or placebo on glycemic and nonglycemic outcomes and on adverse events in T1DM.

Data Extraction. Three reviewers extracted data for study characteristics, outcomes of interest, and risk of bias and summarized strength of evidence using the Grading of Recommendations Assessment, Development and Evaluation approach. Main outcomes were pooled using random-effects model.

Data Synthesis. Among 739 records identified, 6 placebo-controlled RCTs (3238 participants, duration ranging 4-52 weeks) were included. Sotagliflozin reduced HbA1c (WMD:-0.34%[95%CI:-0.41,-0.27], p<0.00001), fasting (WMD:-16.5 mg/dL [-22.1,-10.9] and 2h-postprandial plasma glucose (WMD:-39.2 mg/dL [-50.7, -27.6], and daily total (WMD:-8.99% [-10.93, -7.05]), basal (WMD:-8.03% [-10.14, -5.93]) and bolus (WMD:-9.14%[-12.17, -6.12]) insulin dose. Sotagliflozin improved time-in-range (WMD:+9.73%[6.66, 12.81]) and other continuous glucose monitoring parameters, and reduced body weight(WMD:-3.54% [-3.98,-3.09]), systolic BP(WMD:-3.85 mmHg[-4.76, -2.93]) and albuminuria

(WMD:-14.65 mg/g [-26.72,-2.58]).

Notably, sotagliflozin reduced hypoglycaemia (WMD:-9.09 events per patient-year [-13.82, -4.36]), and severe hypoglycaemia (RR: 0.69[0.49, 0.98]), but increased the risk of ketoacidosis (RR: 3.93[1.94, 7.96]), genital tract infections (RR: 3.12[2.14, 4.54]) diarrhea (RR: 1.50[1.08, 2.10]) and volume depletion events (RR: 2.19[1.10, 4.36]). Initial HbA1c and basal insulin dose adjustment were associated with the risk of DKA. Sotagliflozin 400 mg was more effective that the 200 mg dose for most glycemic and nonglycemic outcomes, but not for adverse events. The quality of evidence was high-to-moderate for most effect and safety outcomes, but low for major adverse cardiovascular events and all-cause death.

Limitations. The relatively short duration of RCTs prevented assessment of long-term outcomes. **Conclusions.** Sotagliflozin provides substantial glycemic and nonglycemic benefits and reduces hypoglycemia in T1DM, Strategies to minimize to risk of DKA and long-term effect on hard outcomes in T1DM patients receiving sotagliflozin warrant future assessment.

KEY-WORDS: sodium glucose co-transport-1/2 (SGLT1/2) inhibitors, LX4211, diabetes treatment, SGLT1, DKA

ABBREVIATIONS

ADA: American Diabetes Association; BP: blood pressure; DKA: diabetic ketoacidosis; EASD: European Association for the Study of Diabetes; EOT: end of treatment; FPG: fasting plasma glucose; LDL: low density lipoprotein; HDL: high density lipoprotein; GTI: genital tract infection. PRISMA: Preferred Reporting Items for Systematic Reviews and Meta-Analyses; RCT: randomized controlled trial; SGLT: sodium glucose co-transporter; T1D: type 1 diabetes mellitus; TID: daily total insulin dose; UTI: urinary tract infection; WMD: weighted mean difference

Introduction

Type 1 diabetes mellitus (T1DM) affects 1.5 million people in the U.S. alone and its prevalence is continuously rising, partly because over 10% of patients initially presumed to have type 2 diabetes (T2DM) at diagnosis subsequently show evidence of islet autoimmunity and progress to insulin dependence in the following years^{1,2}.

The achievement and maintenance of glycemic goals in T1DM proved both difficult and hazardous: in the T1DM Exchange clinic registry the average HbA1c was 8%, only 30% of T1D patients achieved a goal HbA1c of 7% and severe hypoglycemia occurred in up to 20% of patients per-year³; similarly, in the Diabetes Complications and Control Trial (DCCT), patients with T1DM with HbA1c levels within target showed a 2.9-fold increased cardiovascular mortality⁴ and the T1DM patients in the intensive intervention group escalated back to an HbA1c of 8% in the post-trial years⁵.

Insulin is the mainstay of T1DM treatment, but has unwanted effects, including hypoglycemia and weight gain⁶. Severe hypoglycemia in particular is the main factor limiting optimal glucose control in T1DM, is frequent, adds costs to diabetes management, and is a strong predictor of adverse vascular and nonvascular outcomes and death^{7,7,8,9}.

None of the adjunctive therapies approved (i.e., pramlintide) or recently proposed for T1DM [i.e., metformin, incretin analogues, sodium-glucose cotransporter (SGLT)2 inhibitors] has reduced the incidence of hypoglycemia and severe hypoglycemia, which remain the major unsolved issue in the management of these patients ^{10,11,12,13,14,15,16,17,18,19,20}.

SGLT1 is responsible for glucose absorption in the proximal intestine and missense mutations in SGLT1 gene were associated with protection from glucose intolerance, obesity and cardiometabolic risk in population-based studies²¹.

Sotagliflozin (LX4211, SAR439954) is a novel first-in-class dual inhibitor of sodium-glucose cotransporter (SGLT)1 and of SGLT2 (SGLT1/2 inhibitor): while SGLT2 inhibition reduces renal tubule glucose reabsorption, SGLT1 inhibition decreases intestinal glucose absorption. This peculiar dual

mechanism of action may offer incremental benefits over selective SGLT2 inhibitors²² by blunting postprandial glycemic excursions and glycemic variability, lowering the need for bolus insulin correction doses, and eventually reducing hypoglycemic risk²³.

Furthermore, reduced glucose absorption in the proximal intestine increases glucose delivery to the distal intestine, stimulating—incretin glucagon-like peptide 1 (GLP-1)²⁴. In preclinical models, the increased incretin release enhanced weight loss and counteracted—glucagon-induced ketogenesis²⁵, which may reduce the risk of diabetic ketoacidosis (DKA)^{23,24,25}.

Sotagliflozin has recently reached phase 3 development in T1D^{26, 27,28, 29, 30,31} but RCTs evaluating this drug have not been systematically reviewed. To clarify the evidence base of this novel approach, we conducted a meta-analysis of randomized controlled trials (RCTs) evaluating the efficacy and safety of sotagliflozin in adults with T1D.

METHODS

Data Sources and Searches

We searched English and non-English language publications up to January 10th 2019 on the following databases and international and national clinical trial registries: Ovid MEDLINE, Ovid MEDLINE Epub Ahead of Print, Ovid MEDLINE In-Process, EMBASE, Cochrane Database of Systematic Reviews, Epistemonikos, ClinicalTrials.gov, Cochrane CENTRAL Register of Controlled Trials; World Health Organization International Clinical Trials Registry Platform, European Union (EU) Clinical Trials Register, International Standard Randomised Controlled Trial Number (ISRCTN) registry, Australian New Zealand Clinical Trials Registry, and 19 national clinical trial registries (the full list of clinical trial registries is provided in **Supplementary text**). No language restrictions were applied. We also searched the US Food and Drug Administration³², European Medicines Agency³³ and Japanese Pharmaceutical and Medical Devices Agency³⁴ sites and drug manifacturers' websites^{35,36} for relevant documents, and the American Diabetes Association (ADA) and /European Association for the Study of Diabetes (EASD) meeting abstracts, which were subjected to the same assessment as regular articles.

We also contacted by e-mail authors of relevant papers to verify results and methodological quality of retrieved articles and drug manifacturers to inquire about further published and unpublished trials. Additionally, we manually scanned reference lists from trials, review articles and reports to identify any other relevant data.

Search terms: sodium glucose co-transport 1/2 inhibitors, dual sodium-glucose transport inhibitors, SGLT1/2 inhibitors, SGLT1 inhibitors, SGLT1 inhibitors, SGLT1/2 inhibitor, sotagliflozin, LX4211, LP802034, SAR439954, Zynquista, management, therapy, treatment, trial, diabetes, type 1 diabetes (examples of online strategy run are provided in **supplementary text**).

Study Selection

Inclusion criteria: English and non-English (French, Spanish, Portuguese, German, Chinese, Japanese, Korean) articles reporting RCTs with participants aged>18 yrs, of any sex or ethnic origin, comparing sotagliflozin with placebo or active comparators as adjunct therapy to insulin in T1DM.

Exclusion criteria were: non-human studies, non-randomized trials, letters/case reports, articles not reporting outcomes of interest or primary data (editorials, reviews).

Outcome measures

We grouped evaluated outcomes into three broad sets: glycemic efficacy outcomes, non-glycemic outcomes, and safety outcomes.

Glycemic efficacy outcomes were:

- -hemoglobin A1c (<u>HbA1c</u>) changes from baseline (primary outcome)
- -changes in fasting plasma glucose (FPG) levels.
- -changes in <u>2-hour postprandial glycemia (2h-PPG)</u> as measured during an Oral Glucose Tolerance Test (OGTT) or a standardized Mixed Meal Tolerance Test (MTT), as numerous studies link postprandial glucose excursions to the risk of cardiovascular disease (CVD) and report that targeting PPG rather than FPG lowers cardiovascular risk^{37,38}.
- -changes in <u>total</u>, <u>basal</u>, <u>and bolus insulin dose</u>, expressed as % initial insulin dose
- -urinary glucose excretion: we also assessed the effect of SGLT-1/2 inhibitors on daily urinary glucose

excretion.

-continuous glucose monitoring (CGM) parameters: CGM monitoring provides additional information to HbA1c and has been recently recommended for all adult patients with T1D and approved by the Food and Drug Administration (FDA) Advisory Committee³⁹ We therefore assessed the following CGM metrics (described in **supplementary text**): time-in-range (%), average daily glucose, standard deviation (SD) around average daily glucose, mean amplitude of glucose excursion (MAGE)⁴⁰.

.

Non-glycemic outcomes

Non-glycemic outcome measures evaluated were: changes in body weight, systolic and diastolic blood pressure (BP); renal outcomes, defined as changes in estimated glomerular filtration rate (eGFR) and in albuminuria (expressed as urinary albumin/creatinine ratio, ACR), or need for renal replacement therapy; and changes in plasma lipids [triglyceride, low density (LDL)- and high density m(HDL)-cholesterol].

Safety outcomes

Safety measures, were severe hypoglycaemia and any hypoglycaemia, diabetic ketoacidosis (DKA) (definitions provided in **supplementary text**), urinary tract infections (UTIs), genital tract infections (GTIs), other infections; gastrointestinal symptoms, major adverse cardiovascular events (MACE) (cardiovascular death, myocardial infarction, stroke, hospitalization due to heart failure or unstable angina, or coronary revascularization), cancer (overall and type-specific); amputation; bone fracture, volume depletion, renal events, acidosis-related events, drug-induced liver injury, venous thromboembolism, serious adverse events (AEs), AEs leading to treatment discontinuation, all-cause mortality.

Volume depletion, acidosis-related events, renal events and serious AEs were defined according to the Medical Dictionary for Regulatory Activities (MedDRA) preferred items version 14.0⁴¹(supplementary text).

For DKA, we planned to investigate whether the risk of DKA varied across different modes of insulin delivery, i.e. multiple daily injections (MDI) or continuous subcutaneous infusion (CSI).

All measures of dispersion were converted to standard deviations (SDs).

Data extraction and Risk-of-Bias assessment. Two reviewers (GM, RG) extracted data independently and in duplicate by using a predesigned data collection form, based on the Cochrane Handbook for Systematic Reviews of Intervention; discrepancies were arbitrated by a third reviewer and resolved by consensus. The agreement between the 2 reviewers for selection and validity assessment of trials was scored by Kappa coefficient.

The quality of RCTs was assessed by the Cochrane Collaboration Risk-of-Bias Tool⁴². We also assessed sponsorship bias, which we included in the Risk-of-Bias tool. The 2018 Agency for Healthcare Research and Quality (AHRQ) recommendations caution against equating industry sponsorship with high risk of bias and automatically downgrading the evidence for industry sponsorship⁴³. Therefore, for all included trials we systematically assessed a pre-specified list of eight items in trial designing, conducting and reporting, which have been empirically linked to the risk of biased outcomes in industry-funded trials and are not captured by the six domains of the RoB tool^{44,45,46,47,48,49,50} (supplementary Table 1).

Data Synthesis, Analysis and Grading of Evidence. The analysis was carried out in concordance with the Cochrane Handbook of Systematic Reviews of Interventions⁴² using Stata, release 11.2 (StataCorp, College Station, Texas) and RevMan Version 5.3.5(Nordic Cochrane Center, Copenhagen, Denmark⁵¹ and was reported according to PRISMA guidelines⁵²(see **supplementary Appendix**). Treatments were evaluated on an intention-to-treat principle.

We calculated weighted mean differences (WMDs) and 95% CIs for continuous outcomes using an inverse variance random-effects model. For dichotomous outcomes, we calculated Risk Ratios (RRs) and 95% CIs by using the random-effects Mantel–Haenszel approach with significance set at P=0.05. We conservatively used *a priori* a random-effects model assuming a susbtantial variability in treatment effect size across studies.

Statistical heterogeneity was assessed using the I^2 statistic: with I^2 values $\geq 50\%$, we planned to explore

individual study characteristics and those of subgroups of the main body of evidence⁵³.

We planned to conduct sensitivity analyses by repeating the analysis with alternative effect measures (odds ratio vs. relative risk), pooling methods (Peto vs. Mantel-Hanszel⁵⁴), statistical models (fixed vs. random effects), by excluding RCTs where we imputed values and RCTs at high risk of bias in any domains of the RoB tool.

We also planned a priori subgroup analysis to explore potential effects on outcome measures of the following conditions: treatment duration ($\leq 12 \text{ vs.} > 12 \text{ weeks}$), initial HbA1c levels ($\geq 8\% \text{ vs.} < 8\%$), duration of diabetes ($< 20 \text{ yr vs.} \geq 20 \text{ yr}$), background therapy (pre-treatment insulin optimization vs. stable insulin therapy), presence and severity of renal dysfunction.

We explored interactions between different sotagliflozin doses and all outcomes primarily by comparing high dose to low dose arms within head-to-head trials (within-trial approach); we planned to verify robustness of this approach in ruling out dose-response relationship by using also across-trial comparison and meta-regression. Although the "across-trial" approach has a higher risk of ecological bias, it has a higher power that the within-trial approach, thus allowing ruling out dose-response interactions with higher confidence⁵⁵.

When ≥8 comparisons were available, the effect of different doses of SGLT1/2 inhibitor, of baseline HbA1c, of treatment duration and of diabetes duration on each outcome were assessed by meta-regression analysis (random effects model, within-study variance estimated with the unrestricted maximum-likelihood method).

The dose variable in the regression equation was treated categorically, with the starting dose coded as the baseline amount and each doubling of a drug dose was a single increment increase.

Publication bias was examined using funnel plots and the Egger test.

We used the Grading of Recommendations Assessment, Development and Evaluation (GRADE) approach to summarize the strength of evidence at outcome level and determine confidence in summary estimates for clinically relevant comparisons and outcomes ^{56,57}. Three reviewers graded inconsistency, risk of bias, indirectness, imprecision, and publication bias for evidence related to the following areas: glycemic efficacy (outcomes: HbA1c, FPG, 2h-PPG, time-in-range), nonglycemic efficacy (outcomes: body weight, sys BP, eGFR, albuminuria), and adverse events (outcomes: hypoglycemia, severe hypoglycemia, DKA, urinary and genital tract infections, diarrhea, MACE, serious AEs, AEs leading to discontinuation, mortality).

Management of missing data.

We planned to manage missing data by contacting via e-mail the corresponding authors. Where this was unsuccessful, we planned to follow the approach described in Cochrane Handbook of Systematic Reviews of Intervention (chapter 7.6-7.8 and 16.1.3)⁴² (see **supplementary text**).

Role of the Funding Source

This study received no funding.

The protocol of the meta-analysis was submitted as a module assignement fo the Systematic Review module and internally peer-reviewed at HUMANITAS University Gradenigo Hospital Institutional Review Board and is available at our Institution at request.

Patient involvement

No patients were involved in definition of the research question or the outcome measures, and interpretation or writing up of results. Data relating to the impact of the intervention on participants' quality of life were not extracted. Where possible, results of this meta-analysis will be disseminated to the patient community or individual patients and families through the investigators of this meta-analysis.

RESULTS

The flow of study selection is reported in **Figure 1**. At the end of selection, 6 placebo-controlled RCTs

(duration ranging 4-52 weeks) enrolling 3238 T1DM participants were included in the meta-analysis ^{26,27,28,29,30,31,58,59} (main characteristics reported in **supplementary Table 1**).

Twelve phase 1 RCTs conducted in nondiabetic individuals, 18 RCTs enrolling T2DM patients (4 completed, 14 active) and 1 RCT enrolling nondiabetic patients with congestive heart failure were excluded (main characteristics of excluded RCTs reported in **supplementary Table 2**).

All included RCTs compared sotagliflozin with placebo on background insulin treatment. Three RCTs^{28,30,31} compared different sotagliflozin doses (75 mg, 200 mg or 400 mg) with placebo Overall, ten comparison were available for the meta-analysis.

Two RCTs adopted insulin dose optimization (target: FPG 80-130 mg/dL and 2hr-PPG>180 mg/dL) during the 6 weeks preceding randomization ^{30,31}.

Two RCTs excluded patients with impaired renal function (eGFR<60 ml/min/1.73m2)^{26,28}, four RCTs excluded patients with moderate-to-severe (eGFR<45 ml/min/1.73m2) renal impairment^{27,29,30,31}.

Participants' baseline characteristics were equally balanced between the study arms and in all RCTs dropout rates were generally low and balanced across arms. No trial used the last-observation-carried-forward (LOCF) approach to impute missing observations, which were imputed as nonresponse for dichotomous outcomes; for continuous outcomes, mixed-effects model for repeated measures (MMRM) statistics based on the restricted maximum likelihood method for estimation was used.

Two RCTs were clearly funded by non-profit organizations^{26,27}, while a pharmaceutical company funded four RCTs: however, we did not find any evidence of high risk of biased outcomes in trial designing, conducting and reporting.

The overall quality was good for all included RCTs. The risk of bias summary for individual RCTs and the risk of bias graph for each item across included RCTs are detailed in **supplementary Table 1** and summarized in **supplementary Figure 1-2**.

The analysis of Funnel plots and the Egger test (p>0.67 for all outcomes) did not find any evidence of publication bias (supplementary Figure 3 panel A-S).

No values had to be imputed for the meta-analysis during data extraction.

The agreement between the 2 reviewers for study selection was 0.96 and for quality assessment of trials was 0.89.

Glycemic efficacy outcomes

HbA1c

Compared with placebo, sotagliflozin treatment was associated with a significant reduction in HbA1c levels (WMD -0.34%, 95%CI: -0.41 to -0.27%, p<0.00001, I²=20%, N-comparisons=10, 3238 participants)(**Figure 2 panel A**). There was little heterogeneity in the meta-analysis, suggesting a consistent drug effect.

Subgroup and meta-regression analysis revealed the effect was independent of trial duration (β =0. 110; p=0. 28) and baseline HbA1c (β =0. 119; p=0. 384) (Supplementary Table 3).

HbA1c reduction with sotagliflozin 400 mg/d was higher than with 200 mg/d (**Supplementary Table 4**).

Fasting plasma glucose (FPG) and 2h-postprandial plasma glucose (2h-PPG)

Sotagliflozin significantly reduced FPG (WMD -16.98 mg/dL, 95%CI: -22.09 to -11.86 mg/dL, p<0.00001, I²=6%, N-comparisons=10, 3238 participants) and 2h-PPG (WMD -39.24 mg/dL, 95%CI: -50.42 to -28.06 mg/dL, p<0.00001, I²=20%, N-comparisons=9, 539 participants) (**Figure 2 panel B-C**). There was little heterogeneity in the meta-analysis, suggesting a consistent drug effect. The effect was independent of trial duration and baseline HbA1c (Supplementary Table 3)...

Continuous Glucose Monitoring (CGM) parameters

Four RCTs evaluated CGM-derived parameters ^{26,27,30,31}.

Compared with placebo, sotagliflozin significantly increased time-in-range (WMD +9.73%, 95%CI: 6.66 to 12.81%, p<0.00001, I²=24%, N-comparisons=6, 398 participants) and reduced average daily glucose (WMD -15.09 mg/dL, 95%CI: -21.40 to -8.79 mg/dL, p<0.00001, I²=28%, N-comparisons=5, 312 participants), SD around average daily glucose (WMD -6.68 mg/dL, 95%CI: -10.59 to -2.77 mg/dL, p=0.0008, I²=0%, N-comparisons=5, 311 participants) and mean amplitude of glucose excursion (MAGE) (WMD -19.52 mg/dL, 95%CI: -28.91 to -10.54 mg/dL, p<0.0001, I²=0%, N-comparisons=5, 311 participants) (supplementary Figure 4 panel A-D).

There was little heterogeneity in the meta-analysis, suggesting a consistent drug effect.

Sotagliflozin 400 mg/d was significantly more effective than 200 mg/d dose at improving time-in-range, average daily glucose and MAGE (**Supplementary Table 4**).

Daily Total, Basal and Bolus Insulin Dose

Compared with placebo, sotagliflozin reduced daily total (WMD -8.99%, 95%CI: -10.93 to -7.05%, p<0.00001, I^2 =33%, N-comparisons=10, 3238 participants), basal (WMD -8.03%, 95%CI: -10.14 to -5.93%, p<0.00001, I^2 =0%, N-comparisons=10, 3238 participants) and bolus (WMD -9.14%, 95%CI: -12.17 to -6.12%, p<0.00001, I^2 =67%, N-comparisons=10, 3238 participants) insulin dose in T1DM patients (**supplementary Figure 5 panel A-C**).

Heterogeneity for bolus insulin dose was high, and was accounted for by significant subgroup differences between high-dose (400 mg/d) and low-dose (200 mg/d) sotagliflozin (**supplementary Table 4**).

Urinary glucose excretion

Pooled data from two RCTs^{26,28} indicated daily UGE progressively increased with increasing sotagliflozin dose from 75 mg/d to 200 mg/d, but then UGE reached a plateau around 60 g/24 hr with either 200 mg/d and 400 mg/d sotagliflozin (supplementary Figure 6; supplementary Table 4)

Non-glycemic outcomes

Body weight

Compared with controls, sotagliflozin induced a significant weight reduction (WMD -3.54%, 95%CI: -3.98 to -3.09%, p<0.00001, I²=18%, N comparisons=10, 3238 participants) (**Figure 3 panel A**).

On meta-regression analysis, weight change (%) correlated with the magnitude of total insulin dose reduction from baseline (β =0.213; p=0.001).

Blood pressure (BP)

Compared to placebo, sotagliflozin use was associated with a reduction in systolic BP (WMD -3.85 mmHg, 95%CI: -4.76 to -2.93, p<0.00001, I^2 =0%) and in diastolic BP (WMD -1.43 mmHg, 95%CI: -1.98 to -0.89, p<0.00001, I_2 =0%, N comparisons=10, 3238 participants) (**Figure 3 panel B-C**).

These effects were not associated with an increased incidence of ortostatic hypotension (not shown).

Renal effects: eGFR and urinary ACR

Compared with placebo, sotagliflozin treatment was associated with a slight reduction in eGFR as (WMD: -0.80, 95% CI: -1.42 to -0.18 ml/min/1.73 m², p=0.01, I²=0%, N comparisons=10, 3238 participants)(**Figure 4 panel A**).

Urinary ACR was evaluated in 3 phase 3 RCTs (2977 participants, trial duration ranging 24-52 weeks, mean baseline ACR of participants of 52.6, 31.6, 54.3 mg/g, respectively 29,30,31). Pooled analysis of these RCTs showed sotagliflozin was associated with a decrease in ACR (WMD: -14.65, 95% CI: -2.58 to -26.72 mg/g, p=0.02, I₂=0%, N comparisons=5 (**Figure 4 panel A-B**). Subgroup analysis revealed eGFR reduction with sotagliflozin occurred only in RCTs lasting \leq 12 weeks, but not in RCTs of longer duration (**Supplementary Table 4**).

To gain further insight into the effect of time on renal function, we examined the effect of sotagliflozin on eGFR in the 2 RCTs of longest duration (52 weeks) during the initial 24 weeks and during the following 28 weeks. While sotagliflozin continued to reduce ACR throughout the treatment period, the difference

in eGFR between sotagliflozin and placebo varied during follow-up: during the initial 24 weeks patients receiving sotagliflozin experienced a decline in eGFR, while in the following 28 weeks sotagliflozin significantly slowed the eGFR decline as compared with placebo (supplementary Figure 7 panel A-B).

