Efficacy and Safety of Switching From the DPP-4 Inhibitor Sitagliptin to the Human GLP-1 Analog Liraglutide After 52 Weeks in Metformin-Treated Patients With Type 2 Diabetes

A randomized, open-label trial

OBJECTIVE—To assess the efficacy and safety of switching from sitagliptin to liraglutide in metformin-treated adults with type 2 diabetes.

RESEARCH DESIGN AND METHODS—In an open-label trial, participants randomized to receive either liraglutide (1.2 or 1.8 mg/day) or sitagliptin (100 mg/day), each added to metformin, continued treatment for 52 weeks. In a 26-week extension, sitagliptin-treated participants were randomly allocated to receive instead liraglutide (pooled 1.2 and 1.8 mg/day, 1.3; 0.0189). After switching, mostly transient nausea occurred in 21% of participants, and minor hypoglycemia remained low (3; 0.0001). Conversion to liraglutide was associated with reductions in fasting plasma glucose (FPG) (1.2 mg/day, −0.8 mmol/L, P = 0.0004; 1.8 mg/day, −1.4 mmol/L, P < 0.0001) and body weight (1.2 mg/day, −1.6 kg; 1.8 mg/day, −2.5 kg; both P < 0.0001) and with an increased proportion of patients reaching HbA1c <7% (from ~30% to ~50%). Overall treatment satisfaction, assessed by the Diabetes Treatment Satisfaction Questionnaire, improved after switching to liraglutide (pooled 1.2 and 1.8 mg/day, 1.3; P = 0.0189). After switching, mostly transient nausea occurred in 21% of participants, and minor hypoglycemia remained low (3–4% of participants). Continuing liraglutide treatment at 1.2 mg/day and 1.8 mg/day for 78 weeks reduced HbA1c (baseline 8.3 and 8.4%, respectively) by −0.9% (from baseline, additional decreases occurred after switching to liraglutide (1.2 mg/day, −0.2%, P = 0.006; 1.8 mg/day, −0.5%, P = 0.0001). Conversion to liraglutide was associated with reductions in fasting plasma glucose (FPG) (1.2 mg/day, −0.8 mmol/L, P = 0.0004; 1.8 mg/day, −1.4 mmol/L, P < 0.0001) and body weight (1.2 mg/day, −1.6 kg; 1.8 mg/day, −2.5 kg; both P < 0.0001) and with an increased proportion of patients reaching HbA1c <7% (from ~30% to ~50%). Overall treatment satisfaction, assessed by the Diabetes Treatment Satisfaction Questionnaire, improved after switching to liraglutide (pooled 1.2 and 1.8 mg/day, 1.3; P = 0.0189). After switching, mostly transient nausea occurred in 21% of participants, and minor hypoglycemia remained low (3–4% of participants). Continuing liraglutide treatment at 1.2 mg/day and 1.8 mg/day for 78 weeks reduced HbA1c (baseline 8.3 and 8.4%, respectively) by −0.9% and −1.3%, respectively; FPG by −1.3 and −1.7 mmol/L, respectively; and weight by −2.6 and −3.1 kg, respectively, with 9–10% of participants reporting minor hypoglycemia.

CONCLUSIONS—Glycemic control, weight, and treatment satisfaction improved after switching from sitagliptin to liraglutide, although with a transient increase in gastrointestinal reactions.

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Although glucagon-like peptide 1 (GLP-1) receptor agonists (GLP-1RAs) and dipeptidyl peptidase-4 (DPP-4) inhibitors both have a glucose-dependent mechanism of action, distinct differences have emerged in phase 3 trials that, for the most part, have lasted as long as 6 months. In independent trials in patients with type 2 diabetes, the GLP-1RA liraglutide produced reductions in glycated hemoglobin HbA1c as great as 1.6%, in body weight of ~3 kg, and in systolic blood pressure of 2–7 mmHg; it also improved β-cell function (1,2). Two other GLP-1RA regimens, exenatide twice daily and exenatide once weekly, decreased HbA1c by 0.8–0.9% and 1.3–1.9%, respectively, and produced weight reductions similar to liraglutide (up to 3 kg) (3–8). In contrast, smaller reductions in HbA1c (0.4–1.0%) and negligible weight changes have been reported with currently available DPP-4 inhibitors (9–17).

