Retrospective Analysis on the Efficacy, Safety and Treatment Failure Group of Sitagliptin for Mean 10-Month Duration

Won Jun Kim, Cheol-Young Park, Eun Haeng Jeong, Jeong Youn Seo, Ji Soo Seol, Se Eun Park, Eun Jung Rhee, Won Young Lee, Ki Won Oh, Sung Woo Park, Sun Woo Kim
Department of Endocrinology and Metabolism, Kangbuk Samsung Hospital, Sungkyunkwan University School of Medicine, Seoul, Korea

Background: To investigate the clinical results of sitagliptin (SITA) and the characteristics of the treatment failure group or of low responders to SITA.

Methods: A retrospective study of type 2 diabetic patients reviewed 99 cases, including 12 treatment failure cases, who stopped SITA because of worsening patients' condition, and 87 cases, who continued treatment over five visits (total 9.9±10.1 months) after receiving the prescription of SITA from December 2008 to June 2009. Subjects were classified as five groups administered SITA as an initial combination with metformin (MET), add-on to metformin or sulfonylurea, and switching from sulfonylurea or thiazolidinedione. The changes in HbA1c level from the first to last visit (∆HbA1c) in treatment maintenance group were subanalyzed.

Results: The HbA1c level was significantly reduced in four groups, including initial coadministration of SITA with metformin (∆HbA1c=-1.1%, P<0.001), add-on to MET (∆HbA1c=-0.6%, P=0.017), add-on to sulfonylurea (∆HbA1c=-0.5%, P<0.001), and switching from thiazolidinedione (∆HbA1c=-0.3%, P=0.013). SITA was noninferior to sulfonlyurea (∆HbA1c=-0.2%, P=0.63). There was no significant adverse effect. The treatment failure group had a longer diabetes duration (P=0.008), higher HbA1c (P=0.001) and fasting plasma glucose (P=0.003) compared to the maintenance group. Subanalysis on the tertiles of ∆HbA1c showed that low-response to SITA (tertile 1) was associated with a longer diabetes duration (P=0.009) and lower HbA1c (P<0.001).

Conclusion: SITA was effective and safe for use in Korean type 2 diabetic patients. However, its clinical responses and long-term benefit–harm profile is yet to be established.

Keywords: Diabetes mellitus, type 2; Sitagliptin; Treatment outcome

INTRODUCTION

The prevalence of type 2 diabetes, which is caused by mechanisms such as insulin secretory defect failure and insulin resistance, is rapidly increasing worldwide. To prevent complications in the early stages of type 2 diabetes occupying 90% to 95% of South Korean diabetic patients [1], a thorough blood glucose management should be established [2]. According to the American Diabetes Association (ADA) guidelines and the European Association of the Study of Diabetes, initial therapy should involve lifestyle changes and metformin (MET) treatment. However, because of the progressive nature of the disease, combination therapy consisting of two or more drugs or insulin therapy is typically necessary over time [3].

Corresponding author: Cheol-Young Park
Department of Endocrinology and Metabolism, Kangbuk Samsung Hospital, Sungkyunkwan University School of Medicine, 108 Pyeong-dong, Jongno-gu, Seoul 110-746, Korea
E-mail: cydoctor@chol.com
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The main incretin hormones glucagon-like peptide-1 (GLP-1) and glucose-dependent insulinotropic polypeptide (GIP) stimulate insulin synthesis and secretion in the β-cells of the pancreas when blood glucose levels are high. The use of a glucagon suppressing hormone from α-cells [4] has resulted in significant improvements of differentiation and proliferation of β-cells and inhibition of apoptosis in animal model and human islets [5,6]. In type 2 diabetes patients, GLP-1 secretion and GIP function decrease, however, these incretin hormones can be increased by inhibition of incretin-disabling enzyme dipeptidyl peptidase IV (DPP-IV) [7-9].

Sitagliptin (SITA), an oral DPP-IV inhibitor has been proven valid and low risk of side effects, such as hypoglycemia and weight gain in the treatment of type 2 diabetes. Dhillon [10] has reviewed that the addition of SITA to ongoing treatment therapies resulted in significant improvements to HbA1c. Moreover, the report showed that SITA was noninferior to glipizide and generally did not differ from rosiglitazone. SITA can improved HbA1c without weight gain and rarely results in hypoglycemia except for coadministration with sulfonylurea (SU) or insulin.