Plasma lipids

No RCT reported the effect of active treatment or placebo on LDL-C, HDL-C and triglyceride

Safety outcomes

Hypoglycemia and severe hypoglycaemia

The definition of hypoglycemia and severe hypoglycemia was consistent across all RCTs (see online Appendix). Compared with placebo, sotagliflozin treatment was associated with a lower rate of hypoglycemia events (WMD: -9.09 events per patient-year, 95% CI: -13.82 to -4.36 events per patient-year, p=0.0002, I₂=0%, N comparisons=10, 3238 participants) and with a 31% lower risk of severe hypoglycaemia (RR 0.69, 95% CI: 0.49-0.98, p=0.04; N comparisons=10, I²=0%) (**Figure 5 panel A-B**).

Diabetic ketoacidosis (DKA)

Compared with placebo, sotagliflozin was associated with an increased risk of DKA (RR 3.93, 95%CI: 1.94-7.96, p=0.0001; N comparisons=10, I²=0%, 3238 participants, trial duration ranging 4–52 weeks)(**Figure 5 panel C**). Forty-six (69 %) of all cases of DKA occurred at blood glucose>250 mg/dL, while the remaining 21 cases(31%) occurred with blood glucose values ranging 150-250 mg/dL (supplementary Table 5).

The risk for DKA was increased for patients on multiple daily injections (MDI) (RR 3.22, 95%CI: 1.24-

9.09, p=0.01; N comparisons=10, I^2 =0%, 2072 patients) as well as for patients on continuous subcutaneous infusion(CSI) (RR 6.40, 95%CI: 2.82-15.64, p<0.0001; N comparisons=10, I^2 =0%, 1166 patients).

Subgroup analyses revealed the risk of DKA varied according to initial HbA1c of included RCTs: the risk of DKA was increased in RCTs with a mean initial HbA1c<8% (RR 6.62, 95%CI: 2.04-21.48), I^2 =0%, p=0.002, N=3, 1608 participants), but not in RCTs with a mean HbA1c≥8% (RR 2.21, 95%CI: 0.43-11.42, I^2 =0%, p=0.34, N =3, 1630 participants) (supplementary Table 4).

In a meta-regression model including sotagliflozin dose, trial duration, initial HbA1c, initial FPG, changes in HbA1c and FPG, total bolus and basal insulin doses (baseline, changes and end-of-treatment doses) fasting and postprandial glycemia, body weight changes, volume depletion events, the risk of DKA correlated inversely with initial HbA1c (β =-0.331; p=0.009) and with the magnitude of basal insulin dose reduction (β =-0.218; p=0.012) (**supplementary Figure 8**).

Urinary tract infections (UTIs) and genital tract infections (GTI)

Compared with placebo, sotagliflozin did not affect the risk of UTIs (RR 0.97, 95% CI: 0.71-1.33, p=0.84; N comparisons=10, I²=0%, 3238 participants) but was associated with an increased risk of mycotic GTIs (RR 3.12, 95% CI: 2.14-4.54, p<0.00001; N comparisons=10, I²=0%) (**Figure 6 panel A-B**).

In a meta-regression model, the risk of GTI was not related to sotagliflozin dose, urinary glucose excretion, initial HbA1c, initial FPG, changes in HbA1c and FPG (all p-values>0.5).

Gastrointestinal events

Compared with control, sotagliflozin was associated with an increased risk of diarrhea (RR 1.50, 95%CI: 1.08-2.10, p=0.02; N comparisons=10, I²=0%, 3238 participants) (**Figure 6 panel D**), but not of other gastrointestinal symptoms(**supplementary Table 5**).

Other adverse events

Compared with control, sotagliflozin treatment was associated with an increased risk of acidosis-related AEs (RR: 3.85, 95%CI: 2.33-6.36, p<0.00001; N comparisons=10, I^2 2=0%) and of volume depletion events (RR: 2.19, 95%CI: 1.10-4.36, p=0.03; N comparisons=10, I^2 =0%) (**Figure 6 panel D**; **supplementary Table 5**). Subgroup analysis revealed the risk of volume depletion events was increased in the first 12 weeks of treatment, but then subsided (**supplementary Table 3**).

The most common AEs leading to treatment discontinuation were DKA (35.8 % of all patients experiencing DKA discontinued treatment), diarrhea (treatment discontinuation in 6.9% of patients), genital tract infections (treatment discontinuation in 6.3 % of patients), severe hypoglycaemia (treatment discontinuation in 5.6 % of patients), UTIs (treatment discontinuation in 4.4 % of patients) and volume depletion events ((treatment discontinuation in 4.3 % of patients).

Sotagliflozin did not affect the risk of MACE (RR 1.06, 95% CI: 0.40-2.82, p=0.91; N comparisons=10, I²=6%), cancer (RR 0.86, 95% CI: 0.25-2.97, p=0.81; N comparisons=9, I²=0%) or all-cause death (RR 0.35, 95% CI: 0.07-1.71, p=0.19; N comparisons=9, I²=0%) (**supplementary Table 5, supplementary Figure 9 panel B**),

The effect of sotagliflozin on other AEs is summarized in supplementary Table 4.

Dose-response analysis

Three RCTs evaluated the effects of sotagliflozin 400 mg and 200 mg and one RCT assessed also the 75 mg dose-effect. The analysis of dose-response interactions within these 3 RCTs found that the 200 mg dose had a greater glycosuric effect than the 75 mg dose (UGE), but this effect did not increased further with the 400 mg dose.

Sotagliflozin 400 mg/d was associated with a greater improvement than sotagliflozin 200 mg/d in the following outcomes HbA1c, FPG, 2h-PPG, time-in-range, average daily glucose, daily total basal and

bolus insulin dose, body weight, systolic BP, eGFR and ACR (**supplementary Table 5**). We didn't find any relationship between different sotagliflozin doses and adverse events. The results of the within-trial comparison were all confirmed by the across-trial approach.

Sensitivity analyses

Sensitivity analysis conducted using alternative pooling methods, including Peto's Odds Ratio (OR), which has a greater power at event rates below 1%⁵⁴, confirmed the results of the main analysis (supplementary Table X)

Grading of Evidence

Quality of evidence was downgraded to moderate for effect on time-in-range glucose as it was unclear whether the population undergoing CGM substudies was representative of the whole study population, and to low for MACE and all-cause mortality for imprecision (**Table 1-2**).

DISCUSSION

The main findings of our analysis are the following:

- in T1DM patients, sotagliflozin as add-on therapy to insulin ameliorated glycemic efficacy outcomes
 and showed also nonglycemic benefits, including body weight, blood pressure and nephropathy
 marker reduction.
- 2. sotagliflozin treatment was associated with a significant reduction in the incidence of hypoglycaemia and severe hypoglycemia
- 3. DKA was the most serious and frequent adverse event associated with sotagliflozin treatment, which also increased the risk of GTIs, diarrhea, and volume depletion events, but not of UTIs.
- 4. The risk of DKA varied depending on initial HbA1c levels and basal insulin dose reduction.

T1DM patients achieve glycemic goals in 30% of cases, experience severe hypoglycemia in up to 20% of

cases per year and are overweight in 40% of cases³, hence urgently needing adjunctive therapeutic strategies to complement glucose-lowering effects of insulin and mitigate its unwanted effects.

Hypoglycemia, which results from the total dependence of T1D patients on injected insulin therapy, is of particular concern and can be viewed at the basis of highest unmet need in this population^{9,10}, as it is the main factor limiting optimal glucose control; furthermore, severe hypoglycemia is a strong predictor of adverse clinical outcomes and death in diabetic patients^{7,8-18,60}. None of the drugs recently approved for T2DM and seeking an indication for T1DM, including incretin analogues and SGLT2 inhibitors, reduced hypoglycemic risk, which is either unaffected or increased by these therapies^{22,66,61}. Several mechanisms may underlie the observed hypoglycemic risk reduction observed with sotagliflozin. The dual intestinal SGLT1 and renal SGLT2 inhibition blunts acute glucose fluctuations and reduces glycemic variability (supplementary Figure 4C-D), thereby limiting the need for bolus insulin correction doses and the attendant hypoglycemic risk (supplementary Figure 5C)^{15,16,62}. The reduction in the rate of hypoglycemic events may have *per se* contributed to reduce severe hypoglycemia: the recurrence of hypoglycemic episodes blunts autonomic and hormonal responses to subsequent hypoglycemia, impairs hypoglycemia awareness and glucose counterregulation and paves the way to severe hypoglycemia. This functional impairment in counterregolatory mechanisms is distinct from autonomic neuropathy, occurs in the short-term and can be rapidly reversed by reducing hypoglycemia recurrence⁶³.

The analysis of pooled results from phase 3 RCTs disclosed also potential renoprotection for sotagliflozin, which reduced microalbuminuria, a marker of early diabetic nephropathy and an independent cardiovascular risk factor ¹⁹(**Figure 4 panel B**). The transient eGFR decline observed in the initial 12 weeks of treatment is similar to that observed with other SGLT2 inhibitors ⁶⁴ and is consistent with renoprotective mechanisms of SGLT2 inhibition, which enhance afferent arteriolar tone, reduce intraglomerular pressure and relieve glomerular hyperfiltration and barrier damage ⁶⁵. However, in patients receiving sotagliflozin the reduced glomerular perfusion may be aggravated by volume depletion favoured by concomitant osmotic glycosuria (due to renal SGLT2 inhibition) and diarrhea (induced by intestinal SGLT1 inhibition) (**Figure 6 panel D**). Hence it is important to avoid volume depletion in the

early months of treatment with sotagliflozin..

Differently from SGLT2 inhibitors, sotagliflozin did not increase the risk of UTIs (**Figure 6 panel A**): the lower glycosuric effects of sotagliflozin as compared with SGLT2 inhibitors⁶⁶ may have limited the incidence of UTIs, while SGLT1-mediated intestinal glucose malabsorption may have increased diarrhea, usually mild, self-limiting and not inducing treatment discontinuation.

Further supporting the relevance of intestinal SGLT1 inhibition, a dose-response gradient for most glycemic outcomes was observed with increasing sotagliflozin dosage, not paralleled by an increase in glycosuria, which reached a plateau at 60 g/day, 40-50% lower than that reported with full-dose SGLT2 inhibitors ^{67,68} (**supplementary Figure 6**). Whether sotagliflozin maintains unaltered glucose-lowering efficacy in the presence of moderate-to-severe renal failure will be assessed by ongoing trials in T2DM (**supplementary Table 2**)

DKA was the most common relevant adverse event, observed in 61 out of 1912 (3.1%) of sotagliflozintreated patients and inducing treatment discontinuation in 38% of cases (**supplementary Table 5**).

While SGLT2 inhibitor-associated DKA has been reported to occur often at uncharacteristically normal
or mildly elevated (<250 mg/dL) blood glucose levels (euglycemic DKA)⁶⁹, over two thirds of cases of
sotagliflozin-related DKA occurred at high blood glucose levels(**supplementary Table 5**). Notably, our
data indicate a lower initial HbA1c and a greater basal insulin dose reduction during sotagliflozin
treatment increase the risk for DKA (**supplementary Figure 8**; **supplementary Table 3**), possibly
because patients with less deteriorated baseline glycemic control experienced a more rapid insulin dose
down-titration with sotagliflozin. The extent of basal insulin down-titration seems central for DKA
development by allowing unrestricted fasting-induced lipolysis and ketogenesis on a background of
negative glucose balance⁶⁹. Consistently, insulin dose reduction >20% has been found to increase ketone
levels and diminish the glucose-lowering effect of SGLT2 inhibitors⁷⁰.

Clinical and policy implications

In conclusion, sotagliflozin for up to 52 weeks provided consistent glycemic and nonglycemic benefits in T1DM, including the reduction of unwanted effects of insulin therapy, i.e., weight gain and

hypoglycemia. These effects make sotagliflozin an attractive adjunctine therapy to insulin in T1DM patients, which achieve target glycemic goals in 30% of cases, are overweight in 40 % of cases and experience severe hypoglycemia at a rate of up to 20% of patients per-year³. The clinical impact of these benefits may be more appreciable in patients at higher risk of severe hypoglycemia, like those with recurrent hypoglycemia and hypoglycemia unawareness, who represent 17-36% of the general T1DM population⁷¹.

Our analysis may also help minimize the risk of DKA in T1DM treated with sotagliflozin by appropriate patient selection and by defining appropriate protocols for basal insulin dose adjustment. Ketone testing should be performed after each basal insulin dose reduction, rather than relying solely on overt triggering conditions or symptoms of DKA^{28,29,30,31}, which often fail to recognize early DKA⁷². Future research should define safer protocols for basal insulin dose adjustment: as an example, in a recent phase 3 RCT with dapagliflozin reporting no increased risk of DKA, participants were instructed to reduce insulin doses by no more than 20% on treatment initiation, to measure ketonemia whenever glucose readings were consistently elevated, and then subsequently to up-titrate insulin doses back to baseline following positive ketone testing⁷³.

Strengths and limitations

Strengths and limitations of our analysis derive from the characteristics of included evidence: strengths include the thorough assessment of efficacy and safety outcomes, the direct impact of extracted evidence regarding relevant clinical outcomes, like hypoglycemia and DKA, on decision-making in T1DM management. Limitations are the relatively small number and short duration of included trials, not exceeding 52 weeks, which prevented robust assessment of long-term hard outcomes, like MACE and overall mortality. Furthermore, although all included RCTs had good methodological quality, 66% of them were industry-funded, which makes them liable to sponsorship bias⁴⁵. Recent guidelines recommend against automatically downgrading industry-funded trials and we therefore address this issue by verifying a list of items empirically linked by recent literature to biased outcomes in industry-funded trials⁴³

Competing interests: All authors have completed the ICMJE uniform disclosure form at www.icmje.org/coi_disclosure.pdf and declare: no support from any organisation for the submitted work; no financial relationships with any organisations that might have an interest in the submitted work in the previous three years; no other relationships or activities that could appear to have influenced the submitted work. Giovanni Musso takes responsibility for the integrity of the data and the accuracy of the data analysis.

Exclusive license statement.

Giovanni Musso has the right to grant on behalf of all authors and does grant on behalf of all authors, a worldwide license—to the Publishers and its licensees in perpetuity, in all forms, formats and media (whether known now or created in the future), to i) publish, reproduce, distribute, display and store the Contribution, ii) translate the Contribution into other languages, create adaptations, reprints, include within collections and create summaries, extracts and/or, abstracts of the Contribution and convert or allow conversion into any format including without limitation audio, iii) create any other derivative work(s) based in whole or part on the on the Contribution, iv) to exploit all subsidiary rights to exploit all subsidiary rights that currently exist or as may exist in the future in the Contribution, v) the inclusion of

electronic links from the Contribution to third party material where-ever it may be located; and, vi) licence any third party to do any or all of the above.

Funding. This work received no funding

Data sharing statement:

The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

Role of each author:

Giovanni Musso: data collection and elaboration, statistical analysis, writing of the manuscript Roberto Gambino: data collection and discussion, review of the manuscript, approval of manuscript Maurizio Cassader: data collection and discussion, review of the manuscript, approval of manuscript Elena Paschetta: data collection and discussion, writing of the manuscript, approval of manuscript

Acknowledgements

The authors are deeply grateful to Professor Richard D Riley, Centre for Prognosis Research, Research Institute for Primary Care and Health Sciences, Keele University, Keele, United Kingdom, for his precious statistical advice.

Table 1. Quality of evidence for clinically relevant glycemic and nonglycemic effect outcomes: Summary of Findings Table according to the GRADE approach

Sotagliflozin	compared to	nlaceho	for type '	1 diahetes:	alvcemic	effect outcomes
OULAUIIIOZIII	Collibal Eu lo	DIACEDO	IOI LVDC	i ulabetes.	UIVCEIIIIC	CHECK OULCOINES

	Anticipated absolu	te effects* (95% CI)			Certainty of the	
Outcomes	Risk with placebo	Risk with sotagliflozin	Relative effect (95% CI)	№ of participants (studies)	evidence (GRADE)	Comments
Mean change in HbA1c(%) follow up: range 4 weeks to 52 weeks	The mean change in HbA1c ranged from -0.99 to +0.04 %	The mean change in HbA1c in the intervention group was 0,34 % lower (0,41 lower to 0,27 lower)	-	3238 (6 RCTs)	ФФФФ HIGH	Large effect. Dose-response gradient across the 200-400 mg doses
Mean change in fasting plasma glucose (FPG)(mg/dL) follow up: range 4 weeks to 52 weeks	The mean change in FPG ranged from -11 to +39 mg/dL	The mean change in FPG in the intervention group was 16,98 mg/dL lower (22,09 lower to 11,86 lower)	-	3238 (6 RCTs)	ФФФФ HIGH	Large effect Dose-response gradient across the 200-400 mg doses
Mean change in 2hr- postprandial plasma glucose (2h- PPG)(mg/dL) follow up: range 4 weeks to 52 weeks	The mean change in 2h-PPG ranged from -18.5 to +0 mg/dL	The mean change in 2h-PPG in the intervention group was 39,24 mg/dL lower (50,42 lower to 28,06 lower)	-	539 (5 RCTs)	ФФФФ HIGH	Large effect. Dose-response gradient across the 200-400 mg doses
Mean change in % time-in-range (70-180 mg/dL) follow up: range 4 weeks to 52 weeks	The mean mean change in % time-in-range ranged from -1.83 to -0.2	The mean change in % time-in-range in the intervention group was 9,73 % higher (6,66 higher to 12,81 higher)	-	398 (4 RCTs)	⊕⊕⊕○ MODERATE ®	Large effect. Dose-response gradient across the 200-400 mg doses
Sota	gliflozin com	pared to place	ebo for type 1	diabetes: no	n-glycemic	effect outcomes
Mean change in body weight (%) follow up: range 4 weeks to 52 weeks	The mean change in body weight ranged from -0.99 to +0.04 %	The mean change in body weight in the intervention group was 3,54 % lower (3,98 lower to 3,09 lower)	-	3238 (6 RCTs)	ФФФФ HIGH	Dose-response gradient across the 200-400 mg doses
Mean change in systolic blood pressure (BP)(mmHg) follow up: range 4 weeks to 52 weeks	The mean change in systolic BP ranged from -3.8 to 1.7 mmHg	The mean change in systolic blood pressure (BP) in the intervention group was 3,85 mmHg lower (4,76 lower to 2,93 lower)	-	3238 (6 RCTs)	⊕⊕⊕⊕ ніGн	Dose-response gradient across the 200-400 mg doses
Mean change in eGFR (ml/min/1.73 m²) follow up: range 4 weeks to 52 weeks	The mean mean change in eGFR ranged from -1.09 to 0.34 ml/min/1.73 m ²	The mean mean change in eGFR in the intervention group was 0,8 ml/min/1.73 m2 lower (1,42 lower to 0,18 lower)	-	3238 (6 RCTs)	ФФФФ HIGH	Dose-response gradient across the 200-400 mg doses

Mean change in urinary albumin/creatinine ratio (ACR)(mg/g) follow up: range 24 weeks to 52 weeks	The mean mean change in urinary ACR ranged from 4.1 to 14.9 mg/g	The mean change in urinary ACR in the intervention group was 14,57 mg/g lower (26,87 lower to 2,28 lower)	-	2977 (3 RCTs)	⊕⊕⊕⊕ ніgн	Dose-response gradient across the 200-400 mg doses
---	--	---	---	------------------	--------------	--

^{*}The risk in the intervention group (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

CI: Confidence interval; MD: Mean difference; RR: Risk ratio

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate. The true effect is likely to be substantially different from the estimate of effect

Explanations

a. unclear if the population undergoing Continuous Glucose Monitoring substudies was representativ of the whole trial population in the inTandem1 and inTandel2 trials

Table 2. Quality of evidence for clinically relevant adverse events (AEs): Summary of Findings Table according to the GRADE approach

Sotagliflozin compared to placebo for type 1 diabetes: adverse events (AEs)

	Anticipated ab	solute effects* (95% CI)	Relative effect	№ of participants	Certainty of the		
Outcomes	Risk with placebo			(studies)	evidence (GRADE)	Comments	
Mean change in hypoglycemia events(events per patient-year) follow up: range 4 weeks to 52 weeks	The mean change in hypoglycemia events ranged from 69 to 179 events/patient- year	The mean change in hypoglycemia events in the intervention group was 9,09 events/patient-year lower (13,82 lower to 4,36 lower)	-	3238 (6 RCTs)	ФФФ нісн		
Incidence of severe hypoglycemia follow up: range 4 weeks to 52 weeks	43 per 1.000	30 per 1.000 (21 to 42)	RR 0.69 (0.49 to 0.98)	3238 (6 RCTs)	⊕⊕⊕ HIGH		
Incidence of diabetic ketoacidosis (DKA) follow up: range 4 weeks to 52 weeks	5 per 1.000	18 per 1.000 (9 to 36)	RR 3.93 (1.94 to 7.96)	3238 (6 RCTs)	⊕⊕⊕ HIGH	Large effect	
Incidence of urinary tract infections(UTIs) follow up: range 4 weeks to 52 weeks	48 per 1.000	46 per 1.000 (34 to 63)	RR 0.97 (0.71 to 1.33)	3238 (6 RCTs)	⊕⊕⊕ HIGH		
Incidence of genital tract infections(GTIs) follow up: range 4 weeks to 52 weeks	23 per 1.000	73 per 1.000 (50 to 106)	RR 3.12 (2.14 to 4.54)	3238 (6 RCTs)	⊕⊕⊕ HIGH	Large effect	
Incidence of diarrhea follow up: range 4 weeks to 52 weeks	35 per 1.000	52 per 1.000 (37 to 73)	RR 1.50 (1.08 to 2.10)	3238 (6 RCTs)	⊕⊕⊕ ніGн		
Incidence of AEs leading to treatment discontinuation follow up: range 4 weeks to 52 weeks	23 per 1.000	31 per 1.000 (18 to 54)	RR 1.34 (0.78 to 2.30)	3238 (6 RCTs)	ФФФ нібн		
Incidence of serious AEs follow up: range 4 weeks to 52 weeks	69 per 1.000	76 per 1.000 (58 to 99)	RR 1.11 (0.85 to 1.44)	3238 (6 RCTs)	ФФФ нідн		
Incidence of major adverse cardiovascular events (MACE) follow up: range 4 weeks to 52 weeks	5 per 1.000	6 per 1.000 (2 to 15)	RR 1.06 (0.40 to 2.82)	3238 (6 RCTs)	LOM ₅	Few events, OIS not reached	
All-cause mortality follow up: range 4 weeks to 52 weeks	2 per 1.000	1 per 1.000 (0 to 4)	RR 0.34 (0.07 to 1.70)	3238 (6 RCTs)	⊕⊕⊖⊖ LOW ª	Few events, OIS not reached	

Table 2. Quality of evidence for clinically relevant adverse events (AEs): Summary of Findings Table according to the GRADE approach

Sotagliflozin compared to placebo for type 1 diabetes: adverse events (AEs)

0.1	Anticipated ab	solute effects* (95% CI)	Relative effect	№ of participants (studies)	Certainty of the	Comments
Outcomes	Risk with placebo	Risk with sotagliflozin	(95% CI)		evidence (GRADE)	

^{*}The risk in the intervention group (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

CI: Confidence interval; MD: Mean difference; RR: Risk ratio; OIS: optimal information size

GRADE grades Working Group of evidence High certainty: We are very confident that the true effect lies close to that of the estimate of the effect Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is different substantially

