Efficacy and Patient Satisfaction of Dipeptidyl Peptidase-4 Inhibitor After Switching From Once-Daily DPP-4 Inhibitor to Once-Weekly Regimen

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Abstract

Background: We administered once-weekly dipeptidyl peptidase-4 (DPP-4) inhibitor (W) (used omarigliptin as W in this study) to patients with type 2 diabetes taking once-daily DPP-4 inhibitor (D), and investigated efficacy, safety and patient satisfaction before and after switching to W.

Methods: W was administered to 182 patients with type 2 diabetes taking D (used sitagliptin as D in this study), who had been visiting our hospital on an outpatient basis; 164 (90.6\%) of these patients requested to switch medications. Of these 164 patients, this study investigated 153 who requested to continue taking W. Hemoglobin A1c (HbA1c) levels, body weight, blood pressure and a questionnaire survey (Diabetes Treatment Satisfaction Questionnaire (DTSQ)) were evaluated in these patients.

Results: Patient characteristics were as follows: age, 63.9 ± 10.3; male/female ratio, 93:60; duration of diabetes, 14.9 ± 7.7 years; and body mass index (BMI), 25.5 ± 4.2 kg/m\textsuperscript{2}. After switching to W, HbA1c levels changed from 7.41 ± 0.7\% to 7.36 ± 0.9\%, which was not statistically significant. Changes in body weight, BMI, and systolic and diastolic blood pressure were also not significant. On the DTSQ, satisfaction of Q1 significantly increased (P < 0.01). The score for lifestyle assessment did not significantly change, but compliance significantly improved (P < 0.001).

Conclusion: This study revealed that 90\% of patients taking D elected to switch to W. Moreover, patient satisfaction and compliance improved after switching to W. Increased satisfaction appeared to be influenced by improved blood glucose control, but was not associated with compliance. Switching from D to W did not affect HbA1c levels but improved patient adherence.

Keywords: Adherence; Satisfaction; Compliance; Weekly DPP-4 inhibitor; Omarigliptin; Type 2 diabetes

Introduction

Remarkable progress has been made in medications for treating diabetes in parallel with a global increase in the number of diabetic patients. In the realm of diabetes medications, therapeutic drugs with novel mechanisms of action are continually being developed, although existing drugs with prolonged actions are also rapidly appearing on the market. The addition of such therapeutic drugs to the repertoire of treatment options is anticipated to be beneficial for improving patient adherence.

The first dipeptidyl peptidase-4 (DPP-4) inhibitor was approved in Japan in 2009. DPP-4 inhibitors suppress postprandial hyperglycemia, are less likely to induce weight gain or hypoglycemia, and involve relatively few comorbidity-related restrictions. Therefore, they are frequently used in Japan [1]. In 2015, once-weekly DPP-4 inhibitors (W), trelagliptin and omarigliptin, were approved in Japan before global approval. In clinical trials during drug development, trelagliptin was compared to alogliptin and omarigliptin was compared to sitagliptin. Non-inferiority in efficacy and safety was demonstrated for both drugs compared to the existing daily medications [2, 3]. It was also recently reported in a meta-analysis that the efficacy and safety of once-weekly medication are comparable to that of existing daily medications [4]. Because once-weekly DPP-4 inhibitors are only commercially available in Japan, their efficacy and safety in general practice have not been established.

The reduced number of doses and tablets in once-weekly DPP-4 inhibitor (W) therapy may ameliorate patient treatment satisfaction and medication adherence, resulting in better long-term blood glucose control. However, no clear superiority in drug efficacy has been demonstrated compared to the existing and familiar once-daily DPP-4 inhibitor (D), and patient interest and assessment of W have been unclear. In this study, we proposed W to patients with type 2 diabetes taking D (used sitagliptin as D in this study), and investigated the changes in serum levels of hemoglobin A1c (HbA1c) before and after switching to W (used omarigliptin as W in this study). We also conducted a questionnaire survey (Diabetes Treatment...
Satisfaction Questionnaire (DTSQ)) to assess satisfaction and compliance.

**Materials and Methods**

Patients were enrolled between June and August 2017. W was proposed to 182 patients with type 2 diabetes taking D, who had been visiting our hospital on an outpatient basis and had no changes in their diabetes treatment (e.g., diet, exercise, medication) within the past 12 weeks. Clinical examination was performed as usual at the visit. After being notified of their blood test results (e.g., HbA1c levels) of that day, all patients were given the following information about the once-weekly medication: “There is a drug that has essentially the same therapeutic effects as those of the DPP-4 inhibitor (D) you are currently taking. This drug only needs to be taken once a week. There is very little difference in price. We can switch or continue the current medication. If the new medication does not seem suitable for you, we will switch back to the original drug (D). What would you like to do?”