The greater efficacy of GLP-1RAs is probably related to the pharmacological levels of these agonists stimulating GLP-1 receptor activity (18). In contrast, DPP-4 inhibitors modestly affect the levels of endogenous GLP-1, thus producing smaller glycemic reductions and negligible weight effects (19).

Although longer-term, head-to-head trials with the two incretin classes are scarce, we recently reported the results of a 52-week, head-to-head comparison of liraglutide and sitagliptin added to metformin in participants with type 2 diabetes (20). The results showed that liraglutide (1.2 or 1.8 mg/day) produced significantly greater sustained decreases than sitagliptin 100 mg/day in HbA1c (−1.3 and −1.5%, respectively, vs. −0.9%), fasting plasma glucose (FPG) (−1.7 and −2.0 mmol/L, respectively, vs. −0.6 mmol/L), and body weight (−2.8 and −3.7 kg, respectively, vs. −1.2 kg),
with a comparable rate of hypoglycemia although an initially higher frequency of nausea in the liraglutide groups (20). Furthermore, a significantly greater percentage of patients achieved the American Diabetes Association target of HbA1c <7.0% with liraglutide (1.2 and 1.8 mg) than with sitagliptin (30.3 and 63.3% for liraglutide 1.2 and 1.8 mg/day, respectively, vs. 27.1% for sitagliptin) (20). Apart from the higher incidence of gastrointestinal events with liraglutide, as expected with GLP-1RAs, the overall frequencies of adverse events (AEs), serious AEs (SAEs), and minor hypoglycemia were generally comparable between liraglutide and sitagliptin groups (20,21). Significantly greater reductions in HbA1c, FPG, and body weight relative to sitagliptin were also reported with the GLP-1RA exenatide once weekly in a 26-week, head-to-head trial (HbA1c, −0.9% vs. −1.5%; FPG, −0.9 vs. −1.8 mmol/L; body weight, −0.8 vs. −2.3 kg) (6).

Although metformin is the established first-line therapy in type 2 diabetes, there is no consensus regarding optimal second-line therapy or an effective alternative agent when the first second-line therapy used fails to provide adequate glycemic control. Although head-to-head studies have reported that treatment with GLP-1RAs produce greater glycemic and weight benefits compared with DPP-4 inhibitors (6,20,21), there are very limited data available exploring the effects of switching from a DPP-4 inhibitor to a GLP-1RA (22). To address this important clinical question, we conducted an exploratory investigation during which we assessed the efficacy and safety of switching participants from sitagliptin to liraglutide after 52 weeks in a 26-week study extension. The efficacy and safety of continuing liraglutide therapy for 78 weeks were also examined.

**RESEARCH DESIGN AND METHODS**

**Study design and outcome measures**

The 52-week study design and patient inclusion and exclusion criteria were reported previously (21). Briefly, individuals with type 2 diabetes, inadequately controlled with metformin (HbA1c ≥1500 mg/dL after ≥3 months (HbA1c, 7.5–10%)), were randomly allocated to receive either injectable liraglutide 1.2 or 1.8 mg/day or oral sitagliptin 100 mg/day, each added to metformin. After 52 weeks, sitagliptin-treated participants were again randomly allocated (1:1) to receive either liraglutide 1.2 or 1.8 mg/day, through a weekly dose escalation of 0.6 mg/day, and were treated for another 26 weeks. The participants who were randomly assigned to liraglutide at baseline continued the same treatment for the entire 78 weeks (Fig. 1).