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

Explanations

a. downgraded for imprecision

Figure 1: evidence acquisition flow diagram

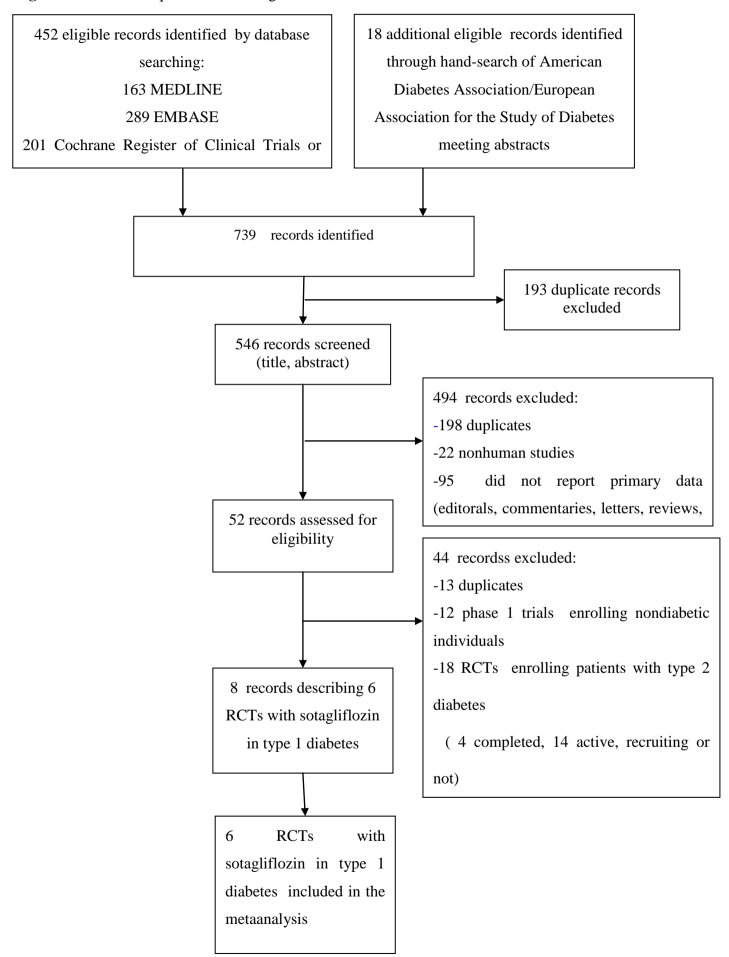
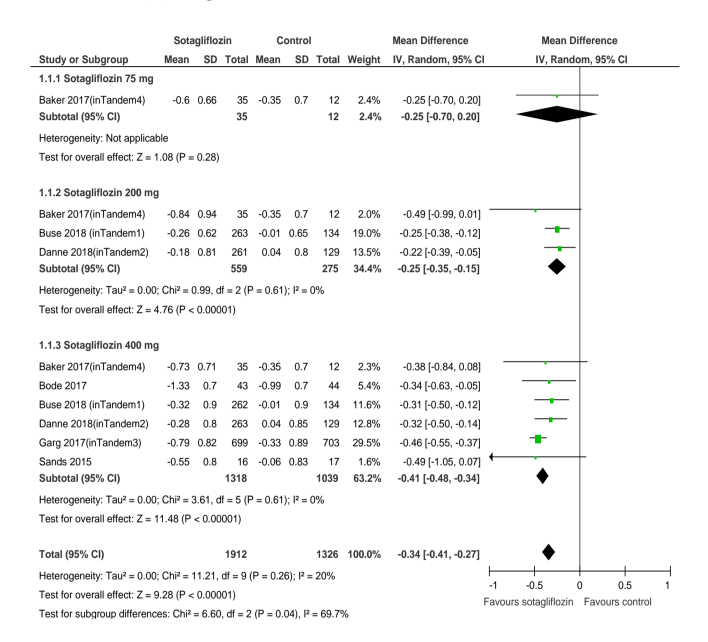
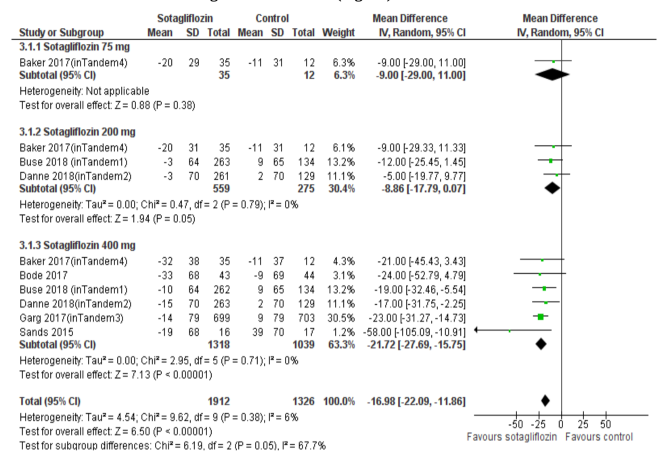


Figure 2. Forest plot of comparison: Sotagliflozin vs. placebo, outcome: HbA1c(%), Fasting Plasma Glucose (FPG) and 2 hour-Postprandial Plasma Glucose (2h-PPG).

Panel A: HbA1c(%) changes from baseline



Panel B: outcome: FPG changes from baseline (mg/dL)

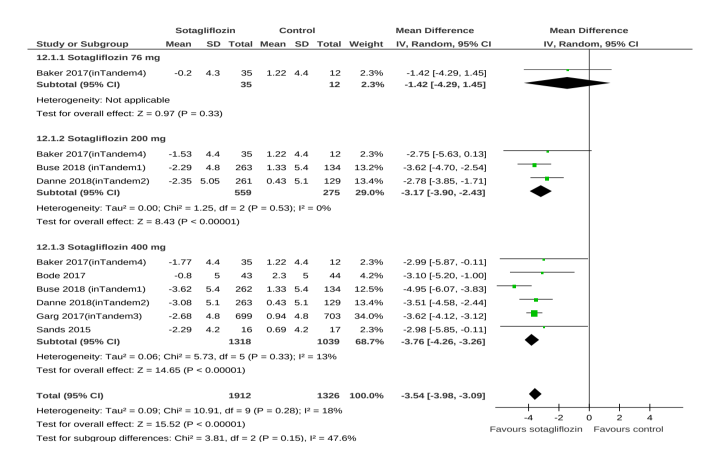


Panel C: outcome: 2h-PPG changes from baseline (mg/dL)

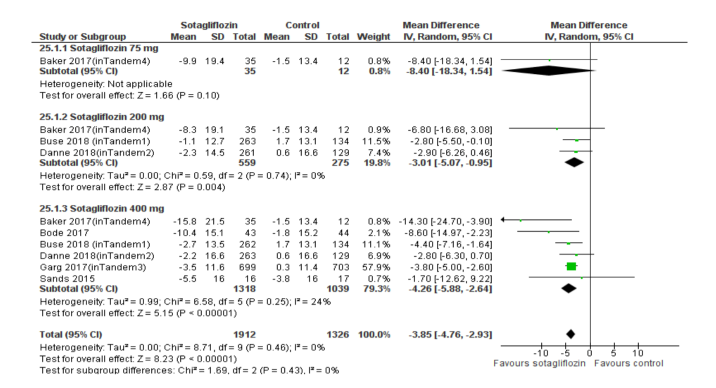
	Sota	gliflo	zin	Control		I		Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
4.1.1 Sotagliflozin 75 mg									
Baker 2017(inTandem4) Subtotal (95% CI)	-20	40	35 35	-0.2	38	12 12	14.6% 14.6 %	-19.80 [-45.06, 5.46] - 19.80 [-45.06, 5.46]	•
Heterogeneity: Not applicat	ole								
Test for overall effect: $Z = 1$.	54 (P = 0	.12)							
4.1.2 Sotagliflozin 200 mg									
Baker 2017(inTandem4)	-28	43	35	-0.2	38	12	14.1%	-27.80 [-53.59, -2.01]	
Buse 2018 (inTandem1)	-35.7	75	44	-18.5	75	22	7.4%	-17.20 [-55.58, 21.18]	
Danne 2018(inTandem2)	-40.1	89	45	10.2	81	24	6.4%	-50.30 [-91.85, -8.75]	
Subtotal (95% CI)			124			58	27.9 %	-29.91 [-48.94, -10.88]	-
Heterogeneity: Tau ² = 0.00;		•	,	= 0.50);	2 = (1%			
Test for overall effect: Z = 3.	.08 (P = 0	.002)							
4.1.3 Sotagliflozin 400 mg									
Baker 2017(inTandem4)	-50	38	35	-0.2	38	12	14.9%	-49.80 [-74.71, -24.89]	
Bode 2017	-56	56	43	0	57	44	15.9%	-56.00 [-79.74, -32.26]	
Buse 2018 (inTandem1)	-40.2	89	47	-18.5	88	23	5.8%	-21.70 [-65.75, 22.35]	
Danne 2018(inTandem2)	-65.5	72	49	0	71	24	8.7%	-65.50 [-100.33, -30.67]	
Sands 2015	-39	42	16	-1	41	17	12.2%	-38.00 [-66.34, -9.66]	
Subtotal (95% CI)			190			120	57.5 %	-48.92 [-61.87, -35.98]	•
Heterogeneity: Tau² = 0.00;			,	= 0.52);	$I^2 = 0$)%			
Test for overall effect: $Z = 7$.	.41 (P < 0	.0000	11)						
Total (95% CI)			349			190	100.0%	-39.24 [-50.42, -28.06]	•
Heterogeneity: Tau² = 57.31	1; Chi ² = 9	9.97, d	df = 8 (F	P = 0.27); ² =	20%			-50 -25 0 25 50
Test for overall effect: $Z = 6$.	.88 (P < 0	.0000	11)						-50 -25 0 25 50 Favours sotagliflozin Favours control
Test for subgroup differenc	es: Chi²=	5.35	. df= 2	(P = 0.0))7), l²	= 62.6	%	'	around dottagninozin i around control

Figure 3. Forest plot of comparison: Sotagliflozin vs. placebo, outcomes: body weight, systolic BP (sysBP) and diastolic BP (diaBP).

Panel A: body weight changes from baseline (%)



Panel B: outcome: sysBP changes from baseline (mmHg)

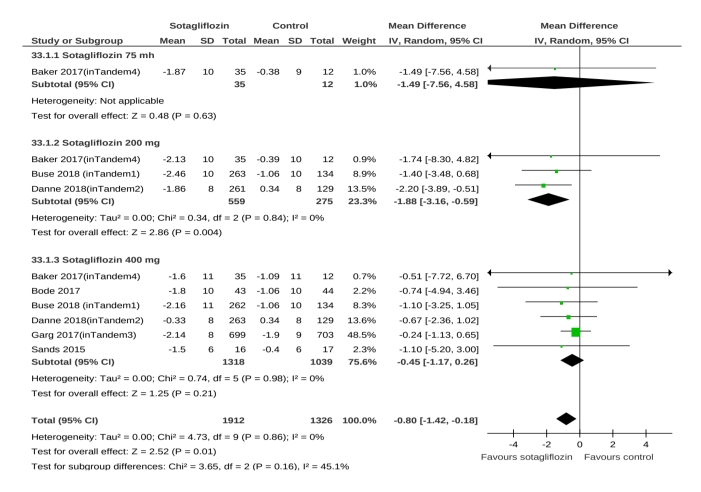


Panel C: outcome: diaBP changes from baseline (mmHg)

	Sota	gliflo	zin	Co	ontro	I		Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
26.1.1 Sotagliflozin 75 mg	3								
Baker 2017(inTandem4)	-1.1	7	35	-0.9	7	12	1.4%	-0.20 [-4.79, 4.39]	
Subtotal (95% CI)			35			12	1.4%	-0.20 [-4.79, 4.39]	
Heterogeneity: Not applica	ble								
Test for overall effect: Z = 0	0.09 (P =	0.93)							
26.1.2 Sotagliflozin 200 m	ng								
Baker 2017(inTandem4)	-2	7.1	35	-0.9	7.1	12	1.4%	-1.10 [-5.76, 3.56]	
Buse 2018 (inTandem1)	-0.6	7.5	263	0.9	7.5	134	12.1%	-1.50 [-3.06, 0.06]	
Danne 2018(inTandem2)	-1.6	7.6	261	-0.3	7.5	129	11.6%	-1.30 [-2.89, 0.29]	
Subtotal (95% CI)			559			275	25.1%	-1.39 [-2.47, -0.30]	•
Heterogeneity: Tau ² = 0.00	; Chi² = 0).05, c	lf = 2 (F	P = 0.98); l² =	: 0%			
Test for overall effect: Z = 2	2.51 (P =	0.01)							
26.1.3 Sotagliflozin 400 m	ng								
Baker 2017(inTandem4)	-3.9	7.1	35	-0.9	7.1	12	1.4%	-3.00 [-7.66, 1.66]	
Bode 2017	-2.5	7	43	0.8	7	44	3.4%	-3.30 [-6.24, -0.36]	
Buse 2018 (inTandem1)	-1.4	7.5	262	0.9	7.5	134	12.1%	-2.30 [-3.86, -0.74]	
Danne 2018(inTandem2)	-0.9	7.6	263	-0.3	7.5	129	11.7%	-0.60 [-2.19, 0.99]	
Garg 2017(inTandem3)	-0.8	7.9	699	0.5	7.7	703	44.1%	-1.30 [-2.12, -0.48]	-
Sands 2015	-1.5	8	16	-0.5	8	17	1.0%	-1.00 [-6.46, 4.46]	
Subtotal (95% CI)			1318			1039	73.5%	-1.47 [-2.10, -0.84]	◆
Heterogeneity: Tau ² = 0.00	; Chi² = 4	1.34, c	lf = 5 (F	P = 0.50); I ² =	: 0%			
Test for overall effect: Z = 4	4.57 (P <	0.000	001)						
Total (95% CI)			1912			1326	100.0%	-1.43 [-1.98, -0.89]	•
Heterogeneity: Tau ² = 0.00	; Chi² = 4	1.68, c	lf = 9 (F	o = 0.86); l² =	: 0%			
Test for overall effect: Z = 5	5.18 (P <	0.000	001)					Four	-4 -2 0 2 4 urs sotagliflozin Favours control
Test for subgroup difference	es: Chi² :	= 0.30), df = 2	P = 0.	86), I	² = 0%		Favo	uis sotagiiiloziii Favouis control

Figure 4. Forest plot of comparison: Sotagliflozin vs. placebo, outcomes: eGFR and urinary Albumin/Creatinine Ratio (ACR)

Panel A: outcome: eGFR changes from baseline (ml/min/1.73m²)



Panel B: outcome: ACR changes from baseline (mg/g)

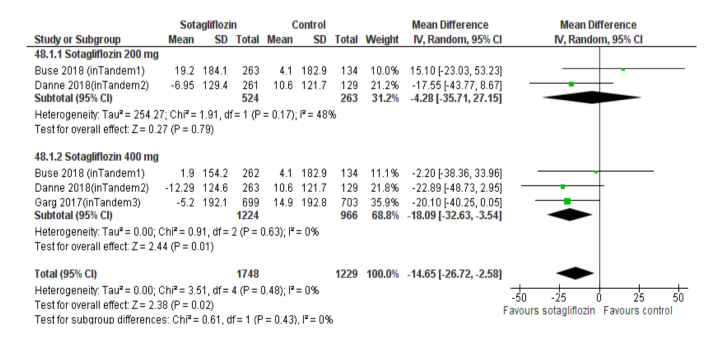
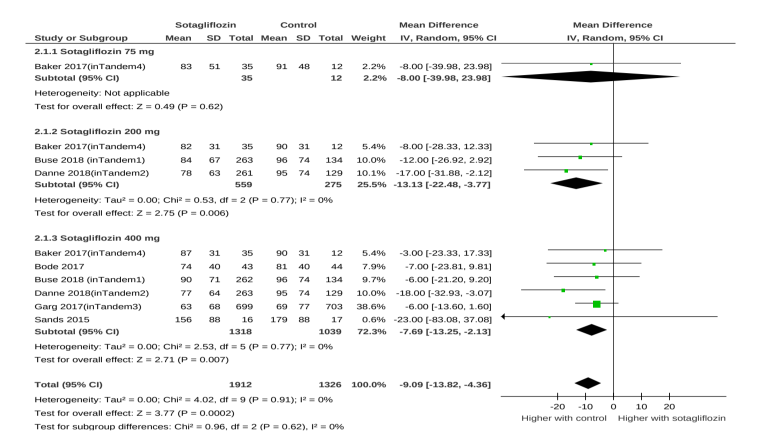
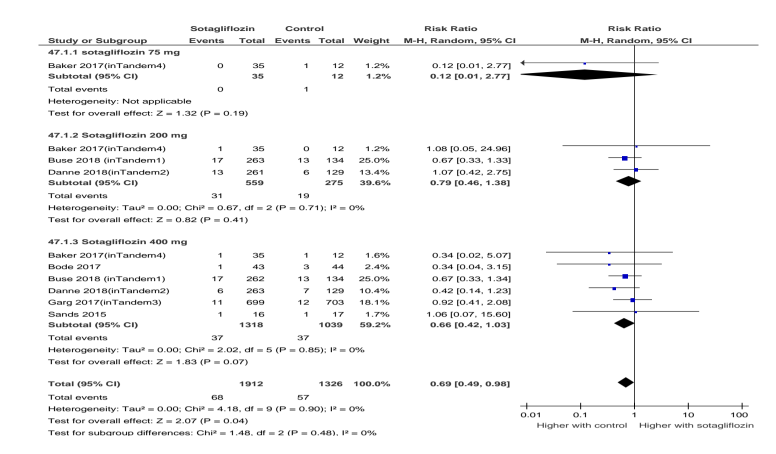


Figure 5. Forest plot of comparison: Sotagliflozin, outcomes: hypoglycemia, severe hypoglycaemia and diabetic ketoacidosis (DKA).

Panel A: outcome: hypoglycemia rate (events per patient-year)



Panel B: outcome: incident severe hypoglycemia

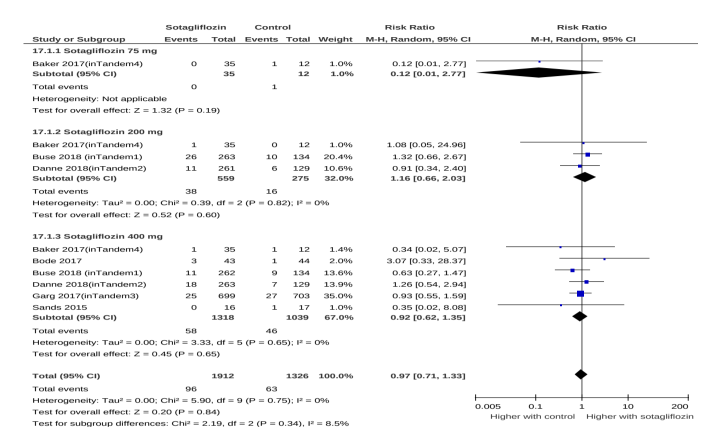


Panel C: outcome: incident DKA

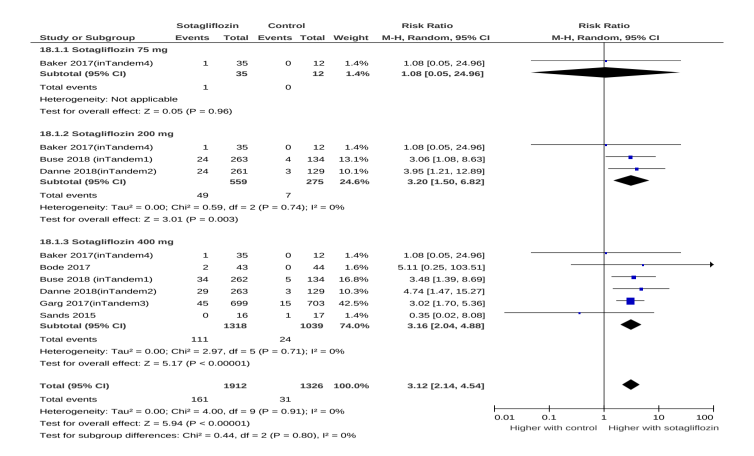
	Sotaglif	lozin	Contr	ol		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% CI	M-H, Random, 95% CI
14.1.1 Sotagliflozin 75 mg							
Baker 2017(inTandem4)	1	35	0	12	5.1%	1.08 [0.05, 24.96]	
Subtotal (95% CI)		35		12	5.1%	1.08 [0.05, 24.96]	
Total events	1		0				
Heterogeneity: Not applicab	le						
Test for overall effect: $Z = 0$.	.05 (P = 0	.96)					
14.1.2 Sotagliflozin 200 mg	9						
Baker 2017(inTandem4)	1	35	0	12	5.1%	1.08 [0.05, 24.96]	-
Buse 2018 (inTandem1)	9	263	1	134	11.8%	4.59 [0.59, 35.82]	
Danne 2018(inTandem2)	6	261	0	129	6.0%	6.45 [0.37, 113.62]	
Subtotal (95% CI)		559		275	22.9%	3.65 [0.83, 15.94]	
Total events	16		1				
Heterogeneity: Tau ² = 0.00;	$Chi^2 = 0.8$	80, df = 2	2 (P = 0.6)	7); I ² =	0%		
Test for overall effect: $Z = 1$.	.72 (P = 0	.09)					
14.1.3 Sotagliflozin 400 mg	9						
Baker 2017(inTandem4)	1	35	0	12	5.1%	1.08 [0.05, 24.96]	
Bode 2017	0	43	1	44	4.9%	0.34 [0.01, 8.14]	· ·
Buse 2018 (inTandem1)	11	262	0	134	6.2%	11.81 [0.70, 198.82]	
Danne 2018(inTandem2)	9	263	0	129	6.2%	9.36 [0.55, 159.50]	-
Garg 2017(inTandem3)	21	699	4	703	44.0%	5.28 [1.82, 15.30]	
Sands 2015	2	16	0	17	5.7%	5.29 [0.27, 102.49]	
Subtotal (95% CI)		1318		1039	72.1%	4.41 [1.92, 10.12]	
Total events	44		5				
Heterogeneity: Tau ² = 0.00;	$Chi^2 = 4.1$	8, df = 5	5 (P = 0.5	2); I ² =	0%		
Test for overall effect: $Z = 3$.	.50 (P = 0	.0005)					
Total (95% CI)		1912		1326	100.0%	3.93 [1.94, 7.96]	
Total events	61		6				
Heterogeneity: Tau ² = 0.00;	$Chi^2 = 5.7$	5, df = 9	9 (P = 0.7)	6); I ² =	0%		0.01 0.1 1 10 100
Test for overall effect: $Z = 3$.	.80 ($P = 0$.0001)					Higher with control Higher with sotagliflozin
Test for subgroup difference	s: Chi² =	0.73, df	= 2 (P = 0	0.69), I ²	9 = 0%		

Figure 6. Forest plot of comparison: Sotagliflozin, outcome: Urinary Tract Infections (UTIs), Genital Tract Infections (GTIs), diarrhea and volume depletion events

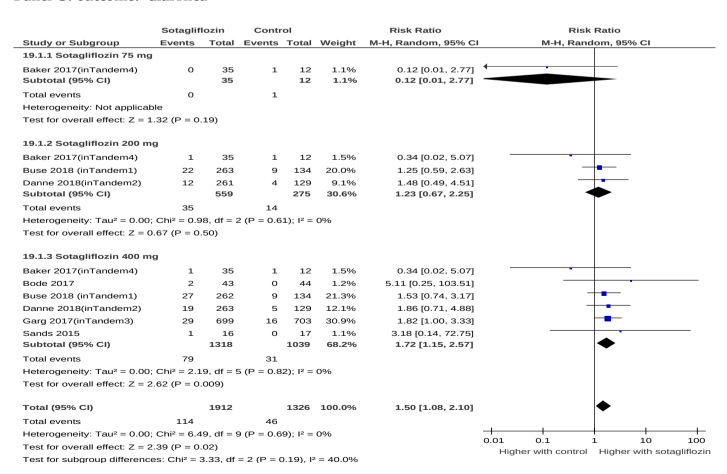
Panel A: outcome: UTIs



Panel B: outcome: GTIs



Panel C: outcome: diarrhea



Panel D: outcome: volume depletion events

	Sotaglif		Conti			Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% CI	M-H, Random, 95% CI
44.1.1 Sotagliflozin 75 mg							
Baker 2017(inTandem4) Subtotal (95% CI)	1	35 35	0	12 12	4.8% 4.8 %	1.08 [0.05, 24.96] 1.08 [0.05, 24.96]	
Total events	1		0				
Heterogeneity: Not applicat	ole						
Test for overall effect: $Z = 0$.	05 (P = 0.9	96)					
44.1.2 Sotagliflozin 200 mg	ı						
Baker 2017(inTandem4)	1	35	0	12	4.8%	1.08 [0.05, 24.96]	-
Buse 2018 (inTandem1)	8	263	2	134	20.2%	2.04 [0.44, 9.46]	- •
Danne 2018(inTandem2)	6	261	1	129	10.7%	2.97 [0.36, 24.37]	- - ·
Subtotal (95% CI)		559		275	35.7%	2.09 [0.66, 6.64]	
Total events	15		3				
Heterogeneity: Tau² = 0.00; Test for overall effect: Z = 1.		•	? (P = 0.8	7); I² = I	0%		
	,						
44.1.3 Sotagliflozin 400 mg	,	0.5		4.0	4.00/	4 00 10 05 04 001	
Baker 2017(inTandem4)	1	35	0	12	4.8%	1.08 [0.05, 24.96]	
Bode 2017	1	43	1	44	6.3%	1.02 [0.07, 15.84]	
Buse 2018 (inTandem1)	4 2	262 263	2	134 129	16.7% 5.2%	1.02 [0.19, 5.51]	
Danne 2018(inTandem2) Garg 2017(inTandem3)	13	203 699	0 2	703	21.5%	2.46 [0.12, 50.91] 6.54 [1.48, 28.86]	_ <u> </u>
Sarg 2017 (in Fanderns) Bands 2015	13	16		17	4.8%	3.18 [0.14, 72.75]	
Subtotal (95% CI)	'	1318	U	1039	59.5%	2.38 [0.97, 5.82]	
Total events	22	1310	5	1055	33.370	2.30 [0.31, 3.02]	
Heterogeneity: Tau² = 0.00;		Q df = F	_	3) · I≅ — I	n 04.		
Test for overall effect: Z = 1.			, , — o.o	∪ _{/1} 1 − 1	J 70		
Total (95% CI)		1912		1326	100.0%	2.19 [1.10, 4.36]	
	20	1912		1320	100.0%	2. 19 [1. 10, 4.30]	
Total events	38	0 46 5	8	41.17	0.07		
Heterogeneity: Tau² = 0.00;) (P = 0.9	1); ==	U 76		0.02 0.1 1 10
Test for overall effect: $Z = 2$.	•		2.00		2 000		Higher with control Higher with sotaglif
Test for subgroup differenc	es: Chi*=	U.23, df	= 2 (P = 1)	U.89), ľ	*= U%		