SITA was first used as a DPP-IV inhibitor in South Korea in early 2008, and the prevalence of SITA prescription in South Korea is increasing. However, results from Korean subjects are limited and so more data and research are required. The present study was performed in tertiary hospitals, and we investigated the clinical characteristics of type 2 diabetic patients who received SITA.

METHODS

Setting and patients
A retrospective study was conducted for patients starting SITA therapy for type 2 diabetes between December 2008 and June 2009, who attended the outpatient clinic at intervals of two to three months for a total of five visits.

The treatment maintenance group was defined as patients who continued a daily dose of 100 mg SITA during follow-up periods, and the treatment failure group was defined as patients who had discontinued SITA treatment because of increasing HbA1c level above 7% and worsening patients’ condition. Among a total of 99 patients, 87 belonged to the treatment maintenance group and 12 were included in the treatment failure group. Exclusion criteria was switch from more than two different kinds of diabetes medications, voluntary discontinuation of SITA treatment, and follow-up loss. The cases of SITA switched from or added-on to previous medications such as meglitinide, α-glucosidase inhibitor or insulin were rare, so they were also excluded.

Patients in the treatment maintenance group were subdivided into five groups: an initial coadministration of SITA with MET group, add-on to MET or SU groups, switch from SU or thiazolidinedione (TZD) groups.

Additionally, for the 87 patients in the treatment maintenance group the changes in HbA1c level from the first to last visit (HbA1c at visit 1–HbA1c at visit 5, ΔHbA1c) were used to divide into three subgroups. The first tertile group had a low response, and the third tertile group had a good response, so they were comparatively analyzed. SITA prescription beyond coverage of medical insurance was paid privately after the consent of the patients.

Explanatory variables and outcome measures
Fasting plasma glucose levels (FPG) and HbA1c were measured in the outpatient clinic. Blood glucose was measured using the glucose oxidase method (CX3 DELTA; Beckman, Brea, CA, USA); HbA1c was measured using high-performance liquid chromatography (Bio-Rad, Richmond, CA, USA); and insulin and C-peptide levels were measured using a gamma counter (Dream Gamma-10; ShinJin Medics Co., Seoul, Korea). Body mass index (BMI) was surveyed within the first year of SITA administration, and outpatient records or prescriptions including personal history, drugs taken, and side effects were examined.

Statistical analysis
All measurements were expressed as mean±standard deviation (SD) or as median (25% confidence interval [CI], 75% CI). For discontinuous variables, a chi-square analysis was used. For continuous variables, ΔHbA1c and the changes in the FPG (ΔFPG) during follow-up periods were examined using the Wilcoxon Signed Rank test because the Kolmogorov-Smirnov test did not demonstrate normality; the Mann-Whitney test was used for the comparative analysis between the treatment failure and maintenance group. For subanalysis, the Kruskal-Wallis test was used to compare tertiles of ΔHbA1c in the treatment maintenance group. The SPSS for Windows, version 17.0, was used for statistical analysis (SPSS Inc., Chicago, IL, USA), and P values less than 0.05 were considered to be statistically significant.
RESULTS

Baseline characteristics
During the study period, 160 patients were newly prescribed with SITA. Over five visits to the outpatient clinic, 87 patients maintained the type and amount of antidiabetic medication and their mean follow-up period was 9.9±10.1 months. We discontinued administration of SITA in the 12 members in the treatment failure group with deteriorating blood glucose. Of the 99 total participants, the mean age was 53.9±12.0, women were 63 (64%), and the duration of diabetes was 4.5±4.9 years.

The initial coadministration of SITA and MET group included 21 patients, the add-on SITA to MET group had 18 patients, the add-on SITA to SU group had 12 patients, the group switched from SU had 22 patients, and the group switched from TZD had 14 patients. MET was prescribed to 86% to 100% of patients in each group. The age, diabetes duration, HbA1c, FPG, fasting insulin level, fasting C-peptide level, and BMI were expressed as mean±SD or median (25% CI, 75% CI) and gender, history of taking antihypertensive and antihyperlipidemic medications were examined as number and percentage (Table 1).