Withdrawal criteria were almost identical to those at 52 weeks (20), with one additional withdrawal criterion (FPG >10.0 mmol/L, with no treatable current cause) applicable during the extension. Efficacy and safety end points were identical to those evaluated at 52 weeks and were assessed as reported previously (20). The protocol, including the extension, was approved by the appropriate institutional review board. The study followed Good Clinical Practice guidelines and conformed to the Declaration of Helsinki, with participants providing written, informed consent. The 78-week trial started on 6 June 2008 and ended on 3 June 2010.

**Statistical methods**

For participants receiving liraglutide from baseline to week 78, efficacy was analyzed with the full analysis set, all randomly allocated participants who were exposed to at least one dose of the trial drug, which was identical to the safety analysis set in this trial. For participants switching from sitagliptin to liraglutide (weeks 52–78), efficacy and safety were analyzed with the extension analysis set, all full analysis set 52-week completers exposed to the trial products during the extension.

As described previously (23), the validated Diabetes Treatment Satisfaction Questionnaire (DTSQ), status version, was used to assess participant satisfaction with treatment (24,25). As in the main trial period, participants from Slovakia, Serbia, and Slovenia were also excluded from the DTSQ analyses during the extension (82 of 419 [19.6%]) because of the lack of linguistically validated questionnaires. The two liraglutide switch groups (those switching from sitagliptin to either 1.2 or 1.8 mg/day liraglutide) were pooled into one group during a post hoc analysis of DTSQ scores. This was done to increase the power of the analysis and to allow a clearer interpretation of score changes between weeks 52 and 78 as differences between oral and injectable treatments. Missing postbaseline data were imputed by means of the last observation carried forward method.

Changes from week 52 to 78 within groups were analyzed by paired t test. Logistic regression was used to analyze the participant proportions achieving HbA1c targets and composite end point (HbA1c <7.0%, with no weight gain and no confirmed major or minor hypoglycemia), with treatment and country as fixed effects and baseline HbA1c, and body weight (for composite) as covariates. The proportions of participants achieving glycemic and composite targets at weeks 52 and 78 were compared with the McNemar test for matched pairs.

Changes from baseline to week 78 were analyzed with ANCOVA, with treatment and country as fixed effects and baseline value as a covariate. Hypoglycemia and serum calcitonin were analyzed as reported previously (21). Mean ± SD values are presented for baseline and week 52 data to demonstrate parameter variability within the population, whereas changes during weeks 52–78 and 0–78 are reported as mean ± SE to illustrate the accuracy of the estimates. P < 0.05 was considered significant.

**RESULTS**—The groups were well matched for demographic and other characteristics at baseline (21). Of 436 total participants completing 52 weeks of treatment, 419 (96.1%) entered the extension, with 381 of the 419 participants (90.9%) completing 78 weeks in total (Fig. 1). Of the 284 combined liraglutide continuers (1.2 and 1.8 mg/day) entering the extension, 259 (91%) finished 78 weeks of treatment. The most common withdrawal reasons during the extension from the liraglutide continuing groups (1.2 and 1.8 mg/day) were meeting the withdrawal criteria (1.8 and 1.4%, respectively) and ineffective therapy (1.3 and 1.4%, respectively). Of the 135 participants who were switched to liraglutide after week 52, 122 (90%) completed the extension. The most common withdrawal reasons from the switch groups (sitagliptin to liraglutide 1.2 mg and sitagliptin to liraglutide 1.8 mg/day) were AEs (9% and 0%, respectively) and meeting the withdrawal criteria (1.5 and 4.4%, respectively). No participants withdrew from either switch group because of ineffective therapy during the extension.