Supplementary text

Online Search strategies

Medline and Cochrane Central Register of Controlled Trials (Central):

1. randomized controlled trial.pt 2. controlled clinical trial.pt 3. randomized.tw 4. clinical trial/ 5. randomly.ab 6. trial.ti 7. placebo.tw 8. 1 OR 2 OR 3 OR 4 OR 5 OR 6 OR 7 9. sodium-glucose transporter 1/2/ 10. sodium-glucose transporter 1/2.tw 11. SGLT1/2.tw 12. SGLT-1/2.tw 13.dual SGLT.tw 14. Sotagliflozin.tw OR LX4211.tw OR LP802034.tw OR SAR439954.tw OR Zynquista.tw 15. LX4211.tw 16. Sotagliflozin.tw 17. LP802034.tw 18. SAR439954.tw 19. Zynquista.tw

20. 9 OR 10 OR 11 OR 12 OR 13 OR 14 OR 15 OR 16 OR 17 OR 18 OR 19

21.8 and 20

EMBASE

- 1. 'randomized controlled trial'/exp OR 'randomized controlled trial'
- 2. 'sodium glucose cotransporter 1/2'/exp OR 'sodium glucose cotransporter 1/2'
- 3. 'sodium glucose cotransporter 1/2 inhibitor'/exp OR 'sodium glucose cotransporter 1/2 inhibitor'
- 4.'sotagliflozin'/exp OR 'sotagliflozin' OR 'LX4211' OR 'LP802034' OR 'SAR439954' OR 'Zynquista'
- 5. 2 OR 3 OR 4
- 22. 1 AND 5

CLINICALTRIALS.GOV

- 1. Sodium-GlucoseTransporter 1/2
- 2. SGLT-1/2
- 3. Sotagliflozin
- 4. LX4211
- 5 LP802034
- 6. SAR439954
- 7. Zynquista

US FDA, EMA, databases

- 1. Sodium-Glucose Transporter 1/2
- 2. SGLT-1/2
- 3. Sotagliflozin
- 4. LX4211
- 5. LP802034,
- 6. SAR439954
- 7. Zynquista

International and National Trial registries search results

- -World Health Organization-International Clinical Trials Registry Platform (http://apps.who.int/trialsearch/): 82 records
- -ClinicalTrials.gov(https://www.clinicaltrials.gov/ct2/home): 37 records
- Cochrane CENTRAL Register of Controlled Trials (https://www.cochranelibrary.com/central/about-central): 47 records
- European Union(EU) Clinical Trials Register (https://www.clinicaltrialsregister.eu/): 13 records
- -ISRCTN (http://www.isrctn.com/): 0 results
- -Epistemonikos (https://www.epistemonikos.org/): 0 records
- -Health Canada Clinical Trial Database (http://www.hc-sc.gc.ca/dhp-mps/prodpharma/databasdonclin/index-eng.php): 11 records
- -German Clinical Trials Register (https://drks-neu.uniklinik-freiburg.de/drks_web/): 0 results
- -Netherlands Trial Register (Dutch) (http://www.trialregister.nl/trialreg/index.asp): 0 results
- -Swiss National Clinical Trials Portal (http://www.kofam.ch/en/swiss -clinical-trials-portal.html) 6 results
- -Australian New Zealand Clinical Trials Registry (http://www.anzctr.org.au/]: 4 records
- -ChineseClinical Trial Register (http://www.chictr.org.cn/enIndex.aspx): 0 records
- -Clinical Trials Registry–India(http://ctri.nic.in/): 1 record
- -Iranian Registry of Clinical Trials (http://www.irct.ir/): 0 records
- **-Japan Primary Registries Network** (http://rctportal.niph.go.jp/): 0 records
- -ClinicalResearch Information Service, Republic of Korea

(https://cris.nih.go.kr/cris/en/use_guide/cris_introduce.jsp): 0 records

- -Philippine Health Research Registry (http://registry. healthresearch.ph/): 0 results
- -Sri Lanka Clinical Trials Registry (http://www.slctr.lk/): 0 records
- -Thai Clinical Trials Registry (http://www.clinicaltrials.in.th/): 0 records
- -Brazilian Clinical Trials Registry (http://www.ensaiosclinicos.gov.br/): 0 records

- -Public Cuban Registry of Clinical Trials (http://registroclinico.sld.cu/en/home): 0 records
- -Peruvian Registry of Clinical Trials (http://www.ins.gob.pe/ensayosclinicos/): 0 records
- -Pan AfricanClinical Trials Registry (http://www.pactr.org/): 0 records
- -South African National Clinical Trials Register: (http://www.sanctr.gov.za/): 0 records
- -Tanzania Clinical Trial Registry (http://www.tzctr.or.tz/): 0 records

Regulatory Agencies sites search results

US Food and Drug Administration (FDA)

https://search.usa.gov/search?utf8=%E2%9C%93&affiliate=fda&query=sotagliflozin&commit=Search: 6 results

European Medicines Agency (EMA)

https://www.ema.europa.eu/en/search/search/search_api_views_fulltext=sotagliflozin): 49 results

Japanese Pharmaceutical and Medical Devices Agency(PMDA)

https://ss.pmda.go.jp/en_all/search.x?q=sotagliflozin&ie=UTF-8&page=1&x=30&y=11:

0 results

Definitions

Hypoglycemia: blood glucose levels ≤ 70 mg/dL documented on self-monitoring blood glucose,
 regardless of symptoms. We evaluated hypoglycaemia asnumber of hypoglycemic events per patient-year ⁷⁴
 Severe hypoglycemia: an event consistent with hypoglycemia (regardless of whether biochemical documentation of a low glucose value was obtained) when any of the following three conditions occurred:

- the patient have an episode of suspected hypoglycemia treated with any form of carbohydrate or with glucagon that required the assistance of others to treat, because the neurologic impairment was severe enough to prevent self-treatment in the opinion of those providing assistance to treat.
- the patient lost consciousness during the episode
- the patient had a seizure during the episode

Diabetic ketoacidosis (DKA): DKA was diagnosed based on evidence of anion-gap metabolic acidosis related to excessive ketone production without a satisfactory alternative cause for anion-gap acidosis, as outlined in Kitabchi et al 2009⁷⁵.

Renal event: defined according to the following Medical Dictionary for Regulatory Activities preferred terms:

Acute prerenal failure; Anuria; Azotemia; Blood creatine abnormal; Blood creatine decreased; Blood creatinine increased; Blood creatinine abnormal; Blood urea increased; Blood urea nitrogen/creatinine ratioincreased

Coma uremic; Computerized tomogram kidney abnormal; Creatine urine abnormal; Creatine urine decreased; Creatine urine increased; Creatinine renal clearance abnormal

Creatinine renal clearance decreased; Creatinine urine abnormal; Creatinine urine decreased

Creatinine urine increased; Cystatin C abnormal; Cystatin C increased, Diabetic end stage renal disease;

Glomerular filtration rate abnormal; Glomerular filtration rate decreased;

Glomerular filtration rate increased; Hypercreatinemia; Hyperparathyroidism secondary

Inulin renal clearance abnormal; Inulin renal clearance decreased; Kidney fibrosis;

Nephrogenic anemia; Nitrogen balance negative; Edema due to renaldisease;

OliguriaPericarditis uremicPhenolsulfonphthalein test abnormal; Postoperative renal failure

Prerenal failure; Renal cortical necrosis; Renal disorder; Renal failure;

Renal failure acute;Renal failure chronic; Renal function test abnormal;Renal impairment;

Renal injury;Renalnecrosis;Renal papillary necrosis;Renal scan abnormal;Renal tubular acidosis;Renal tubular atrophy;Renal tubular disorder;Renal tubular necrosis;Ultrasound kidney

abnormal;Uremicacidosis;Uremicencephalopathy;UremicgastropathyUremic neuropathy;

Uremic pruritus;Urea renal clearance;Urea renal clearance decreased;Urea renal clearance

increased;Uridosis;Urine albumin/creatinine ratio abnormal;Urine albumin/creatinine ratio decreased;

Urine albumin/creatinine ratio increased;Urineoutput;Urine output decreased;

Urine protein/creatinine ratio increased.

Volume depletion event: defined according to the following Medical Dictionary for Regulatory Activities preferred terms:

Acute prerenal failure;Blood pressure abnormal;Blood pressure ambulatory abnormal;Blood pressure decreased;Blood pressure diastolic decreased;Blood pressure fluctuation;Blood pressure immeasurable;Blood pressure inadequately controlled;Blood pressure orthostasisabnormal;Blood pressure orthostatic decreased;Blood pressure systolic abnormal;Blood pressure systolic decreased;Blood pressure systolic inspiratorydecreased;Brachial pulse abnormal; Brachial pulse decreased;Blood pressure systolic inspiratorydecreased;Brachial pulse abnormal; Brachial pulse decreased;Bun/creatinine ratio increased;Capillary nail refill test abnormal;Cardiac index abnormal;Cardiac index decreased;Cardiac output decreased;Cardiovascularinsufficient;Carotid pulse abnormal;Carotid pulse decreased;Central venous pressure abnormal;Central venous pressure decreased;Circulatorycollapse;Decreasedventricularpreload;Dehydration;Diastolichypotension;Femoral

pulse abnormal; Femoral pulse decreased; Hemodynamic test abnormal; Heart rate abnormal; Heart rate decreased:

Heart rate increased; Hypoperfusion; Hypotension; Hypovolemia; Hypovolemic shock;

Labile blood pressure;Left ventricular end-diastolic pressuredecreased;Maximum heart rate decreased;
Mean arterial pressure decreased;Orthostatic heart rate response increased;Orthostatic hypotension;
Orthostatic intolerance;Pedal pulse abnormal;Pedal pulse decreased;Peripheral circulatory
failure;Peripheral coldness;Peripheral pulse decreased;Popliteal pulse abnormal;Popliteal pulse decreased;

Prerenal failure;Presyncope;Pulseabnormal;Pulseabsent;Pulse pressure abnormal;Pulse pressure decreased;Pulse volume decreased;Pulse waveform abnormal;Radial pulse abnormal;Radial pulse decreased;Renalischemia;Schellingtest;Shock;Syncope;Thirst;Tilt table test positive;Urine albumin/creatinine ratio increased;Urine flow decreased;Urine output decreased;
Urine protein/creatinine ratio increased;Vascular test abnormal;Venous pressure abnormal;
Venous pressure decreased;Venous pressure jugular abnormal;Venous pressure jugular decreased;
Volume blood decreased.

Acidosis-related adverse event

Adverse events that satisfy the trigger terms for metabolic acidosis, which are the following Medical Dictionary for Regulatory Activities preferred terms: acetonemia, acidosis, acidosis hyperchloremic, blood ketone body, blood ketone body increased, blood ketone body present, DKA, diabetic hyperglycemia, coma, diabetic ketoacidotichyperglycemic diabetic metabolic decompensation, diabetic coma, hyperglycemic coma, hyperglycemic seizure, hyperglycemic unconsciousness, ketoacidosis, ketosis, lactic acidosis, metabolic acidosis, renaltubularacidosis, uremic acidosis, urine ketone body, and urine ketone body present.

Serious AEs

Serious adverse events were defined as serious if they resulted in death, a life-threaten, patient hospitalization, a persistent or significant incapacity or substantial disruption of the ability to conduct normal life functions, or if they required medical intervention to prevent one of the outcomes listed above. For this meta-analysis, serious AEs were defined as the number of participants experiencing death, cancer (all cancers, bladder cancer, breast cancer), MACE, severe hypoglycaemia, serious acidosis-related adverse events...

Management of missing data.

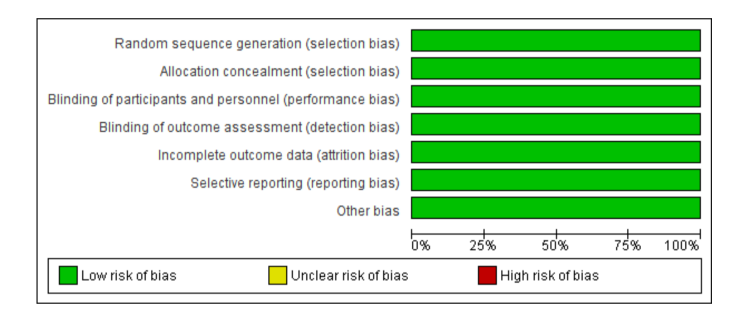
Missing data were managed by contacting via e-mail the corresponding authors of the RCTs. Where this was unsuccessful, we planned to calculate missing data from the raw numbers given in tables and/or estimated from bar charts. For missing standard deviations of mean change in parameters, and where the p value was provided for a comparison between treated and control groups, we planned to calculate the standard deviation by converting the p value into a t value with appropriate degrees of freedom, and then calculating standard error and standard deviation. If neither the standard deviations nor the p values were supplied, we planned to impute a standard deviation from studies with similar measurement methods, duration and measurement error was used if available1 and tested in a sensitivity analysis and reported if the estimate differed meaningfully from previous estimates. If no similar studies were available, a narrative approach would have been used to summarize the data

Supplementary Figures

Supplementary Figure 1. Risk of bias summary: risk of bias item for each included RCT according to Cochrane Risk-of-Bias Tool

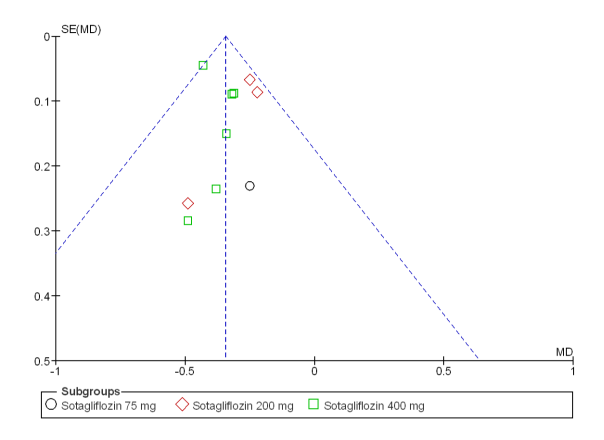
	Random sequence generation (selection bias)	Allocation concealment (selection bias)	Blinding of participants and personnel (performance bias)	Blinding of outcome assessment (detection bias)	Incomplete outcome data (attrition bias)	Selective reporting (reporting bias)	Other bias
Baker 2017(inTandem4)	•	•	•	•	•	•	•
Bode 2017	•	•	•	•	•	•	•
Buse 2018 (inTandem1)	•	•	•	•	•	•	•
Danne 2018(inTandem2)	•	•	•	•	•	•	•
Garg 2017(inTandem3)	•	•	•	•	•	•	•
Sands 2015	•	•	•	•	•	•	•

Supplementary Figure 2. Risk of bias graph: each risk of bias item is presented as percentages across all included RCTs.

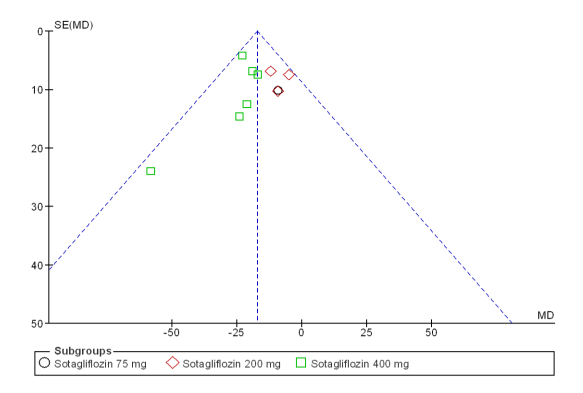


Supplementary Figure 3.

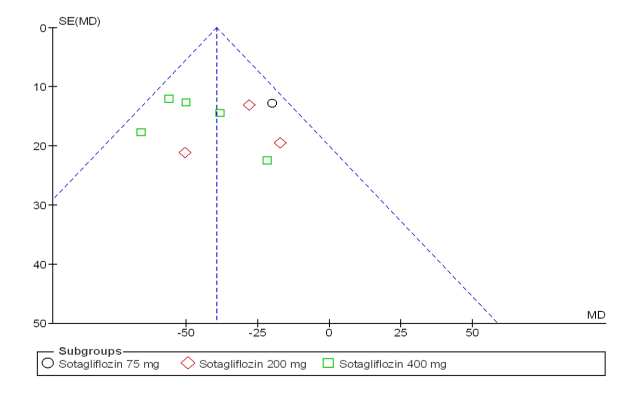
Panel A Funnel plot of comparison: HbA1c(%) outcome: HbA1c(%).



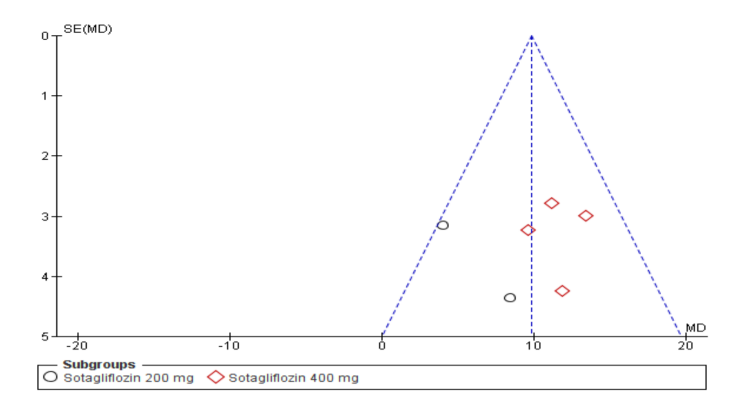
Panel B Funnel plot of comparison: Fasting plasma glucose (FPG; (mg/dL) outcome: FPG(mg/dL).



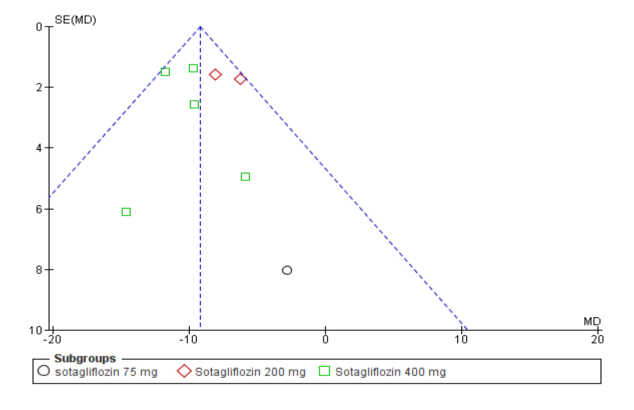
Panel C. Funnel plot of comparison: 2-hr postprandial plasma glucose(PPG) for outcome: 2hr-PPG.



Panel D. Funnel plot of comparison: % time-in-range (70-180 mg/dL) for outcome: % time-in-range

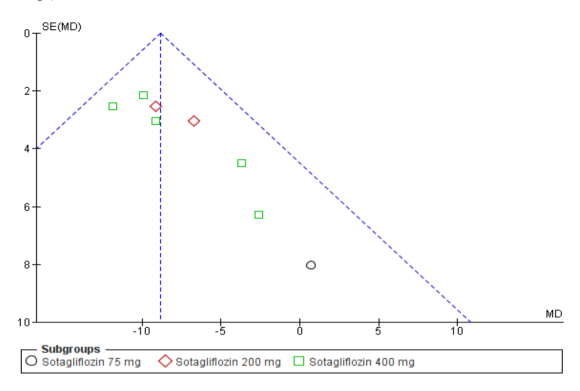


Panel E. Funnel plot of comparison: total daily insulin dose, outcome: total daily insulin dose(% change)

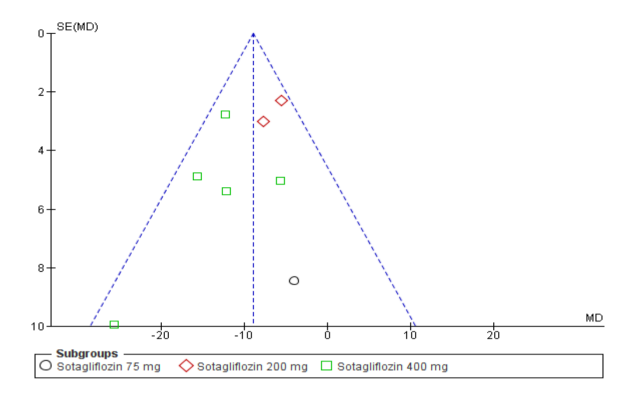


Panel F. Funnel plot of comparison: basal daily insulin dose, outcome: basal daily insulin dose(%

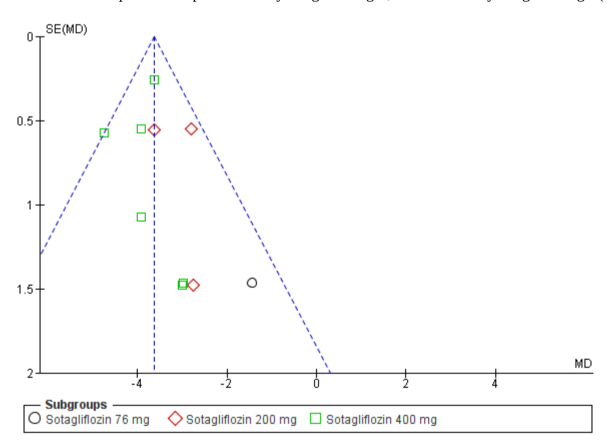
change)



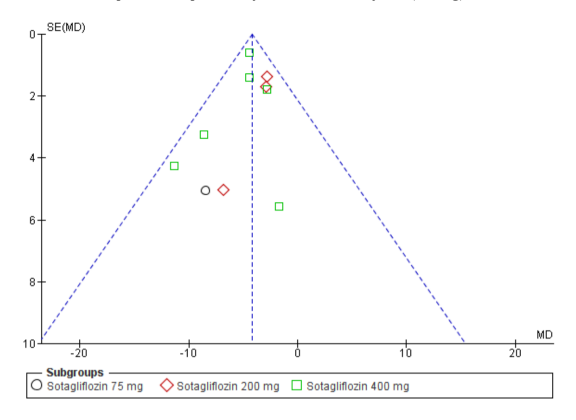
 $\textbf{Panel G}. \ \textbf{Funnel plot of comparison:} \ \ \textbf{bolus daily insulin dose, outcome:} \ \ \textbf{bolus daily insulin dose} (\% \ \ \textbf{change})$



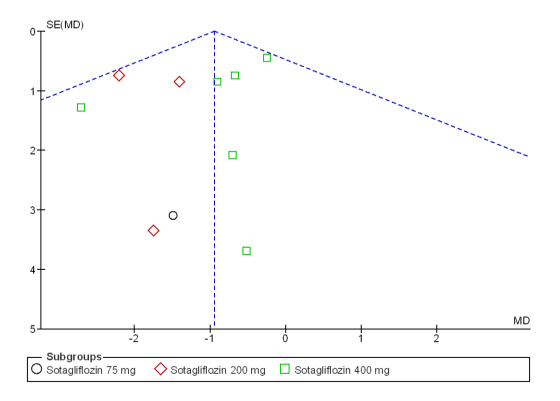
Panel H. Funnel plot of comparison: body weight changes, outcome: body weight changes(%)



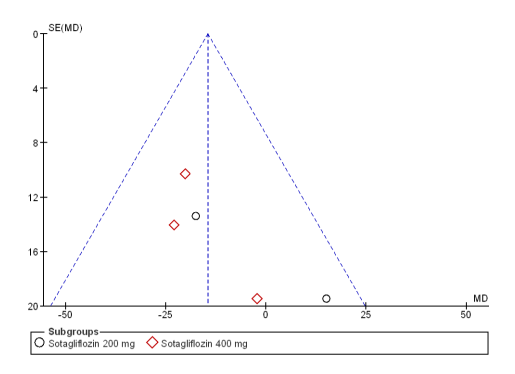
Panel I. Funnel plot of comparison: sys BP, outcome: sys BP(mmHg)



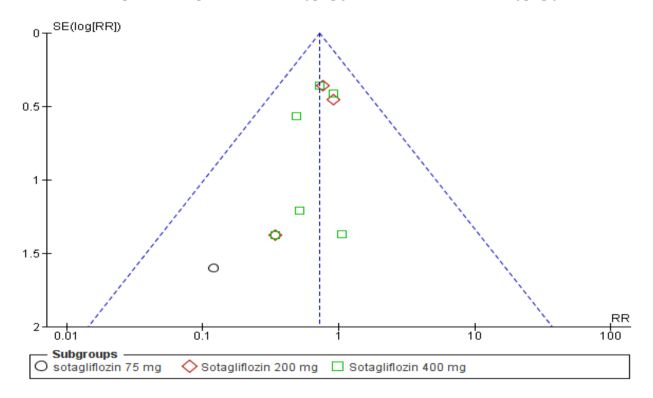
Panel L. Funnel plot of comparison: eGFR changes, outcome: eGFR changes(ml/min/1.73 m2)



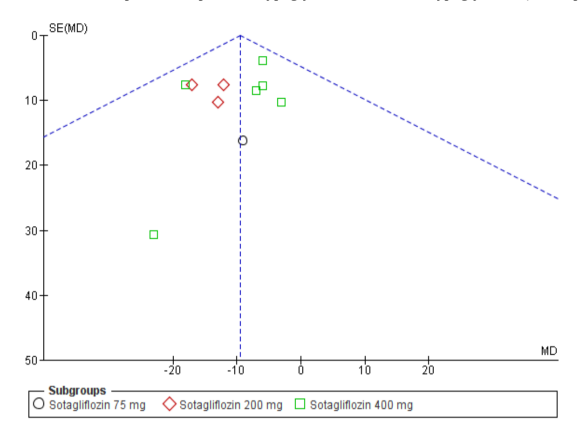
Panel M. Funnel plot of comparison: urinary A/C ratio, outcome: albumin/creatinine ratio(mg/g).