ΔHbA1c and ΔFPG before and after SITA administration
The groups that had statistically significant reductions in HbA1c level between visit 1 and visit 5 (ΔHbA1c) included the initial coadministration of SITA and MET group (ΔHbA1c=-1.1%, P<0.001), the add-on SITA to SU group (ΔHbA1c=-0.5%, P<0.001), the add-on SITA to MET group (ΔHbA1c=-0.6%, P=0.017), and the group switched from TZD (ΔHbA1c=-0.3%, P=0.013). While HbA1c levels decreased significantly at visit 2, those from visit 2 through visit 5 were not significantly different. The mean doses of the group switched from SU were 1.8 mg in the case of glimepiride.

Table 1. Baseline characteristics of subgroups classified by medication regimen

<table>
<thead>
<tr>
<th>Characteristic (at visit 1)</th>
<th>SITA+METa (n=21)</th>
<th>Adding on MET (n=18)</th>
<th>Adding on SU (n=12)</th>
<th>Switching from SU (n=22)</th>
<th>Switching from TZD (n=14)</th>
<th>Total (n=87)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, yr</td>
<td>53.4±9.7</td>
<td>49.8±10.7</td>
<td>55.8±10.4</td>
<td>58.8±13.5</td>
<td>49.8±10.7</td>
<td>54.1±11.5</td>
</tr>
<tr>
<td>Duration of T2DM, yr</td>
<td>1.6±2.0</td>
<td>3.9±2.8</td>
<td>7.9±4.5</td>
<td>3.2±2.2</td>
<td>3.7±2.1</td>
<td>3.7±3.2</td>
</tr>
<tr>
<td>Females, n (%)</td>
<td>13 (62)</td>
<td>15 (83)</td>
<td>9 (75)</td>
<td>10 (46)</td>
<td>9 (64)</td>
<td>56 (64.4)</td>
</tr>
<tr>
<td>BMI, kg/m²</td>
<td>26.6±3.4</td>
<td>24.3±2.5</td>
<td>24.0±4.2</td>
<td>26.0±3.5</td>
<td>27.4±3.3</td>
<td>25.8±3.4</td>
</tr>
<tr>
<td>HbA1c, %</td>
<td>7.3±1.1</td>
<td>7.1±0.8</td>
<td>7.7±0.6</td>
<td>6.8±0.6</td>
<td>7.3±1.1</td>
<td>7.2±0.9</td>
</tr>
<tr>
<td>FPG, mg/dL</td>
<td>158.5±24.3</td>
<td>136.7±25.4</td>
<td>152.1±21.1</td>
<td>127.9±21.8</td>
<td>152.2±31.2</td>
<td>144.3±27.3</td>
</tr>
<tr>
<td>Anti-hypertensive agent, n (%)</td>
<td>9 (42.9)</td>
<td>6 (33.3)</td>
<td>9 (75.0)</td>
<td>10 (45.5)</td>
<td>7 (50.0)</td>
<td>41 (47.0)</td>
</tr>
<tr>
<td>Anti-lipid agent, n (%)</td>
<td>7 (33)</td>
<td>11 (61.1)</td>
<td>9 (75.0)</td>
<td>13 (59.1)</td>
<td>11 (78.6)</td>
<td>55 (63.2)</td>
</tr>
<tr>
<td>MET agent, n (%)</td>
<td>21 (100)</td>
<td>18 (100)</td>
<td>11 (92)</td>
<td>20 (91)</td>
<td>13 (86)</td>
<td>82 (94.3)</td>
</tr>
<tr>
<td>Fasting insulin, μIU/mL</td>
<td>13.3±6.0</td>
<td>10.4±3.5</td>
<td>8.5±5.0</td>
<td>11.9±3.9</td>
<td>11.3±2.2</td>
<td>11.8±4.6</td>
</tr>
<tr>
<td>Fasting C-peptide, ng/mL</td>
<td>4.0±1.6</td>
<td>2.4±2.5</td>
<td>2.1±1.0</td>
<td>3.2±0.8</td>
<td>3.1±0.8</td>
<td>3.3±1.3</td>
</tr>
</tbody>
</table>

Data are presented as mean±standard deviation and median (25% confidence interval [CI], 75% CI) except female, using anti-hypertensive agent and anti-lipid agent and metformin agent (n [%]). SITA, sitagliptin; MET, metformin; SU, Sulfonylurea; TZD, thiazolidinedione; T2DM, type 2 diabetes mellitus; BMI, body mass index; FPG, fasting plasma glucose.

aDenotes initial combination of sitagliptin with metformin.
Treatment outcome of sitagliptin

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(n=18) and 30 mg in the case of gliclazide (modified-release form; MR) (n=4), but no statistically significant reductions were observed (ΔHbA1c=-0.2%, P=0.63) (Table 2, Fig. 1).