We evaluated HbA1c levels, body weight, body mass index (BMI), systolic blood pressure, diastolic blood pressure and patient survey responses before and 8 weeks after switching the patients to W. Three evaluation methods were used in the patient surveys. 1) DTSQ translated by Ishii et al [5] was used to evaluate patient satisfaction. The questionnaire consists of eight questions quantified on a seven-point scale of 0 - 6 and is widely used to evaluate treatment satisfaction in diabetic patients. 2) The evaluations of lifestyle were made according to the following checklist (Table 1), which was modified for this study [6]. The sum of the points represented the lifestyle assessment score. 3) Compliance was evaluated on a visual analog scale (VAS) as shown in Figure 1.

This study was conducted in accordance with the Ethic-
cal Guideline for Clinical Research of the Ministry of Health, Labour and Welfare after obtaining written informed consent from the patients. We obtained approval from the Institutional Review Board of Saiseikai Niigata Daini Hospital (Approval No. E16-38).

All tests were performed using Microsoft Excel 2016 for Mac, and all data are expressed as the mean ± standard deviations. Student’s *t*-test was used for the comparisons of values between before and 8 weeks after switching to W. P values less than 0.05 were considered statistically significant.

**Results**

Patient characteristics are shown in Figure 2. A total of 164 (90.6%) patients requested to switch medications from D to W, after being provided information about W. Of these 164 patients, this study investigated 153 who requested to continue taking W (11 patients requested to switch back to D during the study). Patient characteristics were as follows: age, 63.9 ± 10.3; man/woman ratio, 93:60; duration of diabetes, 14.9 ± 7.7 years; BMI, 25.5 ± 4.2 kg/m²; estimated glomerular filtration rate, 70.2 ± 22.6 mL/min/1.73 m²; HbA1c, 7.4 ± 0.7%; systolic blood pressure, 122.0 ± 11.3 mmHg; diastolic blood pressure, 72.5 ± 7.7 mm Hg; and number of different medications, 5.1 ± 2.2 (all insulin formulations were considered to be one type) (Table 2).

After switching to W, HbA1c levels changed from 7.41 ± 0.7% to 7.36 ± 0.9%, which was not statistically significant.

Changes in body weight, BMI, and systolic and diastolic blood
The questionnaire response rate was 100% because all participants were patients who had agreed to complete the questionnaire survey. For DTSQ, the satisfaction of Q1 significantly increased (P < 0.01; Fig. 3). Recommendation (Q7) tended to increase, but it was not significant. Significant changes in responses were not observed for any of the other questions (Qs).

The association between the number of different medications before switching and the extent of change in satisfaction (Q1) was analyzed, but a correlation was not found (data not shown). Subsequently, patients were divided into two groups: those with increased satisfaction and those with decreased satisfaction on Q1, and the clinical parameters were compared between the two groups (Table 4). In comparing the increased satisfaction (n = 46) and decreased satisfaction groups (n = 24), the former group had significantly improved HbA1c levels compared to the latter group (-0.154 ± 0.32%, 0.146 ± 0.49%; P < 0.01). The score for lifestyle assessment did not significantly change (Fig. 4a), but compliance significantly improved (P < 0.001; Fig. 4b). Compliance and changes in HbA1c were not correlated with each other (data not shown). Switching from D to W did not result in serious adverse events, indicating satisfactory tolerability.

Data are presented as mean ± SD. BMI: body mass index; SBP: systolic blood pressure; DBP: diastolic blood pressure; HbA1c: glycated hemoglobin; DBP: diastolic blood pressure; eGFR: estimated glomerular filtration rate. The insulin preparation counted it for one kind in all.

### Table 2. Patient Characteristics

| Factors               | Mean ± SD (range) |
|-----------------------|-------------------|
| Number of patients (n)| 153               |
| Age (years old)       | 63.9 ± 10.3 (28 - 87) |
| Sex (male/female)     | 94:59             |
| Duration of diabetes (years) | 14.9 ± 7.7 (1 - 43) |
| BMI (kg/m²)           | 25.5 ± 4.2 (16.1 - 39.7) |
| Body weight (kg)      | 65.2 ± 9.5 (49.2 - 84.3) |
| HbA1c (%)             | 7.41 ± 0.75 (5.7 - 9.6) |
| SBP (mm Hg)           | 122.1 ± 11.3 (94 - 143) |
| DBP (mm Hg)           | 72.5 ± 7.8 (59 - 88) |
| eGFR (mL/min/1.73 m²) | 70.23 ± 22.9 (40.9 - 121.2) |
| Number of drugs (n)   | 5.1 ± 2.2 (1 - 12) |
| U-AER (mg/gCr)        | 94.4 ± 187.9 (2.3 - 873.2) |

Data are presented as mean ± SD. AER: albumin excretion rate; BMI: body mass index; SBP: systolic blood pressure; DBP: diastolic blood pressure; eGFR: estimated glomerular filtration rate. The insulin preparation counted it for one kind in all.