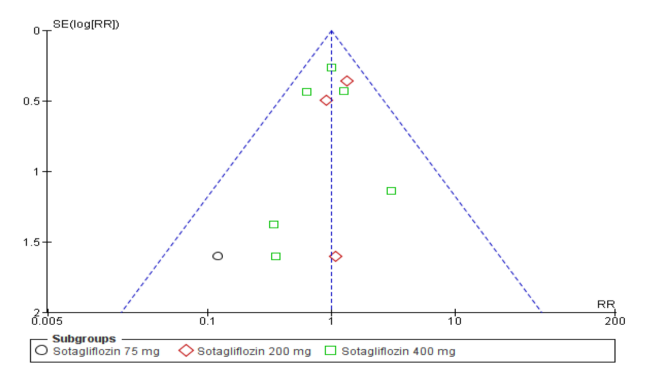
Panel N. Funnel plot of comparison: severe hypoglycemia, outcome: severe hypoglycemia.



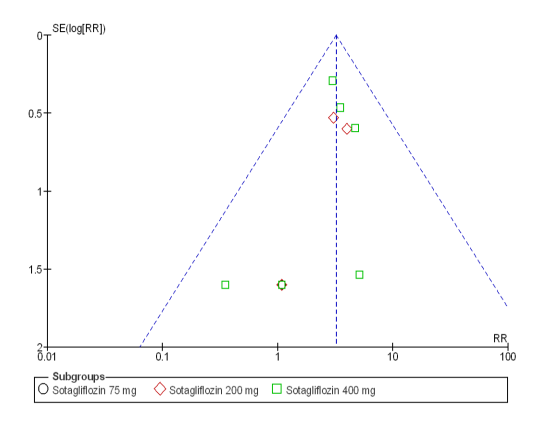
Panel O. Funnel plot of comparison: hypoglycemia, outcome: hypoglycemia (events per patient-year).



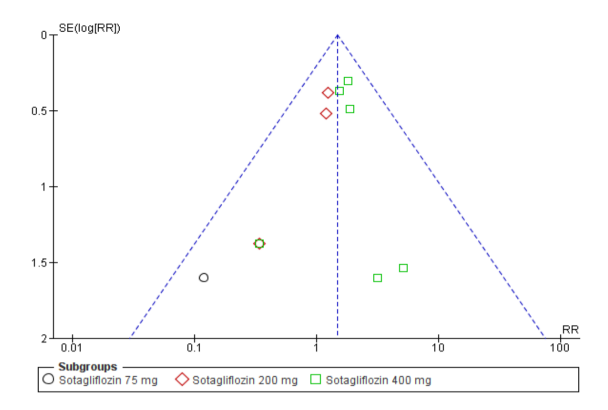
Panel P. Funnel plot of comparison: urinary tract infections, outcome: urinary tract infections.



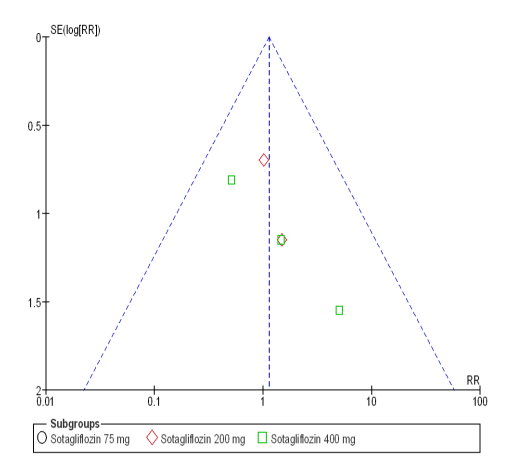
Panel Q. Funnel plot of comparison: genital tract infections, outcome: genital tract infections.



Panel R. Funnel plot of comparison: diarrhea, outcome: diarrhea

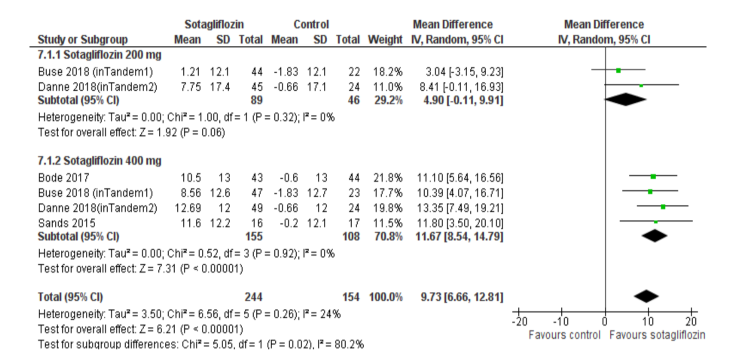


Panel S. Funnel plot of comparison: MACE, outcome: MACE



Supplementary Figure 4. Forest plot of comparison: Sotagliflozin, outcome: Continuous Glucose Monitoring (CGM) parameters.

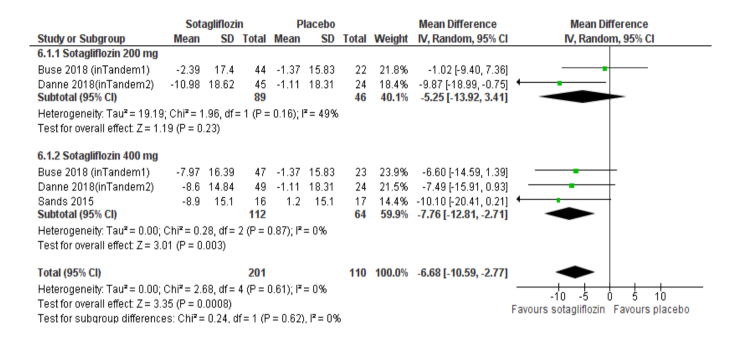
Panel A: outcome: time-in-range (%)



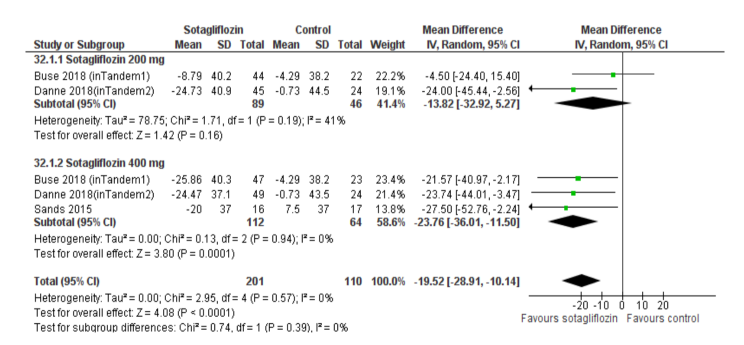
Panel B: outcome: average daily glucose (mg/dL)

	Sotag	glifloz	in	C	ontrol			Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
5.1.1 Sotagliflozin 200 mg									
Buse 2018 (inTandem1)	-2.97	22	44	1.93	22.1	23	22.2%	-4.90 [-16.03, 6.23]	
Danne 2018(inTandem2)	-10	27	45	1.92	26	24		-11.92 [-24.97, 1.13]	_
Subtotal (95% CI)			89			47	39.8%	-7.85 [-16.32, 0.61]	-
Heterogeneity: Tau ² = 0.00; (Chi ² = 0.6	4, df:	= 1 (P =	0.42);1	z = 0%)			
Test for overall effect: $Z = 1.8$	32 (P = 0.0	07)							
5.1.2 Sotagliflozin 400 mg									
Buse 2018 (inTandem1)	-15.57	23	47	1.93	27.8	23	17.5%	-17.50 [-30.63, -4.37]	
Danne 2018(inTandem2)	-19.28	20	49	1.92	20	24	26.3%	-21.20 [-30.97, -11.43]	
Sands 2015	-14	20	16	5.9	20	17	16.4%	-19.90 [-33.55, -6.25]	
Subtotal (95% CI)			112			64	60.2%	-19.89 [-26.68, -13.09]	•
Heterogeneity: Tau ² = 0.00; (Chi ² = 0.2	0, df:	= 2 (P =	: 0.91); [² = 0%)			
Test for overall effect: $Z = 5.7$	74 (P < 0.0	0000	1)						
Total (95% CI)			201			111	100.0%	-15.09 [-21.40, -8.79]	•
Heterogeneity: Tau ² = 14.50;	Chi ² = 5.	56, d	f= 4 (P	= 0.23)	$J^2 = 23$	8%			-20 -10 0 10 2
Test for overall effect: $Z = 4.8$	69 (P < 0.0	0000	1)						20 -10 0 10 2 -20 -20 -20 -20 -20 -20 -20 -20 -20
Test for subgroup difference	s: Chi²=	4.72,	df = 1	P = 0.03	3), l² =	78.8%			ravours solagiiiloziii Favours ci

Panel C: outcome: Standard Deviation (SD) around average daily glucose (mg/dL)

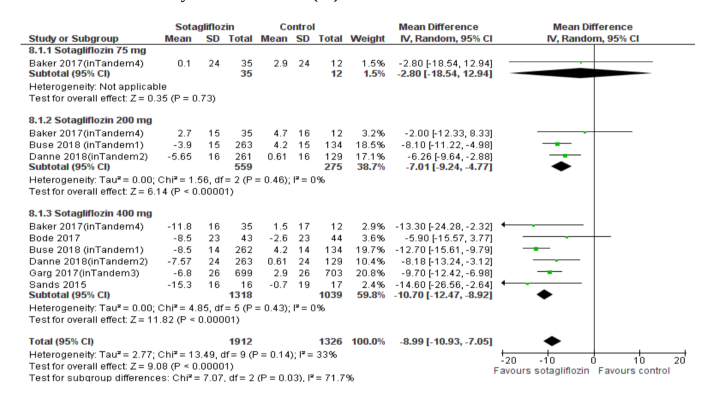


Panel D: outcome: mean amplitude of glucose excursion (MAGE) (mg/dL)

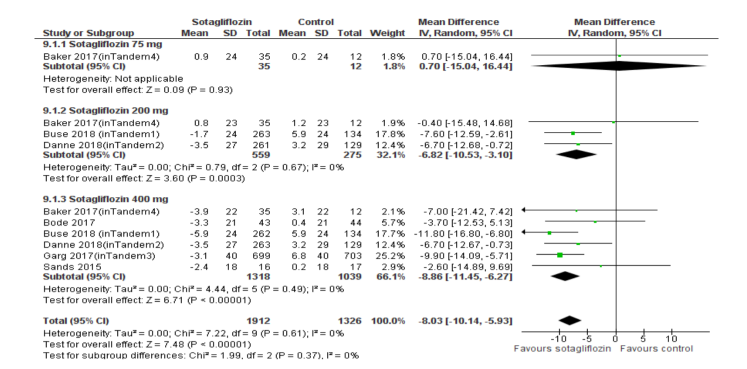


Supplementary Figure 5. Forest plot of comparison: Sotagliflozin, outcome: Daily total, basal and bolus insulin dose (%) changes from baseline.

Panel A: outcome: daily total insulin dose (%)



Panel B: outcome: daily basal insulin dose (%)



Panel C: outcome: daily bolus insulin dose (%)

		glifloz			ntro			Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% C	I IV, Random, 95% CI
10.1.1 Sotagliflozin 75 mg									
Baker 2017(inTandem4)	-0.9	26	35	3.1	25	12	3.2%	-4.00 [-20.56, 12.56]	
Subtotal (95% CI)			35			12	3.2%	-4.00 [-20.56, 12.56]	
Heterogeneity: Not applical									
Test for overall effect: $Z = 0$.47 (P = 0	.64)							
10.1.2 Sotagliflozin 200 m	g								
Baker 2017(inTandem4)	1.9	22	35	3.1	24	12	3.6%	-1.20 [-16.61, 14.21]]
Buse 2018 (inTandem1)	1.5	21	263	7	22	134	25.5%	-5.50 [-10.01, -0.99]]
Danne 2018(inTandem2)	-3.5	28	261	4.2	28	129	18.1%	-7.70 [-13.61, -1.79]	
Subtotal (95% CI)			559			275	47.3%	-6.05 [-9.54, -2.56]	•
Heterogeneity: Tau² = 0.00;	$Chi^2 = 0.$	74, df	= 2 (P :	= 0.69);	$ ^2 = 0$	0%			
Test for overall effect: Z = 3	.40 (P = 0)	.0007)						
10.1.3 Sotagliflozin 400 mg	g								
Baker 2017(inTandem4)	-9.1	25	35	4.5	25	12	3.2%	-13.60 [-29.99, 2.79]	1
Bode 2017	-8	24	43	-2.4	23	44	8.1%	-5.60 [-15.48, 4.28]	i
Buse 2018 (inTandem1)	-8.6	46	262	7	46	134	8.5%	-15.60 [-25.18, -6.02]	i ——
Danne 2018(inTandem2)	-7.9	50	263	4.2	50	129	7.2%	-12.10 [-22.63, -1.57]	i ————————————————————————————————————
Garg 2017(inTandem3)	-5.7	52	699	6.6	52	703	20.2%	-12.30 [-17.74, -6.86]	i —
Sands 2015	-32	29	16	-6.4	28	17	2.3%	-25.60 [-45.07, -6.13]	·
Subtotal (95% CI)			1318			1039	49.6%	-12.38 [-16.15, -8.61]	•
Heterogeneity: Tau² = 0.00;	$Chi^2 = 4$.	04, df	= 5 (P :	= 0.54);	$ ^2 = ($	0%			
Test for overall effect: Z = 6									
Total (95% CI)			1912			1326	100.0%	-9.14 [-12.17, -6.12]	•
Heterogeneity: Tau ^z = 4.02;	Chi ² = 10).94, d	lf = 9 (F	P = 0.28);	18%			
Test for overall effect: Z = 5									-20 -10 0 10 Favours sotagliflozin Favours
i est ior overall ellett. 🗸 🗕 🔾	.00 () - 0								

Supplementary Figure 6. Forest plot of comparison: Sotagliflozin, outcome: daily urinary

glucose excretion (UGE) (g/24 hr)

	Sota	glifloz	zin	Co	ntro	l		Mean Difference	Mean Dif	ference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% C	l IV, Randor	m, 95% CI
37.1.1 Sotagliflozin 75 mg										
Baker 2017(inTandem4) Subtotal (95% CI)	42	25	35 35	0.3	30	12 12	20.5% 20.5 %	41.70 [22.81, 60.59] 41.70 [22.81, 60.59]		•
Heterogeneity: Not applical	ble									
Test for overall effect: Z = 4	.33 (P < 0	0.000	1)							
37.1.2 Sotagliflozin 200 m	g									
Baker 2017(inTandem4) Subtotal (95% CI)	58	30	35 35	0.3	30	12 12	19.3% 19.3 %	57.70 [38.03, 77.37] 57.70 [38.03, 77.37]	- 1	•
Heterogeneity: Not applical	ble									
Test for overall effect: Z = 5	.75 (P < 0	0.000	01)							
37.1.3 Sotagliflozin 400 m	g									
Baker 2017(inTandem4)	71	33	35	0.3	30	12	18.6%	70.70 [50.51, 90.89]]	
Sands 2015	87	15	16	27	15	17	41.6%		·	-
Subtotal (95% CI)			51			29	60.2%	62.19 [53.06, 71.32]	j	•
Heterogeneity: Tau ² = 0.00;	$Chi^2 = 0$.86, d	lf=1 (P	= 0.35)	; l² =	0%				
Test for overall effect: Z = 1	3.35 (P <	0.00	001)							
Total (95% CI)			121			53	100.0%	57.79 [47.72, 67.87]	1	•
Heterogeneity: Tau ² = 36.23	3; Chi² =	4.53,	df = 3 ($P = 0.2^{\circ}$	1); 2:	= 34%				1 50
Test for overall effect: Z = 1									-50 -25 0	25 50
Test for subgroup differenc				2(P = 0.	16). I	² = 45.5	5%		Favours sotagliflozin	ravours control

Figure 7. Forest plot of comparison: Sotagliflozin vs. placebo, outcomes: eGFR changes over week 0-52: pooled analysis of inTandem1 and inTandem2 trials

Panel A: outcome: eGFR changes from baseline during week 0-24 (ml/min/1.73m²)

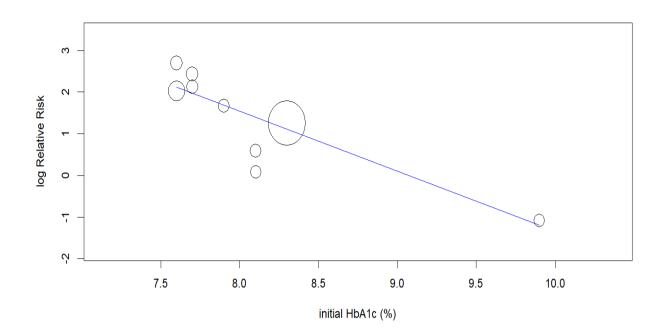
	Sotagliflozin Control			Mean Difference	Mean Difference				
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% C	IV, Random, 95% CI
50.1.1 Sotagliflozin 200 m	ng								
Buse 2018 (inTandem1)	-2.66	8	263	-0.2	8	134	22.0%	-2.46 [-4.12, -0.80]	_ -
Danne 2018(inTandem2)	-2.01	7	261	0.31	7	129	28.0%	-2.32 [-3.80, -0.84]	_ _ _
Subtotal (95% CI)			524			263	50.0%	-2.38 [-3.49, -1.28]	•
Heterogeneity: Tau ² = 0.00); $Chi^2 = 0$.02, c	lf = 1 (F	P = 0.90); l² =	: 0%			
Test for overall effect: $Z = 4$	4.23 (P <	0.000	01)						
50.1.2 Sotagliflozin 400 m	ng								
Buse 2018 (inTandem1)	-2.91	8	262	-0.2	8	134	22.0%	-2.71 [-4.38, -1.04]	_ -
Danne 2018(inTandem2)	-2.66	7	263	0.31	7	129	28.0%	-2.97 [-4.44, -1.50]	
Subtotal (95% CI)			525			263	50.0%	-2.86 [-3.96, -1.75]	•
Heterogeneity: Tau ² = 0.00); $Chi^2 = 0$.05, d	lf = 1 (F	P = 0.82); l² =	: 0%			
Test for overall effect: $Z = \frac{1}{2}$	5.07 (P <	0.000	001)						
Total (95% CI)			1049			526	100.0%	-2.62 [-3.40, -1.84]	•
Heterogeneity: Tau ² = 0.00); $Chi^2 = 0$.42, c	lf = 3 (F	P = 0.94); l² =	: 0%			
Test for overall effect: $Z = 0$	6.57 (P <	0.000	01)						-4 -2 0 2 4 Favours sotagliflozin Favours control
Test for subgroup difference	es: Chi² =	= 0.35	, df = 1	(P = 0.	55), I	² = 0%			ravours solagiilloziii ravours control

Panel B: outcome: eGFR changes from baseline during week 24-52 (ml/min/1.73m²)

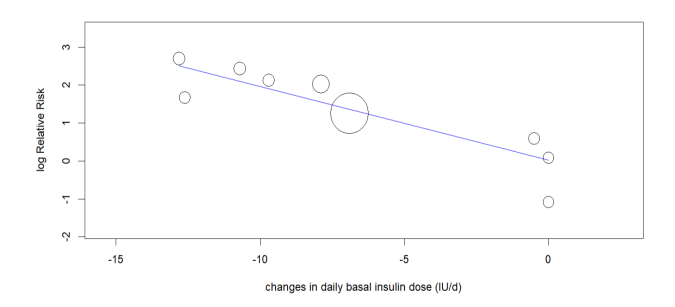
	Sota	gliflo	zin	Co	ontro	I		Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
51.1.1 Sotagliflozin 200 m	g								
Buse 2018 (inTandem1)	0.2	8	263	-0.98	8	134	23.0%	1.18 [-0.48, 2.84]	-
Danne 2018(inTandem2)	0.15	7	261	0.03	7	129	27.0%	0.12 [-1.36, 1.60]	
Subtotal (95% CI)			524			263	50.0%	0.59 [-0.52, 1.69]	
Heterogeneity: Tau ² = 0.00;	Chi ² = 0	.87, d	lf = 1 (F	P = 0.35); l² =	: 0%			
Test for overall effect: Z = 1	.04 (P =	0.30)							
51.1.2 Sotagliflozin 400 m	g								
Buse 2018 (inTandem1)	0.65	8	262	-0.98	8	134	22.9%	1.63 [-0.04, 3.30]	-
Danne 2018(inTandem2)	2.33	7	263	0.03	7	129	27.1%	2.30 [0.83, 3.77]	
Subtotal (95% CI)			525			263	50.0%	2.01 [0.90, 3.11]	
Heterogeneity: Tau ² = 0.00;	Chi ² = 0	.35, d	lf = 1 (F	P = 0.55); l² =	: 0%			
Test for overall effect: Z = 3	3.56 (P =	0.000	04)						
Total (95% CI)			1049			526	100.0%	1.30 [0.35, 2.25]	•
Heterogeneity: Tau ² = 0.30;	Chi ² = 4	.39, d	lf = 3 (F	P = 0.22); l² =	32%		-	
Test for overall effect: $Z = 2$.69 (P =	0.007	')						-2 -1 0 1 2 Favours control Favours sotagliflozin
Test for subgroup difference	es: Chi² =	3.17	, df = 1	(P = 0.	08), I	² = 68.5	5%		ravours control ravours sotagimozin

Supplementary Figure 8. Meta-regression analysis: regression plot of the effect of initial HbA1c(%) (panel A) and of changes in daily basal insulin dose(expressed as IU/d) from baseline (panel B) in relation to the risk (expressed as log risk ratio) of diabetic ketoacidosis (DKA). Each circle represents one comparison group, with the size of each circle representing the weight given to the group in meta-regression.