The statistically significant reduction in FPG between visit 1 and visit 5 (ΔFPG) was only observed in the initial coadministration of SITA and MET group (ΔFPG=-26.8 mg/dL, P=0.002), and there were no significant changes in the group changed from SU (ΔFPG=4 mg/dL, P=0.776), the add-on to SU group (ΔFPG=-5.1 mg/dL, P=1.0), the add-on to MET group (ΔFPG=2.36 mg/dL, P=0.74), or the group changed from TZD (ΔFPG=-7.2 mg/dL, P=0.192) (Table 2, Fig. 2).

Side effects were reported in eight of the 99 patients. One patient showed symptoms of hypoglycemia, one patient had an increase in body weight, five patients showed signs of indigestion, abdominal discomfort and gastrointestinal problems, and one patient reported problems with facial swelling. All patients who complained of gastrointestinal symptoms were being treated with MET combination therapy. There were no serious adverse effects associated with SITA.

Analysis on the treatment failure group

SITA treatment failure was experienced by 12 patients (five males, seven females): four patients changed from or added to insulin treatment, four patients added to MET treatment, one on the patient had initial combination of SITA with MET, one patient switching from SU, and one patient changed from TZD.

Compared to 87 patients in the treatment maintenance group, the diabetes duration of the treatment failure group was longer (7.6 vs. 2.8 year, P=0.008), the HbA1c level at visit 1 was higher (8.6% vs. 7.0%, P=0.001) and the FPG was higher (193 vs. 151 mg/dL, P=0.003). Age, gender, BMI, fasting insulin, and C-peptide level before taking the anti-diabetic drugs, history of hypertension, and hyperlipidemia were not significantly different between the treatment failure and maintenance group (Table 3).

Subanalysis on tertiles of ΔHbA1c in the treatment maintenance group

During treatment, the ΔHbA1c in the 87 patients who continued treatment were categorized into tertiles. Good response group (tertile 3 of ΔHbA1c) had a large improvement in blood glucose level, and low response group (tertile 1 of ΔHbA1c) experienced a small improvement in blood glucose level. Compared to low response group, the HbA1c level at visit 1 of good response group was higher (7.5% vs. 6.6%, P<0.001) and the duration of diabetes was shorter (2.0 vs. 4.0 year, P=0.009); however, age, fasting glucose level, BMI, fasting insulin level, and C-peptide level were not significantly different between

Table 2. Efficacy of subgroups on HbA1c and fasting plasma glucose change (ΔHbA1c and ΔFPG) by sitagliptin from visit 1 to visit 5

<table>
<thead>
<tr>
<th></th>
<th>HbA1c levels, %</th>
<th>FPG, mg/dL</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Baseline</td>
<td>ΔHbA1c</td>
</tr>
<tr>
<td>SITA+METa</td>
<td>7.3</td>
<td>-1.1±0.79</td>
</tr>
<tr>
<td>Adding on MET</td>
<td>7.1</td>
<td>-0.6±0.89</td>
</tr>
<tr>
<td>Adding on SU</td>
<td>7.7</td>
<td>-0.5±0.58</td>
</tr>
<tr>
<td>Swiching from SU</td>
<td>6.8</td>
<td>-0.2±0.67</td>
</tr>
<tr>
<td>Swiching from TZD</td>
<td>7.3</td>
<td>-0.3±0.71</td>
</tr>
</tbody>
</table>

FPG, fasting plasma glucose; SITA, sitagliptin; MET, metformin; SU, Sulfonylurea; TZD, thiazolidinedione.

aDenotes initial combination of sitagliptin with metformin, bStatistical significances were tested by Wilcoxon Signed Rank test.