**Participants switching from sitagliptin to liraglutide**

**Efficacy.** Although 52 weeks of treatment with sitagliptin changed baseline HbA1c by −0.9%, switching to liraglutide (1.2 or 1.8 mg/day) for the next 26 weeks was associated with additional changes...
from the 52-week HbA1c values (Fig. 2A): −0.2% ± 0.1% from 7.23% ± 0.9% (P = 0.006) for liraglutide 1.2 mg/day and −0.5% ± 0.1% from 7.6% ± 1.2% for liraglutide 1.8 mg/day (P = 0.0001). FPG also decreased in both switch groups during weeks 52–78 (Fig. 2B): from 8.6 ± 1.7 mmol/L (154.8 ± 30.6 mg/dL) by −0.8 ± 0.2 mmol/L (−14.4 ± 3.6 mg/dL) for liraglutide 1.2 mg (P = 0.0004) and from 9.2 ± 2.1 mmol/L (165.6 ± 37.8 mg/dL) by −1.4 ± 0.3 mmol/L (−25.2 ± 5.4 mg/dL) for liraglutide 1.8 mg/day (P < 0.0001).

Significant weight reductions occurred after switching to liraglutide for 26 weeks (Fig. 2C): from 92.8 ± 20.6 kg by −1.6 ± 0.4 kg for liraglutide 1.2 mg/day and from 91.6 ± 18.7 kg by −2.5 ± 0.4 kg for liraglutide 1.8 mg/day (both P < 0.0001). Consistent with this observation, significant decreases in waist circumference also occurred after 26 weeks of switching to either dose of liraglutide (Supplementary Table 1).

The proportion of participants achieving HbA1c <7.0% (American Diabetes Association target) increased significantly after switching to liraglutide for both groups (P = 0.0005 for 1.2 mg/day; P = 0.0026 for 1.8 mg/day); (Fig. 2D). A greater percentage of participants reached the American Association of Clinical Endocrinologists target of HbA1c ≤6.5% after switching to liraglutide, with a significant increase observed only for the liraglutide 1.8 mg/day group (P = 0.0117) (Fig. 2D).

Furthermore, changing therapy to liraglutide (both groups) was associated with a significant increase in the percentage of participants reaching the composite end point of HbA1c <7.0%, with no weight gain and no confirmed major or minor hypoglycemia (P = 0.0018 for liraglutide 1.2 mg/day; P = 0.0192 for liraglutide 1.8 mg/day) (Fig. 2E).

Treatment satisfaction was assessed as reported previously in the pooled participant population switched to liraglutide (Fig. 2F) (23). From weeks 52 to 78, overall treatment satisfaction improved significantly for patients switching to liraglutide (+1.3; P = 0.0189), driven mainly by improvements in the categories "current..."
Figure 2—Selected efficacy and safety parameters. Mean HbA1c (A), FPG (B), and change in body weight (C) during 78 weeks for participants originally randomly allocated to receive liraglutide and participants switched to liraglutide after 52 weeks (Wk). D: Proportions of switch group participants (%) reaching target HbA1c <7.0% or ≤6.5% at weeks 52 and 78. E: Proportions of switch group participants reaching composite end point of HbA1c <7.0%, with no weight gain and no confirmed major or minor hypoglycemia, at weeks 52 and 78. F: Changes in the DTSQ scores from Switching from sitagliptin to liraglutide.
Switching to liraglutide significantly increased β-cell function, as determined by homeostasis model assessment of β-cell function (both groups), and significantly reduced the homeostasis model assessment of insulin resistance (sitagliptin to liraglutide 1.8 mg/day) and in LDL cholesterol (both groups) (Supplementary Table 1). There were no significant changes in blood pressure in either switch group during weeks 52–78, but the elevation was statistically significant only for the sitagliptin to liraglutide 1.8 mg/day group (Supplementary Table 1).

Switching to liraglutide was associated with significant reductions in fasting triglycerides and total cholesterol (sitagliptin to liraglutide 1.8 mg/day) and in LDL cholesterol (both groups) (Supplementary Table 1). Mean heart rate increased slightly in both switch groups during weeks 52–78, but the increase was statistically significant only for the sitagliptin to liraglutide 1.8 mg/day group (Supplementary Table 1).