Panel A: effect of initial HbA1c (%) on the RR of DKA

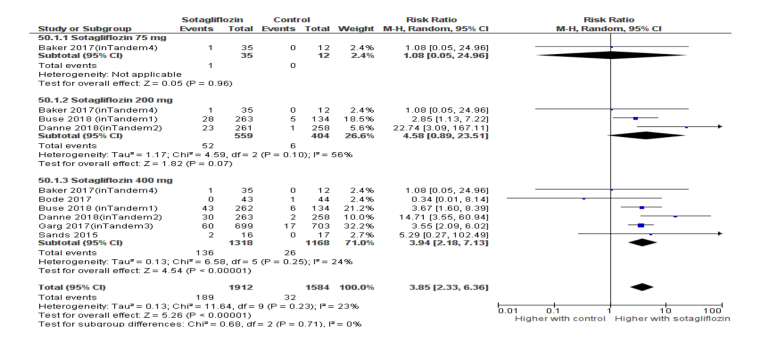


Panel B: effect of changes in daily basal insulin dose (IU/d) from baseline on the RR of DKA



Supplementary Figure 9. Forest plot of comparison: Sotagliflozin, outcome: incidence of acidosis-related and of major adverse cardiovascular events (MACE).

Panel A: outcome: acidosis-related adverse events



Panel B: outcome: major adverse cardiovascular events (MACE)

	Sotaglif		Contr			Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% CI	M-H, Random, 95% CI
22.1.1 Sotagliflozin 75 mg							
Baker 2017(inTandem4)	0	35	0	12		Not estimable	
Subtotal (95% CI)		35		12		Not estimable	
Total events	0		0				
Heterogeneity: Not applical							
Test for overall effect: Not a	pplicable						
22.1.2 Sotagliflozin 200 mg	g						
Baker 2017(inTandem4)	0	35	0	12		Not estimable	
3use 2018 (inTandem1)	6	263	3	134	43.8%	1.02 [0.26, 4.01]	
Danne 2018(inTandem2)	3	261	1	129	17.8%	1.48 [0.16, 14.11]	
Subtotal (95% CI)		559		275	61.6%	1.13 [0.35, 3.64]	-
Fotal events	9		4				
Heterogeneity: Tau² = 0.00;	$Chi^2 = 0.03$	B, df = 1	(P = 0.7)	$B); I^2 = 0$	0%		
Test for overall effect: $Z = 0$.	.20 (P = 0.8)	(4)					
22.1.3 Sotagliflozin 400 mg	a						
Baker 2017(inTandem4)	0	35	0	12		Not estimable	
Bode 2017	ō	43	Ō	44		Not estimable	
Buse 2018 (inTandem1)	1	262	3	134	17.8%	0.17 [0.02, 1.62]	
Danne 2018(inTandem2)	3	263	0	129	10.6%	3.45 [0.18, 66.24]	-
Garg 2017(inTandem3)	2	699	0	703	10.1%	5.03 [0.24, 104.55]	- -
3ands 2015	0	16	0	17		Not estimable	
Subtotal (95% CI)		1318		1039	38.4%	1.18 [0.12, 11.45]	
Fotal events	6		3				
Heterogeneity: Tau² = 2.09;	$Chi^2 = 4.19$	5, df = 2	! (P = 0.1)	3); $I^2 = 6$	52%		
Fest for overall effect: $Z = 0$.	.14 (P = 0.8	18)					
Total (95% CI)		1912		1326	100.0%	1.06 [0.40, 2.82]	-
Total events	15		7				
Heterogeneity: Tau² = 0.08;	$Chi^2 = 4.29$	5, df = 4	P = 0.3	7); $I^2 = 6$	6%		0.01 0.1 1 10 10
Fest for overall effect: $Z = 0$.11 ($P = 0.9$	11)					Higher with control Higher with sotaglifl
			= 1 (P = 1)				

Supplementary Tables

Supplementary Table 1. Characteristics (panel A) and Risk of Bias (panel B) of included trials.

ials	Renal function	eGFR≥60 ml/min/1.73 m²	eGFR≥45 ml/min/1.73 m²	eGFR≥60 ml/min/1.73 m²
ntrolled tr	Dropout rate(%)	%0	4.8%	0% 0% 2.7% 2.7%
ındomized co	Background treatment / Daily TID	Insulin 0.6 IU/kg Insulin 0.6 IU/kg	Insulin 0.8 IU/kg Insulin 0.8 IU/kg	Insulin 0.7 IU/kg Insulin 0.7 IU/kg Insulin 0.7 IU/kg
ncluded ra	Diabetes duration (yr)	18.5	12	24 23 27
istics of ii	Initial HbA1c (%)	7.94	9.9	8.0
Supplementary Table 1 panel A. Characteristics of included randomized controlled trials	Bodyweight (kg) / BMI (kg/m²)	74.2 kg / 27.1 kg/m² 72.7 kg / 26.2	83.8 kg / 29.0 80.7 kg / 27.9	84.1 kg / 29 81.9 kg / 28 78.1 kg / 27
ile 1 par	Gender (%M)	50	49	57 47 40 42
ry Tab	Age (yr)	45	23	45 47 42 48
upplementa	Study Arms	Sota 400 mg placebo	Sota 400 mg placebo	Sota 400 mg Sota 200 mg Sota 75 mg placebo
Σ̄	Study duration (week)	4	12	12

- <u>V</u>	participant s	33	87	141
Author		Sands 2015	Bode 2017	Baker 2017

Author	-N	Study	Study Arms	Age	Gender	(kg) /	Initial	Diabetes	Background	Dropout	Renal
	participants	duration		(yr)	(W%)	BIMI (kg/m²)	HbA1c (%)	duration	treatment /	rate	function
								(yr)	Daily TID		
		(week)							(IU/kg)		
Garg	1402	24	Sota 400 mg	43	51	82.4 kg/ 28.3	8.2	20	insulin 0.7 IU/kg	13%	eGFR≥45 ml/min/1.
2017			placebo	42	48	81.6/28.1	8.2	20	Insulin 0.7 IU/kg	11%	73 m²
Buse			Sota 400 mg	46	46	86.5 kg/29.6	7.6	24	Insulin 0.7 IU/kg	10%	eGFR≥45
2018	793	52	Sota 200 mg	47	48	86.9 kg/29.8	7.6	25	0.7 IU/kg	%6	ml/min/1.
			placebo	45	51	87.3 kg/29.6	7.5	24	Insulin 0.7 IU/kg	12%	
			Sota 400 mg	41	51	81.9 kg/ 27.9	7.7	19	Insulin 0.7 IU/kg	8%	eGFR≥45
Danne 2018	782	52	Sota 200 mg	42	53	81.9 kg/ 27.9	7.7	18	Insulin 0.7 IU/kg	8%	ml/min/1.
			placebo	40	52	81.1 kg/ 27.5	7.7	18	Insulin 0.7 IU/kg	%8	

als	Other:	sponsorship bias		Low risk.	The Robert and	Janice McNair	Foundation	funded the study			Low risk.	JDRF funded the	study			Low risk.	Industry funded		but no high risk	of bias feature	encountered*	
1 panel B Risk of Bias of included randomized controlled trials	Selective reporting			Low risk.	Prespecified outcomes	available on a clinical	trial database and all	reported in publication			Low risk.	Prespecified outcomes	available on a clinical	trial database and all	reported in publication	Low risk.	Prespecified outcomes	available on a clinical	trial database and all		reported in publication	
as of included rande	Incomplete outcome	data		Low risk.	No patients dropped	out					Low risk.	Low dropout rate:				Low risk.	Low dropout rate:					
el B Risk of Bi	Blinding pf	outcome assesment		Low risk.	Quadruple	masking	(Participant,	Care Provider,	Investigator,	Outcomes	Low risk.	Quadruple	masking			Low risk.	Quadruple	masking				
	Blinding of	participants and èpersonnel		Low risk.	Quadruple	masking	(Participant,	Care Provider,	Investigator,	Outcomes	Low risk.	Quadruple	masking			Low risk.	Quadruple	masking				
Supplementary Table	Allocation	concealment		Low risk.	Central	allocation,	web-based	randomization			Low risk.	Central	allocation,	web-based	randomization	Low risk.	Central	allocation,	web-based	randomization		
	Random	eouenbes	generation		Low risk.	7	Computer 1:ct	generated list			Low risk.	Computer	generated list	0		Low risk.	Computer	; ;	generated list			

Author Sands 2015	Bode 2017	Baker 2017
-------------------	--------------	---------------

	Panel B(continued). Risk of	B(continued). Risk of Bias of included randomized controlled trials	ized controlled trials	
Allocation	Blinding of participants and èpersonnel	Blinding pf outcome assesment	Incomplete outcome data	Selective reporting	Other: sponsorship bias
Low risk Central allocation, web- based randomization	Low risk Quadruple masking (Participant, Care Provider, Investigator, Outcomes Assessor)	Low risk Quadruple masking	Low risk Low dropout rate: Missing observations at EOT imputed as nonresponse.	Low risk Prespecified outcomes available on a clinical trial database and all reported in	Low risk Industry funded but no high risk of bias feature encountered*
Low risk Central allocation, web- based randomization	Low risk Quadruple Masking	Low risk Quadruple masking	Low risk Low dropout rate: Missing observations at EOT imputed as	Low risk Prespecified outcomes available on a clinical trial	Low risk Industry funded but no high risk of bias feature encountered*
Low risk Central allocation, web- based randomization	Low risk Quadruple Masking	Low risk Quadruple masking	Low risk Low dropout rate: Missing observations at EOT imputed as nonresponse.	Low risk Prespecified outcomes available on a clinical trial	Low risk Industry funded but no high risk of bias feature encountered*

Abbreviations: eGFR: estimated glomerular filtration rate; JDRF: Juvenile Diabetes Research

Foundation; Sota: sotagliflozin; TID: total insulin dose

^aInsulin dose optimization during the 6 weeks preceding randomization(target: FPG 80-130 mg/dL and 2hr-PPG<180 mg/dL)

*Assessment of sponsorship bias: in the presence of industry sponsorship, the following list of 8 items in trial designing, conducting or reporting, empirically linked by existing literature to biased outcomes in industry-funded trials and not captured by the Cochrane Risk of Bias domains, were assessed: if any one item was present, the trial was downgraded to "high risk of bias".

Item a: unclear clinical relevance of outcome measures: the clinical relevance of trial outcomes is not supported by international guidelines (American Association for the study of Diabetes-ADA or European Association for the Study of Diabetes-EASD guidelines).

Item b: if active comparator was used: inadequacy of doses timing or way of administration,

Item c: -deviations from study protocol or original protocol changes or amendments after trial initiation

Item d: post-hoc selection of the major findings and endpoints

Item e: use of last observation carried forward analysis to impute missing data

Item f:on-treatment outcome reporting /absence of data and safety monitoring board

Item g: absence of sponsor-independent statistician and data analysis

Item h: early trial termination before the endpoint recorded on clinical trial registries

Supplementary Table 2. Characteristics of randomized controlled trials(RTCs) with sotagliflozin excluded from this meta-analysis.

Phase 1	trials			
Official Title	Drug	N-participants	Duration	Year of
(author/ year of publication)	(dose)	(actual or	(week)	registration
ClinicalTrials.gov ID number		anticipated)		
				Status
Effect of Rifampicin on the Pharmacokinetics and	Sota	16	7.5	2017
Pharmacodynamics of Sotagliflozin	400 mg		7.3	Completed
NCT03063580	400 1116			Completed
Oral Contraceptive DDI Study	Sota	30	4	2015
NCT02494609	400 mg			Active, not
				recruiting
PK Study of Sotagliflozin in Subjects With Hepatic	Sota	32	1	2015
Impairment	400 mg			Completed
NCT02471274				
Interaction study to evaluate the Effects of Mefenamic Acid	Sota	16	8	2017
on the Pharmacokinetics and Pharmacodynamics of	400 mg			Completed
Sotagliflozin in Healthy Male and Female Subjects.				
NCT03070678				
A Drug to Drug Interaction Study of Sotagliflozin With	Sota	24	8	2016
Midazolam and Metoprolol.	200 mg			Completed
NCT02940379	or 400			
	mg			
Sotagliflozin Bioequivalence Study	Sota	76	9	2017
NCT03211195	200 mg			Completed
A Study to Evaluate the Effect of Food on the	Sota	14	9	
Pharmacokinetics of Sotagliflozin and to Explore the	200 mg			31/05/2017
Relative Bioavailability in Healthy Subjects. NCT03174548				Completed
A Drug to Drug Interaction Study of Sotagliflozin With	Sota	16	2	2018
Hydrochlorothiazide	200 mg			Completed
NCT03387657				

Comparison of Sotagliflozin Prototype Tablets With	Sota	12	9	2017
Reference Tablet in Healthy Subjects	400 mg			Completed
NCT03310944				
A Bioequivalence Study Testing Two Formulations of	Sota	58	14	2018
Sotagliflozin in Healthy Male and Female Subjects Under	200 mg			Active,
Fasted Conditions	or 400			not yet
NCT03776227	mg			recruiting,
A Phase 1, Open-label, Parallel-group Study to Evaluate	Sota	44	1	2015
Sotagliflozin Safety and Pharmacokinetics in Subjects With	200 mg			Active, Not
Varying Degrees of Renal Function, NCT02647918				recruiting
A Drug-Drug Interaction Study Between Sotagliflozin and	Sota	1	9	2018
Ramipril	400 mg			Completed
NCT03414723				

Randomized trials in type 2 diabetes mellitus(T2DM)

Official Title	Sota dose	N-participants	Duration	Year of
(author/ year of publication)		(actual or	(week)	registration
ClinicalTrials.gov ID		anticipated)		
				Status
A Randomized, Open-Label, Three-Way Crossover Study	Sota	15	4	2012
of Two Oral Formulations of LX4211 in Subjects With	150 mg			Completed
Type 2 Diabetes Mellitus	or 300			
NCT01188863	mg			
A Study to Evaluate the Pharmacodynamic Effects of Single-	Sota	18	3	2015
Dose Co-Administration of LX4211 With Januvia® in Type 2	400 mg			Completed
Diabetics				
NCT01441232				
Pharmacodynamic and Pharmacokinetic Effects of LX4211	Sota	31	1	2015
in Subjects With Type 2 Diabetes and Renal Impairment	400 mg			Completed
NCT01555008				
Safety and Efficacy of LX4211 With Metformin in Type 2	Sota 75	299	12	2015
Diabetes Patients With Inadequate Glycemic Control on	mg, 200			completed
Metformin	mg, 400			
NCT01376557	mg			

	l	1	l	
Efficacy and Safety of Sotagliflozin Versus Placebo in	Sota	369	24	29/11/2018
Chinese Patients With Type 2 Diabetes Mellitus Not	200 mg			Recruiting,
Adequately Controlled by Diet and Exercise				
NCT03760965	mg			
Efficacy and Safety of Sotagliflozin Versus Placebo in	Sota	369	24	Recruiting
Chinese Patients With Type 2 Diabetes Mellitus Not	200 mg			29/11/2018
Adequately Controlled by Metformin With or Without	or 400			
Sulfonylurea	mg			
NCT03761134				
Effect of Sotagliflozin on Cardiovascular Events in Patients	Sota	4000	32	Recruiting30
With Type 2 Diabetes Post Worsening Heart Failure	200 mg			/04/2018
(SOLOIST-WHF Trial)	or 400			
NCT03521934	mg			
Comparison of Pharmacodynamic Effects of Sotagliflozin	Sota	40	8	Recruiting
and Empagliflozin in T2DM Patients With Mild to Moderate	400 mg			06/03/2018
Hypertension				
NCT03462069				
Efficacy and Bone Safety of Sotagliflozin Dose 1 and Dose 2	Sota	360	24	Active, not
Versus Placebo in Subjects With Type 2 Diabetes Mellitus	200 mg			recruiting
Who Have Inadequate Glycemic Control. (SOTA-BONE Trial)	or 400			21/12/2017
NCT03386344	mg			
Efficacy and Safety of Sotagliflozin Versus Glimepiride and	Sota	930	52	Active, Not
Placebo in Subjects With Type 2 Diabetes Mellitus That Are	200 mg			recruiting
Taking Metformin Monotherapy(SOTA-GLIM trial)	or 400			02/11/2017
NCT03332771	mg			
Efficacy and Safety of Sotagliflozin versus Placebo and	Sota	700	26	Active, not
Empagliflozin in Subjects with Type 2 Diabetes Mellitus who	400 mg			recruiting
have Inadequate Glycemic Control while taking a DPP4				
Inhibitor Alone or with Metformin(SOTA-EMPA trial)				
NCT03351478				
Effect of Sotagliflozin on Cardiovascular and Renal Events in	Sota	1500	5 years	Active,
Patients with Type 2 Diabetes and Moderate Renal	200 mg			recruiting
Impairment Who Are at Cardiovascular Risk(SCORED trial)	vs. 400			04/10/2017
NCT03315143	mg			
				4

200 mg vs. sota 400 mg Sota 200 mg vs. sota 400 mg Sota 200 mg vs. sota 200 mg vs. sota	780	52	recruiting 2017 Active, Not recruiting 03/08/2017 Active, Not
Sota 200 mg vs. sota 400 mg Sota 200 mg			Active, Not recruiting 03/08/2017
Sota 200 mg vs. sota 400 mg Sota 200 mg			recruiting 03/08/2017
200 mg vs. sota 400 mg Sota 200 mg			recruiting 03/08/2017
200 mg vs. sota 400 mg Sota 200 mg			recruiting 03/08/2017
200 mg vs. sota 400 mg Sota 200 mg			recruiting 03/08/2017
vs. sota 400 mg Sota 200 mg	276	52	03/08/2017
400 mg Sota 200 mg	276	52	
Sota 200 mg	276	52	Active, Not
200 mg	276	52	Active, Not
200 mg	276	52	Active, Not
200 mg	276	52	Active, Not
			,
vs. sota			recruiting
			03/08/2017
400 mg			
Sota	500	26	Active,
400 mg			Not
			recruiting
			24/02/2017
Sota	400	26	Active, Not
400 mg			recruiting
			05/10/2016
Sota	500	26	Active, Not
200 mg			recruiting
vs. sota			05/10/2016
400 mg			
gestive I	Heart Failure	e	
rug	N-participants	Duration	Year of
			registration
į	anticipated)		
			Sstatus
41 Si	ota	s. sota 00 mg ota 500 ota 400 ota 400 ota 500 ota 500 ota 500 ota 500 ota 600 mg s. sota oo mg sestive Heart Failure ug N-participants	00 mg s. sota 00 mg ota ota 00 mg s. sota 00 mg s. sota 00 mg s. sota 00 mg sestive Heart Failure ug ose) (actual or anticipated)

Safety, Tolerability and Pharmacodynamic Activity of	Sota	81	5	Active,
Sotagliflozin in Hemodynamically Stable Patients With	200 mg			Recruiting
Worsening Heart Failure.				04/12/2017
NCT03292653	mg			

Abbreviations: UGE: urinary glucose excretion; T2D: type 2 diabetes mellitus; OAD: Oral Antidiabetic Agents;

Sota: sotagliflozin

Supplementary Table 3. Results of subgroup and sensitivity analysis.

Treatment duration		
Outcome	treatment duration ≤12	treatment duration >12 weeks
HbA1c (%)	-0.37 (-0.56, -0.18), I ² =0%, p=0.0001, N =5 comparisons, 261 participants	-0.36(-0.47, -0.26), I ² =12%, p<0.00001, N =5 comparisons, 2977 participants
FPG (mg/dL)	-16.74 (-28.49, -5.00), I ² =10%, p=0.005, N = 5, 261 participants	-16.77 (-23.05, -10.49), I ² =25%, p<0.00001, N=5, 2977 participants
2h-PPG (mg/dL)	-38.72 (-52.27, -25.16), I ² =20%, p<0.00001, N=5, 261 participants	-40.10(-63.73, -16.47), I ² =30%, p=0.001, N=5, 278 participants
Total insulin dose (IU/d)	-9.51 (-17.91, -1.81), I ² =0%, p=0.009, N=5, 261 participants	-9.16 (-11.40, -6.92), I ² =36%, p<0.00001, N=3, 2977 participants
Basal insulin dose (IU/d)	-5.33 [-10.49,-1.49], I ² =0%, p=0.03, N=3, 261 participants	-8.89 (-11.16, -6.61) I ² =0%, p<0.00001, N=5, 2977 participants

Bolus insulin dose	$[-13.77 [-23.04, -3.50] I^2 = 34\%,$	$-9.51 (-13.10, -5.92), I^2=24\%, p<0.00001,$
(IU/d)	p=0.0004, N =5, 261 participants	N=5, 2977 participants
Time-in-	11.31(6.75,15.87) I ² =0%, p<0.00001,	8.88(4.25, 13.51) I ² =36%, p=0.0002, N=4,
Range (%)	N=2, 120 participants	278 participants
Body weight	-2.63(-4.09, -1.17), I ² =0%, p=0.0004,	-3.67(-4.25, -3.10), I ² =0%, p<0.00001, N=5,
change (%)	N=5, 261 participants	2977 participants
Systolic BP (mmHg)	-8.65(-12.49, -4.81), I ² =34%,	-3.61(-4.55, -2.66), I ² =0%, p<0.00001,
	p=0.0004, N=5, 285 participants	N=5, 2977 participants
Diastolic BP (mmHg)	-2.13 (-4.00, -0.27), I ² =0%, p=0.02,	-1.36 (-1.93, -0.80), I ² =0%, p<0.00001,
	N=3, 285 participants	N=3, 2977 participants
eGFR	-2.26(-4.41, -0.11), I ² =0%, p=0.04,	-0.42(-1.15, 0.32), I ² =0%, p=0.26, N=5,
(ml/min/1.73 m ²)	N=5, 261 participants	2977 participants
Albumin-creatinine	No studies	-14.57(-26.87, -2.28), I ² =0%, p=0.02,
ratio (ACR)(mg/g)		N=3, 2977 participants
Hypoglycemia	-9.82(-16.00, -1.48), I ² =0%, p=0.01,	$-9.71(-15.05, -4.38), I^2=0\%, p<0.00001,$
(events per patient-	N=3, 261 participants	N=3, 2977 participants
year)		
Severe	$0.41(0.13, 1.28), I^2=0\%, p=0.12,$	$0.72(0.51, 1.04), I^2=0\%, p=0.08, N=5,$
hypoglycemia	N=5, 261 participants	2977 participants
JI - 8 J	$1.23(0.31 4.04) I^2-0% p-0.77$	$5.89(2.60, 13.36), I^2=0\%, p<0.00001,$
DKA	1.23(0.31, 4.34) 1 –070, p–0.77,	5.05(2.00, 15.50), 1 = 070, p \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \
DKA	, , , , , , , , , , , , , , , , , , , ,	N=5, 2977 participants
DKA UTI	N=5, 261 participants	
	N=5, 261 participants	N=5, 2977 participants
	N=5, 261 participants 0.70(0.20, 2.42), I ² =0%, p=0.57, N=5, 261 participants	N=5, 2977 participants 0.99(0.71, 1.37), I ² =0%, p<0.00001, N=5, 2977 participants
UTI	N=5, 261 participants 0.70(0.20, 2.42), I ² =0%, p=0.57, N=5, 261 participants	N=5, 2977 participants 0.99(0.71, 1.37), I ² =0%, p<0.00001, N=5, 2977 participants
UTI	N=5, 261 participants 0.70(0.20, 2.42), I ² =0%, p=0.57, N=5, 261 participants 1.21(0.30, 4.86), I ² =0%, p=0.79, N=35 261 participants	N=5, 2977 participants 0.99(0.71, 1.37), I ² =0%, p<0.00001, N=5, 2977 participants 3.36(2.27, 4.96), I ² =0%, p<0.00001, N=5,
UTI	N=5, 261 participants 0.70(0.20, 2.42), I ² =0%, p=0.57, N=5, 261 participants 1.21(0.30, 4.86), I ² =0%, p=0.79, N=35 261 participants	N=5, 2977 participants 0.99(0.71, 1.37), I ² =0%, p<0.00001, N=5, 2977 participants 3.36(2.27, 4.96), I ² =0%, p<0.00001, N=5, 2977 participants
UTI	N=5, 261 participants 0.70(0.20, 2.42), I ² =0%, p=0.57, N=5, 261 participants 1.21(0.30, 4.86), I ² =0%, p=0.79, N=35 261 participants 1.70(1.08, 2.77), I ² =0%, p=0.04, N=5, 261 participants	N=5, 2977 participants 0.99(0.71, 1.37), I ² =0%, p<0.00001, N=5, 2977 participants 3.36(2.27, 4.96), I ² =0%, p<0.00001, N=5, 2977 participants 1.59(1.12, 2.24), I ² =0%, p=0.009, N=5,
UTI GTI Diarrhea	N=5, 261 participants 0.70(0.20, 2.42), I ² =0%, p=0.57, N=5, 261 participants 1.21(0.30, 4.86), I ² =0%, p=0.79, N=35 261 participants 1.70(1.08, 2.77), I ² =0%, p=0.04, N=5, 261 participants	N=5, 2977 participants 0.99(0.71, 1.37), I ² =0%, p<0.00001, N=5, 2977 participants 3.36(2.27, 4.96), I ² =0%, p<0.00001, N=5, 2977 participants 1.59(1.12, 2.24), I ² =0%, p=0.009, N=5, 2977 participants
UTI GTI Diarrhea Volume depletion	N=5, 261 participants 0.70(0.20, 2.42), I ² =0%, p=0.57, N=5, 261 participants 1.21(0.30, 4.86), I ² =0%, p=0.79, N=35 261 participants 1.70(1.08, 2.77), I ² =0%, p=0.04, N=5, 261 participants 2.62 (1.18, 5.82), I ² =3%, p=0.02, N=5, 261 participants	N=5, 2977 participants 0.99(0.71, 1.37), I ² =0%, p<0.00001, N=5, 2977 participants 3.36(2.27, 4.96), I ² =0%, p<0.00001, N=5, 2977 participants 1.59(1.12, 2.24), I ² =0%, p=0.009, N=5, 2977 participants 1.37 (0.30, 2.19), I ² =0%, p=0.68, N=5, 2977 participants
UTI GTI Diarrhea Volume depletion events	N=5, 261 participants 0.70(0.20, 2.42), I ² =0%, p=0.57, N=5, 261 participants 1.21(0.30, 4.86), I ² =0%, p=0.79, N=35 261 participants 1.70(1.08, 2.77), I ² =0%, p=0.04, N=5, 261 participants 2.62 (1.18, 5.82), I ² =3%, p=0.02, N=5, 261 participants	N=5, 2977 participants 0.99(0.71, 1.37), I ² =0%, p<0.00001, N=5, 2977 participants 3.36(2.27, 4.96), I ² =0%, p<0.00001, N=5, 2977 participants 1.59(1.12, 2.24), I ² =0%, p=0.009, N=5, 2977 participants 1.37 (0.30, 2.19), I ² =0%, p=0.68, N=5,