DISCUSSION

In this study, the treatment outcomes of SITA were analyzed at one university hospital for an average duration of ten months. The patients were divided into groups based on the classification of treatment regimens and were retrospectively reviewed. Significant improvements in HbA1c and blood glucose levels were observed after a few months in the initial coadministration of SITA with MET group, the add-on to MET or SU groups and in the group switched from TZD. The efficacy of the SITA showed a similar effect as that of the SU, usually the minimum amount of commercial units.

Although there were many complaints of gastrointestinal discomfort, all of them were from the group coadministered with MET. Among those who complained, no clinically significant adverse effects were observed. Compared to the treatment maintenance group, the duration of diabetes of the treatment failure group was longer and the HbA1c and FPG were higher. However, the age, gender, BMI, fasting insulin, fasting C-peptide, antihypertensive medication, antihyperlipidemia medication history, and smoking history were not significantly different in these groups.

After the recent development of drugs that increase the levels of the incretin hormones GLP-1 and GIP, it was found that commercialized DPP IV-inhibitor prolonged and enhanced incretin hormone activity. As a result of these drugs, insulin secretion from pancreatic β cells is stimulated based on blood sugar level; glucagon secretion from pancreatic α-cells is suppressed to manage blood glucose, pancreatic cell dysfunction...
is prevented, and hypoglycemia and weight gain are rare [4,11,12].

SITA (Januvia®) is the first DPP IV-inhibitor that has been licensed for use in type 2 diabetes patients; It has been reported to not affect the pharmacokinetics of other oral diabetes drugs such as MET, SU, and rosiglitazone [10,13]. The validity and reliability of SITA have been demonstrated in type 2 diabetes patients because blood glucose level is improved and side effects from SU or insulin-like treatment drugs such as hypoglycemia and weight gain are very uncommon. FDA approved the use of SITA in type 2 diabetes and drug indications are as follows: SITA monotherapy, combination therapy with MET, dual add-on therapy to SU, MET, TZD, triple oral therapy with MET, SU/TZD, and add-on to insulin. Domestically, however, only the add-on to MET has been acknowledged for uncontrolled type 2 diabetic patients.

Several randomized double-blind clinical trials have evaluated the efficacy of oral sitagliptin in type 2 diabetic patients with HbA1c level ranged from 6.5-8% to 10-12%. Initial coadministration therapy of SITA with MET significantly lowered HbA1c, FPG and two-hour postprandial blood glucose levels more than did MET monotherapy after 24 weeks of treatment, and maintained the effect for 54 to 104 weeks [14-16]. When SITA was added to MET, the reductions in HbA1c and FPG were sustained during an 18- to 24-week observation period [17-19], and when added to SU, improvements in blood glucose were maintained for 24 weeks [20]. Additionally, when 100 mg of both SITA and MET were combined, there was no comparative difference in the level of reduction in HbA1c level between 52 weeks of MET and 5 mg glipizide [21] or 18 weeks of an 8-mg rosiglitazone and MET combination therapy [19]. In the present study, significant improvements in HbA1c level were found in every group except the group switched from SU. Two to three months after the initial treatment, the significant decrease in HbA1c level plateaued and was maintained for ten months. The group switched from SU was administered an average dosage of 1.8 mg glimepiride and an average dosage of 30 mg gliclazide MR.

Unlike HbA1c, FPG showed an immediate improvement only in the initial MET and SITA combination group. However, the values of FPG in other four groups were less recorded and the standard deviation of FPG was higher than the standard deviation of the HbA1c. These results indicate a limitation to our retrospective study.

The average body weight change in the patients in the group switched from TZD during the observation period was 1.1 kg, but it was not significant (n=14, P=0.124). The 18-week study by Scott et al. [19] demonstrated that, while the addition of TZD to MET gained an average of 1.5 kg, the addition of SITA to MET lost an average of 0.4 kg, a 1.9-kg difference between the two groups. Therefore, SITA can be used as an alternative to avoid the weight gain associated with TZD.