There were no significant changes in waist circumference in participants originally randomly allocated to receive liraglutide during weeks 0–78, both in the 1.8 mg/day group. One previously reported death from pancreatic carcinoma (diagnosed after 8 days of liraglutide treatment) occurred during the first 52 weeks and was classified by the investigator as unlikely to be related to the trial drug. The second death (bile duct cancer) occurred during the extension in a participant treated for 401 days (diagnosed after 316 days) and with a known choleodochal stenosis in the medical history and cholecystectomy with stent implant since 2006. The event was classified by the investigator as unlikely to be related to the trial drug.

During the 78 weeks, two events of major hypoglycemia occurred in a 39-year-old female participant originally randomly allocated to receive liraglutide 1.2 mg/day. The first episode occurred during the main trial period and was
Switching from sitagliptin to liraglutide

described previously (21). The second episode occurred during the extension, after 486 treatment days. Finger-stick glucose concentration was too low to register. The participant was hospitalized overnight for glucose stabilization; no seizures or loss of consciousness occurred. The participant recovered after 1 day, and the loss of consciousness occurred. The participant was admitted for glucose stabilization; no seizures or concentration was too low to register. The episode occurred during the extension, after the participant switched to liraglutide for 1.2 mg/day. The rates of minor hypoglycemia, serious and possibly related to the trial drug. The rates of minor hypoglycemia, after exclusion of an extreme outlier in the liraglutide 1.8 mg/day group with 23 minor episodes, were 0.156 and 0.130 episodes/participant/year for liraglutide 1.2 and 1.8 mg/day, respectively.

During the 78 weeks, mean serum calcitonin levels remained below the upper limit of normal for both sexes in the two treatment groups, and no cases of malignant thyroid neoplasm were reported. One case of pancreatitis (nonacute) occurred during the first 52 weeks and was described previously; no additional cases occurred after week 52 (20).

CONCLUSIONS—Switching from sitagliptin after 52 weeks to liraglutide for the next 26 weeks, both in combination with metformin, was associated with significant improvements in glycemic control, body weight, β-cell function, and overall treatment satisfaction. Apart from transient increases in gastrointestinal reactions, as is common when initiating GLP-1RA therapy, no additional or unexpected safety or tolerability concerns were raised by switching to either dose of liraglutide. Participants treated with liraglutide for 78 weeks had clinically significant reductions in HbA1c (0.9–1.3%) and weight (2.5–3.1 kg), with safety and tolerability profiles consistent with previous reports and no new safety concerns (1).

These results are consistent with other long-term studies of liraglutide. In one trial, 2 years of treatment with liraglutide 1.2 or 1.8 mg/day changed HbA1c by −0.6 and −0.9%, respectively, and body weight by −2.3 and −2.8 kg, respectively (26). In another, 26-week improvements in HbA1c were sustained for 2 years of treatment with liraglutide, and weight reductions were also maintained after 2.5 years (−3.0 kg with 1.2 mg/day and −2.9 kg with 1.8 mg/day) (27).

The greater efficacy of liraglutide than sitagliptin observed in the first 52 weeks of this study is consistent with other head-to-head trials comparing a GLP-1RA with sitagliptin (the DURATION-2 [Duration therapy Utilization: Researching changes in A1c, weight, and other factors Through Intervention with exenatide ONce weekly] trial of exenatide once weekly vs. sitagliptin and the T-emerge-4 trial of taspoglutide vs. sitagliptin) (6,28). The DURATION-2 trial also included a switch phase after 26 weeks. The 26-week reductions in HbA1c (−0.3%) and body weight (−1.1 kg) after switching from sitagliptin to exenatide once weekly in the DURATION-2 trial extension were similar to the reductions observed in participants switched to liraglutide 1.2 mg/day in our study; however, greater mean decreases in these glycemic control indicators and weight occurred in patients switched to liraglutide 1.8 mg/day (Fig. 2A–C) (22).