Outcome	initial HbA1c levels	initial HbAlc levels
	< 8%	≥8%
HbA1c (%)	-0.27 (-0.35, -0.19), I ² =0%, p<0.00001,	-0.44(-0.52, -0.36), I ² =0%, p<0.00001, N =5
	N =5 comparisons, 1608 participants	comparisons, 1630 participants
FPG (mg/dL)	-14.77 [-23.25, -6.30], I ² =25%,	-19.83 [-26.51, -13.15], I ² =0%, p<0.00001, N
	p=0.0006, N =3, 1608 participants	=3, 1630 participants
2h-PPG (mg/dL)	-39.82(-56.70, -22.94), I ² =8%,	-38.74 [-55.81, -21.67], I ² =4%, p<0.00001, N
	p<0.00001, N =5, 311 participants	=4, 228 participants
Total insulin dose	-9.23 (-12.12, -6.33), I ² =39%,	-9.04(-11.48, -6.59), , I ² =0%, p<0.00001, N
(IU/d)	p<0.00001, N =5 comparisons, 1608	=5 comparisons, 1630 participants
(10/4)	participants	
Basal	-8.19 (-10.84, -5.55), I ² =0%, p<0.00001,	-7.76 (-11.23, -4.29), I ² =0%, p<0.00001, N =5
insulin dose (IU/d)	N = 5 comparisons, 1608 participants	comparisons, 1630 participants
Bolus insulin dose	-9.94(-14.84, -5.05), I ² =32%,	-9.77(-14.01, -5.52), I ² =0%, p<0.00001, N =5
(IU/d)	p<0.00001, N =5 comparisons, 1608	comparisons, 1630 participants
(10/4)	participants	
Time-in-	8.88(4.25, 13.5), I ² =0%, p=0.0002, N	11.31(6.75, 15.87), I ² =0%, p<0.00001, N =2,
Range (%)	=4, 278 participants	120 participants
Body weight	-3.66(-4.44, -2.87), I ² =30%, p<0.00001,	-3.50(-3.96, -3.03), I ² =0%, p<0.00001, N =5
change (%)	N = 5 comparisons, 1608 participants	comparisons, 1630 participants
Systolic BP	-3.27 (-4.76, -1.78), I ² =0%, p<0.0001, N	-6.67(-10.38, -2.96), I ² =0%, p=0.0004, N =5
(mmHg)	=5 comparisons, 1608 participants	comparisons, 1630 participants
Diastolic BP	-1.42(-2.20, -0.65), I ² =0%, p=0.0003, N	-1.44(-2.20, -0.69), I ² =0%, p=0.0002, N =5
(mmHg)	=5 comparisons, 1608 participants	comparisons, 1630 participants
eGFR	-1.35 (-2.26, -0.44), I ² =0%, p=0.004,	-1.07 (-2.35, -0.29), I ² =0%, p=0.21, N =5,
(ml/min/1.73 m ²)	N =5, 1608 participants	1630 participants
Albumin-creatinine	-13.92(-27.36, -0.48),	-20.10(-40.25, -0.63), I ² =NA, p=0.04, N =1,
ratio (ACR)(mg/g)	I ² =0%, p=0.04, N=4, 1608 participants	1402 participants
Tauto (ACR)(mg/g)	, , , , , , , , , , , , , , , , , , , ,	
Hypoglycemia	-13.47(-20.90, -6.03), I ² =0%, p=0.004,	-6.12(-10.96, -1.28), I ² =0%, p=0.01 N=5,
(orienta non notice)	N=5, 1608 participants	1630 participants
(events per patient- ar)		
Severe	0.69(0.46, 1.02), I ² =0%, p=0.07, N=5,	$0.71(0.36, 1.43), I^2=0\%, p=0.34, N=5, 1630$
Hypoglycemia	1608 participants	participants

DKA	6.62(2.04, 21.48), I ² =0%, p=0.002, N=5,	2.21(0.43, 11.42), I ² =0%, p=0.34, N =5, 1630
	1608 participants	participants
UTI	0.86(0.48, 1.56), I ² =0%, p=0.62, N =3,	0.96(0.57, 1.59), I ² =0%, p=0.86, N =3, 1630
	1608 participants	participants
GTI	3.39(1.53, 7.52), I ² =14%, p<0.003, N	2.97(1.71, 5.19), I ² =0%, p=0.0001, N =5,
	=5, 1608 participants	1630 participants
Diarrhea	1.50 (0.97, 2.29), I ² =0%, p=0.07, N =5,	0.98 (0.32, 3.01), I ² =0%, p=0.98, N =5, 1630
	1608 participants	participants
Volume depletion	1.89 (0.76, 4.68), I ² =0%, p=0.17, N	2.68 (0.93, 7.73), I ² =0%, p=0.0001, N=5,
Events	=5, 1608 participants	1630 participants
MACE	0.89(0.33, 2.44), I ² =0%, p=0.82, N =5,	5.03(0.24, 104.55), I ² =0%, p=0.30, N =5,
	1608 participants	1630 participants
	Duration of diabe	tes
Outcome	duration of diabetes<20	duration of diabetes≥20
	yr	yr
HbA1c (%)	-0.33(-0.44, -0.22), I ² =0%, p<0.00001,	-0.36(-0.46, -0.25),, I ² =0%, p<0.00001, N =6,
	N =4 comparisons, 902 participants	2336 participants
FPG (mg/dL)	-17.18(-31.70, -2.66), I ² =0%, p=0.01, N	-18.19(-23.76, -12.62), I ² =0%, p<0.00001, N
	=4 comparisons, 902 participants	=6, 2336 participants
2h-PPG (mg/dL)	-51.96(-67.00, -36.92), I ² =0%,	-29.94(-42.98, -16.89), I ² =16%, p<0.00001, N
	p<0.00001, N =4 comparisons, 262	=5, 277 participants
	participants	
Total insulin dose	-7.16(-9.79, -4.53), I ² =0%, p<0.00001,	-9.75(-12.21, -7.28),, I ² =0%, p<0.00001, N
(IU/d)	N =4 comparisons, 902 participants	=6, 2336 participants
Basal	-5.83 (9.47, -2.19), I ² =0%, p=0.002, N	-9.14(-11.72, -6.56),, I ² =0%, p<0.00001, N
insulin dose (IU/d)	=4 comparisons, 902 participants	=6, 2336 participants
Bolus insulin dose	-9.42(-14.79, -4.04), I ² =0%, p=0.0006,	-9.18 (-13.47, -4.90),, I ² =20%, p<0.00001, N
(IU/d)	N =4 comparisons, 902 participants	=6, 2336 participants
Time-in-	11.53(8.21, 14.84), I ² =0%, p<0.00001,	7.69(1.52, 13.89), I ² =0%, p=0.02, N =2, 136
Range (%)	N =4 comparisons, 262 participants	participants
Body weight	-3.13(-3.82, -2.44), I ² =0%, p<0.00001,	-3.13(-3.82, -2.44), I ² =0%, p<0.00001, N =6,
change (%)	N =4 comparisons, 902 participants	2336 participants
Systolic BP (mmHg)	-3.50(-5.72, -1.28), I ² =0%, p=0.0002, N	-4.01(-5.33, -2.70), I ² =13%, p<0.00001, N =6,
	=4 comparisons, 902 participants	2336 participants
Diastolic BP (mmHg)	-1.24(-2.27, -0.21), I ² =0%, p=0.02, N =4	-1.51(-2.14, -0.87), I ² =0%, p<0.00001, N =6,
	comparisons, 902 participants	2336 participants
	<u> </u>	

eGFR	-1.36(-2.47, -0.26), I ² =0%, p=0.02, N =4	-0.66(-1.36, -0.04), I ² =0%, p=0.04, N=6,
(ml/min/1.73 m ²)	comparisons, 902 participants	2336 participants
Albumin-creatinine	-20.45(-33.12, -7.77), I ² =0%, p=0.002,	-15.71(-32.62, 1.21), I ² =0%, p=0.01, N =3,
rotio (ACD)(mg/g)	N = 2 comparisons, 782 participants	1798 participants
ratio (ACR)(mg/g) Hypoglycemia	-13.68(-21.90, -5.46), I ² =0%, p=0.001,	-7.58(-11.24, -1.91), I ² =0%, p=0.006, N =6,
	N =4 comparisons, 902 participants	2336 participants
(events per patient-		
year)		
Severe	0.68(0.36, 1.31), I ² =0%, p=0.25, N =4	$0.70(0.47, 1.05), I^2=0\%, p=0.08, N=6, 2336$
Hypoglycemia	comparisons, 902 participants	participants
DKA	4.60(1.82, 15.73), I ² =0%, p=0.006, N =4	4.30(1.98, 9.31), I ² =0%, p=0.0002, N=6,
	comparisons, 902 participants	2336 participants
UTI	1.13(0.62, 2.07), I ² =0%, p=0.69, N=4	$0.91(0.63, 1.32), I^2=0\%, p=0.73, N=6, 2336$
	comparisons, 902 participants	participants
GTI	3.76(1.73, 8.16), I ² =0%, p=0.0008, N =4	2.95(1.92, 4.52), I ² =0%, p<0.00001, N=6,
	comparisons, 902 participants	2336 participants
Diarrhea	1.85 (0.93, 3.68), I ² =0%, p=0.08, N =4	1.39 (0.92, 2.09), I ² =0%, p=0.12, N =6,
	comparisons, 902 participants	2336 participants
Volume depletion	1.55 (0.63, 3.83), I ² =0%, p=0.34, N =4	2.10 (0.92, 4.85)
events	comparisons, 902 participants	, I ² =0%, p=0.12, N =6, 2336 participants
MACE	2.02(0.34, 12.13), I ² =0%, p0.44, N =4	$0.82(0.17, 3.92), I^2=0\%, p0.80, N=6, 2336$
	comparisons, 902 participants	participants
	Background thera	ру
Outcome	stable insulin therapy	pre-randomization insulin
		optimization
HbA1c (5)	-0.44(-0.52, -0.36), I ² =0%, p<0.00001,	-0.37(-0.45, -0.29), I ² =0%, p<0.00001, N =4,
	N =6, 1663 participants	1575 participants
FPG (mg/dL)	$-20.21(-27.60, -12.83), I^2=0\%,$	-13.46(-20.49, -6.43), I ² =0%, p=0.0002, N =4,
	p<0.00001, N =6, 1663 participants	1575 participants
2h-PPG (mg/dL)	-38.72(-52.27, -25.16), I ² =19%,	-40.10 (-63.73, -16.47), I ² =0%, p=0.0009, N
_	p<0.00001, N =5, 261 participants	=4, 278 participants
Total insulin dose	-9.26(-11.66, -6.87), I ² =0%, p<0.00001,	-8.94(-11.98, -5.89), I ² =0%, p<0.00001, N =4,
(111/4)	N =6, 1663 participants	1575 participants
(IU/d) Basal	-7.38(-10.71, -4.04), I ² =0%, p<0.00001,	-8.47(-11.18, -5.76), I ² =0%, p<0.00001, N =4,
	N =6, 1663 participants	1575 participants
insulin dose (IU/d)	, L L	* *

Bolus insulin	-10.12(-15.07, -5.16), I ² =0%, p<0.0001,	-8.51(-12.57, -4.45), I ² =0%, p<0.0001, N =4,
dose(IU/d)	N =6, 1663 participants	1575 participants
Time-in-	11.31(6.75, 15.87), I ² =0%, p<0.00001,	9.35(5.50, 13.21), I ² =0%, p<0.00001, N =4,
Range (%)	N =6, 120 participants	311 participants
Body weight	-3.48(-3.95, -3.02), I ² =0%, p<0.00001,	-3.70(-4.58, -2.83), I ² =0%, p<0.00001, N =4,
1 (0/)	N =6, 1663 participants	1575 participants
change (%) Systolic BP (mmHg)	-6.67(-10.38, -2.96), I ² =0%, p=0.0004,	-3.27([-4.76, -1.78), I ² =0%, p<0.0001, N =4,
	N =6, 1663 participants	1575 participants
Diastolic BP (mmHg)	-1.43(-2.18, -0.69), I ² =0%, p=0.0002, N	-1.43(-2.22, -0.65), I ² =0%, p=0.0004, N =4,
	=6, 1663 participants	1575 participants
eGFR	-0.98(-1.70, -0.23), I ² =0%, p=0.03, N	-1.37(-2.22, -0.52), I ² =0%, p=0.002, N
(ml/min/1.73 m ²)	=6, 1663 participants	=4, 1575 participants
Albumin-creatinine	-20.10(-39.57, -0.63), I ² =0%, p=0.04, N	-13.92(-27.36, -0.48), I ² =0%, p=0.04, N
ratio (ACR)(mg/g)	=1, 1402 participants	=4, 1575 participants
Hypoglycemia	-7.23(-12.05, -2.40), I ² =0%, p=0.01, N	-13.32(-20.81, -5.83), I ² =0%, p=0.0005, N =4,
(avents now notion)	=6, 1663 participants	1575 participants
(events per patient-		
year)		
Severe	$0.70 (0.37, 1.04), I^2=0\%, p=0.08, N=6,$	0.68(0.46, 1.02), I ² =0%, p=0.06, N =4, 1575
Hypoglycemia	1663 participants	participants
DKA	$3.08(1.32, 7.17), I^2=0\%, p=0.009, N=6,$	6.90(1.91, 24.89), I ² =0%, p=0.003, N =4,
	1663 participants	1575 participants
UTI	0.89 (0.54, 1.45), I ² =0%, p=0.64, N=6,	1.03(0.68, 1.55), I ² =0%, p0.90, N =4, 1575
	1663 participants	participants
GTI	2.64(1.55, 4.49), I ² =0%, p=0.0003, N	3.68(2.17, 6.24), I ² =0%, p<0.00001, N =4,
	=6, 1663 participants	1575 participants
Diarrhea	1.59 (1.03, 2.46), I ² =0%, p=0.04, N	1.51 (1.07, 2.26], I ² =0%, p=0.04, N =4,
	=6, 1663 participants	1575 participants
Volume depletion	2.23 [0.90, 7.44], I ² =0%, p=0.08, N	1.80 (0.70, 4.65), I ² =0%, p=0.22, N =4, 1575
events	=6, 1663 participants	participants
MACE	0.89(0.33, 2.44), I ² =0%, p=0.82, N=6,	1.03 (0.24, 10.55) I ² =0%, p=0.78, N =4, 1575
	1663 participants	participants
	Renal function at bas	eline
Outcome	eGFR≥60 m1/min/1.73 m²	eGFR≥45 ml/min/1.73m ²
HbA1c (%)	-0.39 (-0.63, -0.14), I ² =0%, p=0.0002, N	-0.37 (-0.46, -0.27), I ² =0%, p<0.00001, N=6,
	=4, 174 participants	3064 participants
	, 1	1 1

FPG (mg/dL)	-18.29 (-32.87, -3.71), I ² =28%, p=0.01,	-17.46(-23.00, -11.92), I ² =6%, p<0.00001, N
	N =4, 174 participants	=6, 3064 participants
2h-PPG (mg/dL)	-33.81(-46.92, -20.69), I ² =2%,	-45.63(-63.51, -27.75), I ² =21%, p<0.00001, N
	p<0.00001, N =4, 174 participants	=5, 365 participants
Total insulin dose	-8.46 (-15.13, -1.79), I ² =20%, p=0.01,	-9.03(-11.14, -6.92), I ² =9%, p<0.00001, N =6,
(IU/d)	N =4, 174 participants	3064 participants
Basal	-8.51 [-15.60,- 0.59], I ² =8%, p=0.03,	-8.57 (-10.77, -6.36), I ² =0%, p<0.00001,
insulin dose (IU/d)	N =4, 174 participants	N =6, 3064 participants
Bolus insulin dose	-17.55 (-26.14, -8.96), I ² =0%, p=0.01,	-9.04 (-12.21, -5.86),
(IU/d)	N =4, 174 participants	I ² =6%, p<0.00001, N =4, 3064 participants
Time-in-	11.80 (3.50, 20.10), I ² =NA, p=0.005, N	9.44 (5.88, 12.99), I ² =17%, p<0.00001, N
Range (%)	=1, 33 participants	=5, 365 participants
Body weight	-2.98 (-5.02, -0.95), I ² =0%, p=0.0006,	-3.64 (-4.16, -3.11), I ² =35%, p<0.00001, N
change (%)	N =4, 174 participants	=6, 3064 participants
Systolic BP	-7.93(-13.06, -2.80), I ² =0%, p=0.0002,	-3.71(-4.64, -2.78), I ² =0%, p<0.00001, N
(mmHg)	N =4, 174 participants	=6, 3064 participants
Diastolic BP	-1.53(-2.59, -0.46), I ² =28%, p=0.005,	-1.51(-2.33, -0.70), I ² =0%, p<0.00001, N
(II-)	N =4, 174 participants	=6, 3064 participants
(mmHg) eGFR	-1.21(-3.99, -0.57), I ² =0%, p=0.04, N	-0.78 [-1.42, -0.15], I ² =0%, p=0.02, N =6,
(1/ ² /1 72 ²)	=4, 174 participants	3064 participants
(ml/min/1.73 m ²) Albumin-creatinine	No study	-14.57(-26.87, -2.28), I ² =0%, p=0.02, N =5,
(* (A CP)(,	2977 participants
ratio (ACR)(mg/g) Hypoglycemia	-9.70 [-19.50, -3.11], I ² =0%, p=0.01,	-9.47 (-14.55, -4.38), I ² =0%, p<0.00001, N
	N =4, 174 participants	=6, 3064 participants
(events per patient-	17 – 4, 174 participants	-0, 3004 participants
year)		
Severe	0.49 (0.11, 2.06), I ² =0%, p=0.33, N	0.71 (0.50, 1.01), I ² =0%, p=0.06, N=6,
Hypoglycemia	=4, 174 participants	3064 participants
DKA	8.06(1.04, 22.25), I ² =0%, p=0.04, N	4.72 (1.99, 11.21), I ² =0%, p=0.0002, N =6,
	=4, 174 participants	3064 participants
UTI	0.35 (0.08, 1.59), I ² =0%, p=0.91, N	1.01 (0.73, 1.40), I ² =0%, p=0.76, N=6,
	=4, 174 participants	3064 participants
GTI	2.29 (1.07, 7.71), I ² =0%, p=0.04, N =4,	3.38 (2.30, 4.98), I ² =0%, p<0.00001, N=6,
	174 participants	3064 participants
Diarrhea	1.50 [1.08, 3.10], I ² =0%, p=0.04, N	1.53 (1.09, 2.14), I ² =0%, p=0.03, N=6,
	=4, 174 participants	3064 participants
	1	

Volume depletion	3.85 (0.89, 6.48), I ² =0%, p=0.13, N	2.23 (0.91, 4.60), I ² =0%, p=0.33, N =6,
events	=4, 174 participants	3064 participants
MACE	No events, N =4, 174 participants	1.06 (0.40, 2.82), I ² =0%, p=0.91, N =6,
		3064 participants
Sensitivity	v analysis: Peto Odds Ra	tio, fixed-effect model
Outcome	<u> </u>	significance, N-comparisons,
	011(00)(01), 1, 50001501001	organization, it comparisons,
	parti	icipants
Severe	0.68(0.46, 0.98), I ² =0%, p=0.04,N=10, 32	38 narticipants
Severe	0.08(0.40, 0.38), 1 =0%, p=0.04,N=10, 32	38 participants
Hypoglycemia	2 02 (2 27 C 47) 1 ² 00(0 00001 N 1	0.2020
DKA	3.92 (2.37, 6.47), I ² =0%, p<0.00001, N=1	0, 3238 participants
UTI	0.98(0.71, 1.37), I ² =0%, p=0.92, N=10, 32	238 participants
GTI	2.85(2.10, 3.87), I ² =0%, p<0.00001, N=10	0, 3238 participants
Diarrhea	1.55 (1.11, 2.16), I ² =0%, p=0.01, N=10, 3	238 participants
Nausea-vomiting	0.97(0.32, 2.96), I ² =0%, p=0.96, N=10, 32	238 participants
Headache	1.69(0.26, 11.04), I ² =0%, p=0.58, N=10, 3238 participants	
Sinusitis	1.07(0.06, 18.62), I ² =0%, p=0.91, N=10, 3	3238 participants
Nasopharyngitis	1.07(0.14, 8.39), I ² =0%, p=0.91, N=10, 32	238 participants
Renal events	1.19(0.57, 2.45),I ² =0%, p=0.65, N=10, 32	38 participants
Acidosis-related	3.70 (2.80, 4.90), I ² =0%, p<0.00001, N=	10, 3238 participants
Events		
Volume depletion	2.64 (1.44, 4.83), $I^2=0\%$, $p=0.01$, $N=10$, Ω	3238 participants
events		
Bone fractures	$0.70(0.39, 1.25), I^2=0\%, p=0.23, N=10, 32$	238 participants
Amputation	3.40(0.26, 18.38)I ² =0%, p=0.38, N=10, 3238 participants	
Suspected drug- duced liver injury	1.01(0.09, 11.13), I ² =0%, p=0.99, N=10, 3238 participants	
Serious AEs	1.13(0.86, 1.48), I ² =0%, p=0.39, N=10, 3	238 participants
AEs leading to	1.57 (1.06, 2.34), I ² =0%, p=0.02, N=10, 3	3238 participants
Discontinuation		
MACE	1.15(0.48, 2.80), I ² =0%, p=0.75, N=10, 32	238 participants

Cancer	0.67(0.22, 2.11), I2=0%, p=0.75, N=10, 3238 participants
All-cause deaths	0.19 (0.03, 1.51), I ² =0%, p=0.12, N=10, 3238 participants

Abbreviations: AE: adverse events; FPG: fasting plasma glucose; MACE: major adverse cardiovascular outcomes DKA: diabetic ketoacidosis; GTI: genital tract infections; PPG: postprandial plasma glucose; UTI: urinary tract infections

Supplementary Table 4. Dose-response interactions: within-trial analysis of the pooled data from three RCTs (Baker et al; Buse et al; Danne et al.). Only statistically significant interactions between evaluated outcomes and sotagliflozin doses are reported.