In the meta-analysis of SITA, the risk of hypoglycemia is associated with the combination with SU or insulin treatment and gastrointestinal side effects is associated with MET [20,22,23]. In the present study, one patient who reported having hypoglycemia had previously taken SU, and the five patients who had reported gastrointestinal side effects were all taking MET;

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**Table 4.** Baseline demographic and clinical differences of treatment maintenance group according to HbA1c change (ΔHbA1c, HbA1c at visit 1–HbA1c at visit 5) tertiles, excluding 2nd tertile group

<table>
<thead>
<tr>
<th>Variable</th>
<th>1st tertile (ΔHbA1c ≤0.2%)</th>
<th>3rd tertile (ΔHbA1c &gt;0.7%)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. (%)</td>
<td>29 (33.3)</td>
<td>30 (34.5)</td>
<td></td>
</tr>
<tr>
<td>Female, n (%)</td>
<td>15 (52.0)</td>
<td>22 (73.0)</td>
<td>0.198</td>
</tr>
<tr>
<td>Age, yr</td>
<td>56.0 (48.0, 60.0)</td>
<td>55.0 (46.0, 65.0)</td>
<td>0.525</td>
</tr>
<tr>
<td>Duration of T2DM, yr</td>
<td>4.0 (2.5, 7.4)</td>
<td>2.0 (0.9, 3.0)</td>
<td>0.009*</td>
</tr>
<tr>
<td>HbA1c, %</td>
<td>6.6 (6.2, 6.9)</td>
<td>7.5 (7.1, 8.2)</td>
<td>&lt;0.001*</td>
</tr>
<tr>
<td>FPG, mg/dL</td>
<td>131 (118, 151)</td>
<td>151 (137, 170)</td>
<td>0.082</td>
</tr>
<tr>
<td>BMI, kg/m²</td>
<td>26.7 (23.9, 29.1)</td>
<td>25.2 (23.8, 28.6)</td>
<td>0.861</td>
</tr>
<tr>
<td>Fasting insulin, μIU/mL</td>
<td>12.3 (8.7, 16.8)</td>
<td>10.4 (8.4, 13.4)</td>
<td>0.791</td>
</tr>
<tr>
<td>Fasting C-peptide, ng/mL</td>
<td>2.8 (2.3, 3.7)</td>
<td>3.4 (2.8, 3.9)</td>
<td>0.762</td>
</tr>
</tbody>
</table>

Data are presented as median (25% confidence interval [CI], 75% CI) or number (%).
*Statistical significances were tested by Kruskal-Wallis and Bonferroni post-hoc analysis.
the side effects of SITA monotherapy could not be determined. The duration of diabetes ($P=0.008$), HbA1c ($P=0.001$) and FPG ($P=0.003$) in the treatment failure group were significantly higher than those of the treatment maintenance group (Table 3). However, only a small number of patients were classified into the failure group, and more domestic participants are needed for further analysis.

When $\Delta$HbA1c of the treatment maintenance group were divided into three parts and analyzed, the low response group had a longer diabetes duration ($P=0.009$) as well as a lower initial HbA1c level ($P<0.001$) (Table 4). However, the low response group did not mean treatment failure and our results suggest that the efficacy of sitagliptin therapy is noninferior to that of other common diabetes drugs such as SU. Additionally, the ghrelin hormone and the parasympathetic nervous system are involved in the regulation of GLP-1 [24,25] and genetic polymorphisms of GLP-1 receptors have been reported [26]. Further understanding the individual responses to fats and carbohydrates and GLP-1 receptor will help predict non- or less-responders to SITA.

This study had several limitations. First, we only selected subjects at a single university hospital and the number of available participants in each group was small due to the selection criteria and classification. Second, the retrospective approach lost some data of clinical and biochemical parameters. Third, the preservation of pancreatic $\beta$-cell function from SITA administration could not be confirmed clinically since oral glucose tolerance tests were not usually performed after administration of SITA. However, according to most of the other reports, significant improvements in the homeostasis model of assessment of $\beta$-cell function (HOMA-$\beta$), the proinsulin-to-insulin ratio, and the insulinogenic index were reported [10, 14,17,18,23]. SITA was found to have no effect on insulin resistance or sensitivity as reflected by HOMA-IR and the quantitative insulin sensitivity check index (QUICKI) [18,20,27]; on the other hand, some reports showed significant improvements in insulin resistance or sensitivity [17,28].

In summary, SITA is efficient for glycemic controls in the cases of the initial coadministration with MET, the add-on therapy to MET or SU and administration switched from TZD or SU. It is also relatively safe over an average of ten months in Korean type 2 diabetes. However, well-designed, prospective and long-term studies are needed to understand the treatment failure or response of SITA.

CONFLICTS OF INTEREST

No potential conflict of interest relevant to this article was reported.

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