### Table 3. Changes in Clinical Parameters After Switching From Sitagliptin to Omarigliptin

|                          | Baseline       | 8 weeks       | P value |
|--------------------------|----------------|---------------|---------|
| BMI (kg/m²)              | 25.5 ± 4.2     | 25.3 ± 3.6    | NS      |
| Body weight (kg)         | 65.2 ± 9.5     | 65.7 ± 9.6    | NS      |
| HbA1c (%)                | 7.41 ± 0.75    | 7.36 ± 0.79   | NS      |
| SBP (mm Hg)              | 122.1 ± 11.3   | 127.6 ± 11.9  | NS      |
| DBP (mm Hg)              | 72.5 ± 7.8     | 75.5 ± 9.7    | NS      |

Data are presented as mean ± SD. BMI: body mass index; SBP: systolic blood pressure; DBP: diastolic blood pressure; NS: not significant by Student’s paired t-test.
In this study, we proposed switching from D to W; accordingly, 90.6% of patients requested to switch to W while 10.4% did not. A previous survey in Japanese type 2 diabetic patients regularly visiting the outpatient clinic reported that 55.3% of patients requested a switch to W [7]. The request rate may have been lower than that in our study because the survey was conducted during a 2-week limited prescription period (In Japan, only 2 weeks’ worth of new drug can be prescribed the first year after its approval). Okazaki et al [8] recently reported that 81% of the 68 Japanese type 2 diabetic patients taking D requested to switch to W, and that the elevation in HbA1c levels before switching led patients to request the switch. In other words, patients with worse blood glucose levels requested switching to W. In this study, changes in HbA1c levels were not evaluated before the medication switch. However, our study took place between June and August when there is seasonal variation (HbA1c levels are highest in winter-spring and lowest in summer-autumn in most patients in Japan) [9]. Considering this, it is unlikely that HbA1c was exacerbated in many patients in this study. Thus, although most patients requested switching to W, it is unknown if the reason was due to increased HbA1c levels.

Table 4. Comparison of Clinical Parameters Between Patients With Increased and With Decreased Satisfaction Based on Q1 Questions

|                        | Satisfied (+) N = 46 | Satisfied (-) N = 24 | P value |
|------------------------|----------------------|----------------------|---------|
| Sex (male/female)      | 27:19                | 14:10                |         |
| Age (years old)        | 63.6 ± 12.8          | 63.4 ± 9.9           | NS      |
| Duration of diabetes (year) | 15.8 ± 7.9         | 14.0 ± 7.7           | NS      |
| eGFR                   | 82.94 ± 24.8         | 77.25 ± 15.9         | NS      |
| U-AER                  | 73.5 ± 167.6         | 42.0 ± 65.7          | NS      |
| BMI (kg/m²), 0 weeks   | 25.7 ± 4.3           | 24.9 ± 4.0           | NS      |
| BMI (kg/m²), 8 weeks   | 25.9 ± 4.3           | 24.6 ± 4.2           | NS      |
| Body weight (kg), 0 weeks | 68.8 ± 13.0         | 66.4 ± 12.7          | NS      |
| Body weight (kg), 8 weeks | 68.8 ± 13.0         | 66.2 ± 13.0          | NS      |
| SBP (mm Hg), 0 weeks   | 121.3 ± 12.5         | 124.9 ± 12.0         | NS      |
| SBP (mm Hg), 8 weeks   | 122.5 ± 15.1         | 126.2 ± 12.7         | NS      |
| DBP (mm Hg), 0 weeks   | 72.1 ± 12.2          | 74.4 ± 10.6          | NS      |
| DBP (mm Hg), 8 weeks   | 72.1 ± 11.1          | 74.9 ± 13.1          | NS      |
| HbA1 (%) , 0 weeks     | 7.57 ± 0.77          | 7.32 ± 0.54          | NS      |
| HbA1c (%) , 8 weeks    | 7.41 ± 0.73          | 7.47 ± 0.67          | NS      |
| ΔHbA1c                 | -0.154 ± 0.32        | 0.146 ± 0.49         | <0.01   |
| Number of drugs        | 5.3 ± 2.0            | 6.0 ± 2.9            | NS      |

Data are presented as mean ± SD. BMI: body mass index; SBP: systolic blood pressure; DBP: diastolic blood pressure; NS: not significant by Student’s paired t-test.