Outcome	Sotagliflozin 200 mg vs. 75 mg	Sotagliflozin 400 mg vs. 200 mg		
HbA1c (%)	$-0.24 (-0.62, 0.14) I^2=NA, p=0.22, N=1,$	-0.22 (-0.28, -0.12)		
	70 participants	$I^{2}=0\%$, p=0.001, N =3, 1119 participants		
FPG (mg/dL)	0.0 (-14.06, 14.06), I ² =NA, p=1.00,	-9.82 (-17.05, -2.58), I ² =0%, p=0.008, N =3,		
	1.0 N =1, 70 participants	1119 participants		
2h-PPG (mg/dL)	-8.00(-27.46, 11.46), I ² =NA, p<0.00001,	-20.51 (-33.98, -7.03), I ² =0%, p=0.003, N =3,		
	N =1, 70 participants	1119 participants		
Total insulin dose	2.60(-6.78, 11.98), I ² =0%, p=0.77, N=1,	-5.25(-7.66, -2.84), I ² =0%, p<0.0001, N =3,		
(%)	70 participants	1119 participants		
Basal insulin	-0.10(-11.11, 10.91), I ² =0%, p=0.99, N	-4.64(-8.64, -0.64), I ² =0%, p=0.01, N =3, 1119		
dose (%)	=1, 70 participants	participants		
Bolus insulin dose (%)	-2.80(-8.48, 14.08), I ² =0%, p=0.89, N =1,	-7.85(-11.96, -3.75), I ² =0%, p=0.0002, N =3		
	70 participants	1119 participants		
Time-in-range(%)	No study	6.48(2.97, 9.99), I ² =0%, p=0.0003, N =2, 185		
		participants		
Average daily	No study	-11.02(-17.70, -4.33), I ² =0%, p=0.001, N =2,		
Glucose (mg/dL)		185 participants		
Urinary glucose	16.00(3.06, 28.94), p=0.03, N=1, 70	13.00(-1.78, 27.78), p=0.20, N=1, 70		
Excretion (g/24 hr)	participants	participants		
Body weight (%)	-1.33(-3.37, 0.71), p=0.20, N =1, 70	-0.96 (-1.55, -0.37), I ² =0%, p=0.001, N =3,		
	participants	1119 participants		
Systolic BP(mmHg)	1.60(-7.42, 10.62), p=0.53, N =1, 70	-2.51 (-3.83, -1.20), I ² =0%, p=0.0002,		
	participants	N =3, 1119 participants		
eGFR	-0.26(-4.95, 4.43), p=0.91, N=1, 70	1.05(0.11, 2.12], p=0.03, N=1, N=3, 1119		
2	participants	participants		
(ml/min/1.73 m ²) Urinary	No study	-12.29 (-26.81, -1.23), I ² =0%, p=0.03, N =3,		
•		1049 participants		
albumin/creatinine		20.5 paracipanto		
ratio (ACR)(mg/g)				

Supplementary Table 5: Summary of main findings of meta-analysis for safety outcomes in included RCTs

Outcome	Studies	Events/Participants (n/N)		Effect estimate	[1 ² (%)
	(n)				
		Sotagliflozin	Control	[95%CI]	
Hypoglicemia (events per patient-year)	6	87/1912	98/1326	MD: -7.69 (-13.25, -2.13)	0
Severe hypoglycemia	6	68/1912	57/1326	RR: 0.69 (0.49, 0.98)	0
Diabetic ketoacidosis (DKA)	6	61/1912	6/1326	RR: 3.93 (1.94, 7.96)	0
Occurring at blood glucose>250 mg/dL n(% total events)		42 (69%)	4 (67%)		
Occurring at blood glucose≥150-250 mg/dL n(% total events)		19(31%)	2(33%)		
Occurring at blood glucose<150-mg/dL n(% total events)		0 (0%)	0 (0%)		
Urinary tract infections (UTIs)	6	96/1912	63/1326	RR: 0.97 (0.71, 1.33)	0
Genital mycotic infections (GTIs)	6	161/1912	31/1326	RR: 3.12 (2.14, 4.54)	0
Diarrhea	6	114/1912	46/1326	RR: 1.50 (1.08, 2.10)	0
Nausea-vomiting	6	8/ 1912	7/1326	RR: 0.60 (0.12, 2.94)	0
Headache	6	3/1912	2/1326	RR: 1.59 (0.30, 8.33]	0
Sinusitis	6	1/1912	1/1326	RR: 1.07 [0.06, 15.62)	0
Nasopharingytis	6	2/1912	2/1326	RR: 1.07 (0.13, 8.67)	0

Renal events	6	21/1912	11/1326	RR: 1.16 (0.56, 2.40)	0
Acidosis-related events	6	187/1912	32/1326	RR: 3.85 (2.33, 6.36)	23
Volume depletion events	6	38/1912	8/1326	RR: 2.19 (1.10, 4.36)	0
Bone fractures	6	29/1912	23/1326	RR: 0.71 (0.40, 1.24)	0
Amputation	6	2/1912	0/1326	RR: 3.02 (0.31, 29.09)	0
Suspected drug-induced liver injury	6	2/1912	1/1326	RR: 0.44 (0.07, 2.76)	0
Venous thromboembolism	6	0/1877	0/1888	-	-
Serious AEs	6	109/1912	143/1326	RR: 1.29 (0.89, 1.82)	0
AEs leading to discontinuation	6	81/1912	31/1326	RR: 1.34 (0.78, 2.30)	25
Hypoglycemia		1 (1%)*	3(3%)*		
Severe hypoglycemia		4(6%)*	3(5%)*		
Diabetic ketoacidosis (DKA)		23(38%)*	1(17%)*		
Urinary tract infections (UTIs)		3(3%)*	4(6%)*		
Genital tract infections (GTIs)		9(6%)*	3(10%)*		
Diarrhea		8(7%)*	3(7%)*		
Volume depletion events		1(4%)*	1(12%)*		
Major adverse cardiovascular outcomes (MACE)	6	15/1912	7/1326	RR: 1.06 (0.40, 2.82)	0
AMI		8	3		
Stroke		1	2		
Hospitalization for HF/UA		0	0		
Coronary revascularization		6	2		
Cancer	6	7/1912	4/1326	RR: 0.86 (0.25, 2.97)	0
Breast		2	2		
Lung		3	2		
Thyroid		1	0		

Melanoma		1	0		
All-cause death	6	1/1912	3/1326	RR: 0.35 (0.07, 1.70)	0

Abbreviations: AE: adverse events; VTE: Venousthromboembolism; Sota: sotagliflozin; TID: total daily insulin dose; plcb: placebo; HF: heart failure; UA: unstable angina.

*the percentage refers to the percentage of all patients experiencing that AE

For all outcomes, the length of follow-up ranged 4 to 52 weeks

REFERENCES

http://www.cdc.gov/diabetes/pubs/statsreport14/diabetesinfographic.pdf; accessed November 30th 2017.

- ² Turner R, Stratton I, Horton V, Manley S, Zimmet P, Mackay IR, et al. UKPDS 25: autoantibodies to islet-cell cytoplasm and glutamic acid decarboxylase for prediction of insulin requirement in type 2 diabetes. UK Prospective Diabetes Study Group. Lancet 1997; 350: 1288-93.
- ³ Miller KM, Foster NC, Beck RW. Current state of Type 1 diabetes treatment in the U.S.: updated data from the T1D Exchange Clinic Registry. Diabetes Care 2015; 38: 971-8.
- ⁴ Lind M, Svensson AM, Kosiborod M, Gudbjörnsdottir S, Pivodic A, Wedel H, et al. Glycemic control and excess mortality in type 1 diabetes. N Engl J Med 2014;371:1972–1982
- ⁵Nathan DM; DCCT/EDIC Research Group. The Diabetes Control and Complications Trial/ Epidemiology of Diabetes Interventions and Complications study at 30 years: overview. Diabetes Care 2014;37:9–16
- ⁶ American Diabetes Association. Standards of Medical Care in Diabetes-2018. Diabetes Care 2018; 41(Suppl. 1): S73-S85.
- ⁷Gerstein HC, Miller ME, Byington RP, Goff DC Jr, Bigger JT, Buse JB, et al. Action to Control Cardiovascular Risk in Diabetes Study Group. Effects of intensive glucose lowering in type 2 diabetes. N Engl J Med 2008; 358: 2545–2559
- ⁸ Zoungas S, Patel A, Chalmers J, de Galan BE, Li Q, Billot L, et al. Severe hypoglycemia and risks of vascular events and death. N Engl J Med. 2010;363:1410-8.
- ⁹ Seaquist ER, Anderson J, Childs B, Cryer P, Dagogo-Jack S, Fish L, et al Hypoglycemia and diabetes: a report of a workgroup of the American Diabetes Association and the Endocrine Society. Diabetes Care. 2013;36:1384-95
- ¹⁰ Cefalu WT, Tamborlane WV, Skyler JS. Type 1 diabetes at a crossroads! Diabetes Care 2015; 38: 968-70.
- ¹¹ Monnier L, Colette C, Owens D. The glycemic triumvirate and diabetic complications: is the whole greater than the sum of its component parts. Diabetes Res Clin Pract 2012;95:303–11.
- ¹² Ceriello A, Esposito K, Piconi L, Ihnat MA, Thorpe JE, Testa R, et al. Oscillating glucose is more deleterious to endothelial function and oxidative stress than mean glucose in normal and type 2 diabetic patients. Diabetes 2008:57:1349–54.
- ¹³ Soupal J, Skrha J, Jr., Fajmon M, Horová E, Mráz M, Škrha J, et al.: Glycemic variability is higher in type 1 diabetes patients with microvascular complications irrespective of glycemic control. Diabetes Technol Ther 2014;16: 198–203.

¹US Center for Disease Control and Prevention;

¹⁴ Snell-Bergeon JK, Roman R, Rodbard D, Garg S, Maahs DM, Schauer IE, et al.: Glycaemic variability is associated with coronary artery calcium in men with Type 1 diabetes: the Coronary Artery Calcification in Type 1 Diabetes study. Diabet Med 2010; 27:1436–1442.

- ¹⁵ Zinman B, Marso SP, Poulter NR, Emerson SS, Pieber TR, Pratley RE, et al. Day-to-day fasting glycaemic variability in DEVOTE: associations with severe hypoglycaemia and cardiovascular outcomes (DEVOTE 2). Diabetologia 2018;61:48–57.
- ¹⁶ Pieber TR, Marso SP, McGuire DK, Zinman B, Poulter NR, Emerson SS, et al. DEVOTE 3: temporal relationships between severe hypoglycaemia, cardiovascular outcomes and mortality. Diabetologia 2018;61:58–65.
- ¹⁷ Wright LA, Hirsch IB. Metrics Beyond Hemoglobin A1C in Diabetes Management: Time in Range, Hypoglycemia, and Other Parameters. Diabetes Technol Ther. 2017;19:S16-S26
- ¹⁸ Danne T, Nimri R, Battelino T, Bergenstal RM, Close KL, DeVries JH, et al. International Consensus on Use of Continuous Glucose Monitoring. Diabetes Care. 2017;40:1631-1640
- ¹⁹American Diabetes Association. Standards of Medical Care in Diabetes-2018. Diabetes Care 2018; 41(Suppl. 1): S73-S85.
- ²⁰ American Diabetes Association. 6. Glycemic targets: *Standards of Medical Care in Diabetes*—2018. Diabetes Care 2018;41(Suppl. 1):S55–S64
- ²¹ Seidelmann SB, Feofanova E, Yu B, Franceschini N, Claggett B, Kuokkanen M, et al. Genetic Variants in SGLT1, Glucose Tolerance, and Cardiometabolic Risk. J Am Coll Cardiol. 2018;72:1763-1773.
- ²² Yamada T, Shojima N, Noma H. Sodium-glucose co-transporter-2 inhibitors as add-on therapy to insulin for type 1 diabetes mellitus: Systematic review and meta-analysis of randomized controlled trials. Diabetes Obes Metab. 2018;20:1755-1761.
- ²³ Powell DR, DaCosta CM, Smith M: Effect of LX4211 on glucose homeostasis and body composition in preclinical models. J Pharmacol Exp Ther 2014;350:232–242.
- ²⁴ Powell DR, Smith M, Greer J, Harris A, Zhao S, DaCosta C, et al. LX4211 increases serum glucagonlike peptide 1 and peptide YY levels by reducing sodium/glucose cotransporter 1 (SGLT1)-mediated absorption of intestinal glucose. J Pharmacol Exp Ther. 2013;345:250-9
- ²⁵ Meek TH, Dorfman MD, Matsen ME, Fischer JD, Cubelo A, Kumar MR, et al.: Evidence that in uncontrolled diabetes, hyperglucagonemia is required for ketosis but not for increased hepatic glucose production or hyperglycemia. Diabetes 2015;64:2376–2387.
- ²⁶ Sands AT, Zambrowicz BP, Rosenstock J, Lapuerta P, Bode BW, Garg SK² et al. Sotagliflozin, a Dual SGLT1 and SGLT2 Inhibitor, as Adjunct Therapy to Insulin in Type 1 Diabetes. Diabetes Care. 2015;38:1181-8.

²⁷. Bode B, Phillip Banks Paul Strumph Sangeeta Sawhney The Sotagliflozin JDRF Study Writing Group Efficacy and safety of sotagliflozin, a dual SGLT1 and SGLT2 inhibitor, as adjunct to insulin in young adults with poorly controlled type 1 diabetes (JDRF Study) Diabetologia 2017; 60: S87-88

²⁸ Baker C, Wason S, Banks P. A 12-week dose-ranging study of sotagliflozin, a dual SGLT1 and SGLT2 inhibitor, as adjunct therapy to insulin in type 1 diabetes (inTandem4) Diabetologia 2017; 60: S409

²⁹ Garg SK, Henry RR, Banks P, Buse JB, Davies MJ, Fulcher GR, et al. Effects of Sotagliflozin Added to Insulin in Patients with Type 1 Diabetes. N Engl J Med. 2017;377:2337-2348

³⁰Buse JB, Garg SK, Rosenstock J, et al. Sotagliflozin in Combination With Optimized Insulin Therapy in Adults With Type 1 Diabetes: The North American inTandem1 Study. Diabetes Care. 2018 Jun 24. pii: dc180343. doi: 10.2337/dc18-0343. [Epub ahead of print]

³¹Danne T, Cariou B, Banks P, et al. HbA_{1c} and Hypoglycemia Reductions at 24 and 52 Weeks With Sotagliflozin in Combination With Insulin in Adults With Type 1 Diabetes: The European inTandem2 Study. Diabetes Care. 2018 Jun 24. pii: dc180342. doi: 10.2337/dc18-0342. [Epub ahead of print]

³² https://search.usa.gov/search?utf8=%E2%9C%93&affiliate=fda&query=sotagliflozin&commit=Search

³³ https://www.ema.europa.eu/en/search/search?search api views fulltext=sotagliflozin

³⁴ https://ss.pmda.go.jp/en_all/search.x?q=sotagliflozin&ie=UTF-8&page=1&x=30&y=11

³⁵ www.lexpharma.com

³⁶ www.sanofi.com

³⁷ Esposito K, Giugliano D, Nappo F, Marfella R. Campanian Postprandial Hyperglycemia Study Group. Regression of carotid atherosclerosis by control of postprandial hyperglycemia in type 2 diabetes mellitus. Circulation. 2004;110:214-9

³⁸ Raz I, Ceriello A, Wilson PW, Battioui C, Su EW, Kerr L, et al.: Post hoc subgroup analysis of the HEART2D trial demonstrates lower cardiovascular risk in older patients targeting postprandial versus fasting/premeal glycemia. Diabetes Care 2011;34: 1511–1513.

³⁹https://www.fda.gov/NewsEvents/Newsroom/PressAnnouncements/ucm611454.htm

⁴⁰ Baghurst PA. Calculating the mean amplitude of glycemic excursion from continuous glucose monitoring data: an automated algorithm. Diabetes Technol Ther 2011;13:296–302

⁴¹ https://www.meddra.org/ Last accessed on July 21st, 2018

⁴²Higgins JPT, Green S, eds. Cochrane Handbook for Systematic Reviews of Interventions, Version 5.2.0. The Cochrane Collaboration; updated March 2011. Accessed at www.cochrane-handbook.org on 14 July 2017.

- ⁴⁶ Safer DJ. Design and reporting modifications in industry sponsored comparative psychopharmacology trials. J Nerv Ment Dis 2002;190:583-92.
- ⁴⁷National Research Council. Panel on Handling Missing Data in Clinical Trials. Committee on National Statistics, Division of Behavioral and Social Sciences and Education. Washington, DC: The National Academies Press; 2010. The prevention and treatment of missing data in clinical trials.
- ⁴⁸ Psaty BM, Kronmal RA. Reporting mortality findings in trials of rofecoxib for Alzheimer disease or cognitive impairment: a case study based on documents from rofecoxib litigation. JAMA. 2008; 299: 1813-7.
- ⁴⁹ Henry DA, Kerridge IH, Hill SR, McNeill PM, Doran E, Newby DA, et al. Medical specialists and pharmaceutical industry-sponsored research: A survey of the Australian experience. MJA. 2005;182:557-60.
- ⁵⁰ Gøtzsche PC, Hr´objartsson A, Johansen HK, Haahr MT, Altman DG, Chan A-W. Constraints on publication rights in ndustry-initiated clinical trials. JAMA. 2006;295:1645-6.
- ⁵¹ Review Manager (RevMan) [computer program]. Version 5.3. Copenhagen, Denmark: Nordic Cochrane Center, Cochrane Collaboration; 2012.
- ⁵²Liberati A, Altman DG, Tetzlaff J, Mulrow C, Gøtzsche PC, et al. The PRISMA Statement for Reporting Systematic Reviews and Meta-Analyses of Studies That Evaluate Health Care Interventions: Explanation and Elaboration. BMJ 2009;339:b2700.
- ⁵³ Guyatt GH, Oxman AD, Kunz R. GRADE Working Group. GRADE guidelines: 7. Rating the quality of evidence—inconsistency. J Clin Epidemiol. 2011;64:1294-302.
- ⁵⁴ Bradburn MJ, Deeks JJ, Berlin JA, Russell Localio A. Much ado about nothing: a comparison of the performance of meta-analytical methods with rare events. Stat Med. 2007;26: 53-77.
- ⁵⁵ Fisher DJ, Carpenter JR, Morris TP, Freeman SC, Tierney JF. Meta-analytical methods to identify who benefits most from treatments: daft, deluded, or deft approach? BMJ. 2017; 356:j573.

⁴³Viswanathan M, Patnode CD, Berkman ND, Bass EB, Chang S, Hartling L, et al. Recommendations for assessing the risk of bias in systematic reviews of health-care interventions. J Clin Epidemiol. 2018;97:26-34.

⁴⁴Lachin JM. Fallacies of last observation carried forward analyses. Clin Trials. 2016;13:161-8.

⁴⁵ Lexchin J.Sponsorship bias in clinical research. Int J Risk Saf Med. 2012;24: 233-42

⁵⁶ Guyatt G, Oxman AD, Akl EA. GRADE guidelines: 1. Introduction-GRADE evidence profiles and summary of findings tables. J Clin Epidemiol. 2011;64:383-94.

⁵⁷Guyatt GH, Oxman AD, Kunz R, Brozek J, Alonso-Coello P, Rind D, et al. GRADE guidelines 6. Rating the quality of evidence--imprecision. J Clin Epidemiol. 2011 Dec;64(12):1283-93.

⁵⁸https://www.easd.org/virtualmeeting/home.html#!resources/efficacy-and-safety-of-sotagliflozin-a-dual-sglt1-and-sglt2-inhibitor-as-adjunct-to-insulin-in-young-adults-with-poorly-controlled-type-1-diabetes-jdrf-study-15ae18d4-b8f6-439c-87ea-2f19dcad388e

⁵⁹https://www.easd.org/virtualmeeting/home.html#!resources/a-12-week-dose-ranging-study-of

sotagliflozin-a-dual-sglt1-and-sglt2-inhibitor-as-adjunct-therapy-to-insulin-in-type-1-diabetes

intandem4Dose-ranging Study in Patients With Type 1 Diabetes Mellitus (inTandem4)

- ⁶⁰ Lee AK, Warren B, Lee CJ. The association of severe hypoglycemia with incident cardiovascular events and mortality in adults with type 2 diabetes. Diabetes Care 2018;41:104–11.
- ⁶¹ NovoNordisk Company Announcement 51/2015; https://www.novonordisk.com/bin/getPDF.1947182.pdf; Accessed on: 24 October 2018.
- ⁶²Monnier L, Colette C, Wojtusciszyn A, et al. Toward defining the threshold between low and high glucose variability in diabetes. Diabetes Care 2017;40:832–8.
- ⁶³ Fanelli CG, Epifano L, Rambotti AM, Pampanelli S, Di Vincenzo A, Modarelli F, et al. Meticulous prevention of hypoglycemia normalizes the glycemic thresholds and magnitude of most of neuroendocrine responses to, symptoms of, and cognitive function during hypoglycemia in intensively treated patients with short-term IDDM. Diabetes. 1993;42:1683-9.
- ⁶⁴ Neuen BL, Ohkuma T, Neal B, Matthews DR, de Zeeuw D, Mahaffey KW, et al. Cardiovascular and Renal Outcomes With Canagliflozin According to Baseline Kidney Function. Circulation. 2018:138:1537-1550.
- ⁶⁵ Heerspink HJ, Perkins BA, Fitchett DH, Husain M, Cherney DZ. Sodium glucose cotransporter 2 inhibitors in the treatment of diabetes mellitus: cardiovascular and kidney effects, potential mechanisms, and clinical applications. *Circulation*. 2016;134:752–772.

⁶⁶ Musso G, Gambino R, Cassader M, Pagano GF. A novel approach to control hyperglycemia in type 2 diabetes: sodium glucose co-transport (SGLT) inhibitors: systematic review and meta-analysis of randomized trials. Ann Med. 2012;44:375-93.

⁶⁷ Janssen Research & Development, LLC. Canagliflozin as an adjunctive treatment to diet and exercise alone or co-administered with other antihyperglycemic agents to improve glycemic control in adults with type 2 diabetes mellitus, 11 December 2012, http://www.fda.gov/downloads/

AdvisoryCommittees/CommitteesMeetingMaterials/Drugs/EndocrinologicandMetabolicDrugsAdvisoryCommittee/ UCM334551.pdf (accessed 13 September 2014).

⁶⁸ Neumiller JJ. Empagliflozin: a new sodium-glucose cotransporter 2 (SGLT2) inhibitor for the treatment of type 2 diabetes. *Drugs Context* 2014; 3: 212–262

⁶⁹Rosenstock J, Ferrannini E. Euglycemic Diabetic Ketoacidosis: A Predictable, Detectable, and Preventable Safety Concern With SGLT2 Inhibitors. Diabetes Care. 2015;38:1638-42.

 70 Henry RR, Dandona P, Pettus J, Mudaliar S, Xu J, Hansen L.. Dapagliflozin in patients with type 1 diabetes: A post hoc analysis of the effect of insulin dose adjustments on 24-hour continuously monitored mean glucose and fasting β-hydroxybutyrate levels in a phase IIa pilot study. Diabetes Obes Metab. 2017:19:814-821.

⁷¹ Olsen SE1, Asvold BO, Frier BM, Aune SE, Hansen LI, Bjørgaas MR Hypoglycaemia symptoms and impaired awareness of hypoglycaemia in adults with Type 1 diabetes: the association with diabetes duration. Diabet Med. 2014;31:1210-7.

⁷² Peters AL, Buschur EO, Buse JB. Euglycemic Diabetic Ketoacidosis: A Potential Complication of Treatment With Sodium-Glucose Cotransporter 2 Inhibition. Diabetes Care. 2015; 38:1687-93.

Dandona P, Mathieu C, Phillip M, D, Hansen L, Griffen SC, Tschöpe D, et al. Efficacy and safety of dapagliflozin in patients with inadequately controlled type 1 diabetes (DEPICT-1): 24 week results from a multicentre, double-blind, phase 3, randomised controlled trial. Lancet Diabetes Endocrinol. 2017;5: :864-876.

⁷⁴Workgroup on Hypoglycemia, American Diabetes Association.Defining and reporting hypoglycemia in diabetes: a report from the American Diabetes Association Workgroup on Hypoglycemia.

Diabetes Care. 2005;28:1245-9

⁷⁵Kitabchi AE, Umpierrez GE, Miles JM, Fisher JN. Hyperglycemiccrises in adultpatients with diabetes.Diabetes Care. 2009;32:1335-43