These glycemic and nonglycemic improvements after switching therapy are likely due to the different modes of action exhibited by DPP-4 inhibitor sitagliptin and the GLP-1RA liraglutide. Direct stimulation of GLP-1 receptors by GLP-1RAs dosed to pharmacological levels results in glucose-dependent insulin secretion, inhibition of glucagon release, increased satiety, and reduced food intake, thus translating to effective glycemic control and weight loss observed in multiple studies (18,19). In contrast, DPP-4 inhibitors act indirectly by preventing enzymatic degradation of the incretin hormones, GLP-1 and glucose-dependent insulinotropic polypeptide (18,19). Although DPP-4 inhibitors also stimulate glucose-dependent insulin secretion and inhibit glucagon release, lower levels of endogenous GLP-1 achievable with these agents translate into reduced glycemic efficacy and negligible weight effects relative to GLP-1RAs.

The assessment of treatment satisfaction can provide valuable information on patient adherence and thus may help predict long-term treatment outcomes (29). Our results show that the switch from an oral therapy to an injectable was associated with an increase in overall treatment satisfaction, while patients’ perceptions of treatment convenience and flexibility remained essentially unchanged. The improved treatment satisfaction may result from weight loss or actual or perceived improvements in treatment efficacy. These results complement the greater treatment satisfaction with liraglutide compared with sitagliptin reported in the first 52 weeks of this trial (23), and similar results have been reported in another study comparing sitagliptin with an injectable GLP-1RA (30).

The 78-week safety profile of liraglutide was generally consistent with that observed during the development program, with gastrointestinal disorders being the most prevalent AEs (1). The liraglutide continuing groups had more gastrointestinal events than the sitagliptin group during the initial 4–8 weeks of the 78 treatment weeks. Similarly, gastrointestinal events were the most frequently reported events after switching to liraglutide, as expected with GLP-1RAs (1,2). In accordance with previous reports, nausea incidence peaked transiently after switching and declined after several weeks (1,21).

Postmarketing reports of pancreatitis with exenatide, and later sitagliptin, resulted in the addition of pancreatitis as a withdrawal criterion in this trial as well, and a single case of chronic pancreatitis was reported during the first year (20). In addition, during the trial, calcitonin levels remained below the upper normal limits for both sexes with both liraglutide and sitagliptin, consistent with findings from all phase 3 trials with liraglutide (31).

Overall study design limitations included the absence of a placebo group and the lack of double blinding. Limitations that applied specifically to the switch period were the small number of patients per switch group and the lack of a control group (e.g., a sitagliptin continuing group) to serve as a reference and comparison for the observed efficacy changes in the groups switched to liraglutide. Because the number of patients was small, it was decided not to keep a sitagliptin continuing group and instead switch all sitagliptin recipients to liraglutide; however, another 2-year study with sitagliptin added to metformin has shown that efficacy tends to decline after the first year (32), and thus no further improvement in sitagliptin-treated patients would have been expected had they not switched to liraglutide.

In conclusion, switching treatment from sitagliptin to liraglutide was associated with improvements in glycemic control, body weight, and treatment satisfaction, while the rates of hypoglycemia remained low. Overall, the switch was well tolerated, although it was accompanied by a transient rise in gastrointestinal reactions (mainly nausea), as expected with GLP-1 receptor agonists. The additional glycemic and weight reductions observed after switching second-line treatment to liraglutide show that this switch may be especially beneficial for those patients not achieving glycemic goals with sitagliptin, without
the need for a third-line antihyperglycemic agent. It is reasonable to hypothesize that similar findings may be observed when comparing other long-acting GLP-1RAs and DPP-4 inhibitors. This exploratory study sets the stage for larger, more robust studies to examine the effects of switching from one second-line type 2 diabetes therapy to another.

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R.P. participated in study design, enrolled patients, reviewed and interpreted the clinical trial report and data, and reviewed and edited the manuscript. M.A.N., T.B., E.M., S.Fi., A.J.G. and M.D. were investigators, and they participated in the interpretation of data, as well as in review of the manuscript. A.B.T. was International Medical Director for the trial, and she participated in trial conduct, planning of analyses, data interpretation, and drafting and revision of the manuscript. S.Fu. participated in data interpretation and review of the manuscript. R.P. is the guarantor of this work and, as such, had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.


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