Discussion

In this study, we proposed switching from D to W; accordingly, 90.6% of patients requested to switch to W while 10.4% did not. A previous survey in Japanese type 2 diabetic patients regularly visiting the outpatient clinic reported that 55.3% of patients requested a switch to W [7]. The request rate may have been lower than that in our study because the survey was conducted during a 2-week limited prescription period (In Japan, only 2 weeks’ worth of new drug can be prescribed the first year after its approval). Okazaki et al [8] recently reported that 81% of the 68 Japanese type 2 diabetic patients taking D requested to switch to W, and that the elevation in HbA1c levels before switching led patients to request the switch. In other words, patients with worse blood glucose levels requested switching to W. In this study, changes in HbA1c levels were not evaluated before the medication switch. However, our study took place between June and August when there is seasonal variation (HbA1c levels are highest in winter-spring and lowest in summer-autumn in most patients in Japan) [9]. Considering this, it is unlikely that HbA1c was exacerbated in many patients in this study. Thus, although most patients requested switching to W, it is unknown if the reason was due to increased HbA1c levels.

In this study, significant changes in HbA1c, body weight and BMI were not observed 8 weeks after switching from D to W. A meta-analysis of omarigliptin (five studies) and trelagliptin (two studies) [4] (follow-up of 12 weeks (n = 2) or 24 weeks (n = 5)) also showed similar results. W was originally developed to have the same therapeutic effects as D. All previous reports involved clinical trials, which took place in a specialized environment where participants strictly followed the medication schedule; thus, the effects in patients were
postulated to be equivalent. In actual clinical practice, on the other hand, it has been reported that drug adherence rates are less than 70% in diabetic patients [10]. Thus, a decrease in HbA1c level was anticipated through improved adherence and decreased incidence of forgetting to take the medication by switching to W; however, the results were unchanged. There are two plausible reasons for this. First, the effects of W on adherence may not be substantial enough to improve blood glucose. Second, participants of this study already had good compliance, so switching to W may not have induced significant changes in HbA1c.

Patient adherence to W was evaluated with DTSQ. A significant difference was only observed for Q1 regarding satisfaction. Tosaki et al [11] investigated 26 type 2 diabetic patients and found significant differences in convenience (Q4) and flexibility (Q5) when patients switched from various types of D to W. In their study, D included twice-daily DPP-4 inhibitors. With the exception of the effect of switching from twice-daily to once-weekly medications, their study may not have reflected the true advantage of W. In our study, all patients (n = 153) switched from D (once-daily) to W (once-weekly), reflecting the effects of W in a simple manner. This may have led to divergent results between different studies.

Patients with increased satisfaction after switching from D to W exhibited improved HbA1c levels. It is possible that patient satisfaction increased because the post-switch survey was conducted after informing them of the HbA1c levels measured that day, and patients with improved blood glucose levels may have been in a better mood. It is unclear if this was an effect of W itself.

Because type 2 diabetic patients often present with symptoms such as hypertension and dyslipidemia, the number of medications inevitably increase. We previously investigated drug adherence in patients who switched from taking two drugs to one combination tablet [12]. In the question, “Was your mental burden alleviated with one fewer medications?” approximately 80% of patients responded that the burden “greatly decreased” or “somewhat decreased.” Before conducting this survey, healthcare providers did not anticipate any changes by reducing the number of tablets taken by patients by just one; however, the results indicated that patients were much more satisfied, even with this small decrease. Participants of this study took an average of 5.1 types of medications. Many patients who switched to W commented that, “this is so much easier!” By switching from D to W, patients took one fewer tablet for 6 days of the week, which may have alleviated their mental burden and contributed to their increased satisfaction.

W did not affect the score for lifestyle assessment. It has been reported that this score decreased in patients who showed attenuated therapeutic effects of D during the treatment course [13]. In other words, when there is a disturbance in a patient’s lifestyle, the therapeutic effect of D declines. In this study, we determined whether W could improve lifestyle; however, it did not appear to have enough power to change this parameter.

Although medication adherence significantly improved with W, HbA1c levels did not improve. Moreover, medication adherence was not correlated with satisfaction. Because medication adherence was already high (≥ 80%) when patients were taking D, even if adherence further improved by switching to W, it is unlikely that the changes would be substantial enough to affect HbA1c or satisfaction.

Both D and W are DPP-4 inhibitors with the same mechanisms of action, and as such, are viewed as the same type of drug by healthcare providers. However, they may be completely different drugs (once-daily drug versus once-weekly drug) from the patient’s perspective. The results of this study showed that patient satisfaction and compliance improved with W compared to D. Therefore, it is important to present options to patients before prescribing medications.

Conclusions

This study revealed that 90% of patients taking D elected to switch to W. Moreover, patient satisfaction and compliance improved after switching to W. Increased satisfaction appeared to be influenced by improved blood glucose levels but was not associated with compliance. Switching from D to W did not affect HbA1c levels but improved patient adherence.

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Conflicts of Interest

The authors have no conflicts of interest to declare.

Disclosure

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