Management of dyslipidemia and hyperglycemia with a fixed-dose combination of sitagliptin and simvastatin

Helmut Steinberg
Matt S Anderson
Thomas Musliner
Mary E Hanson
Samuel S Engel

Merck Sharp & Dohme Corp., Whitehouse Station, NJ, USA

Abstract: The risk of death due to heart disease and stroke is up to four times higher in individuals with diabetes compared to individuals without diabetes. Most guidelines that address treatment of dyslipidemia in patients with diabetes consider diabetes a cardiovascular disease (CVD) “risk equivalent” and recommend intensive treatment of dyslipidemia for the purpose of CVD prevention. Statins (3-hydroxy 3-methylglutaryl coenzyme A reductase [HMG-CoA reductase] inhibitors) are first-line agents in achieving lipid goals as an adjunct to diet and exercise and should be used in most patients. In addition to lipid management and blood pressure control, glycemic control is a basic component in the management of diabetes. Glycemic control is achieved by combining diabetes self-management education, diet and exercise, and, where required, antihyperglycemic agents (OHAs). Persistence and adherence to therapy are critical in achieving recommended treatment goals. However, overall compliance with concomitantly prescribed OHAs and statins is low in patients with type 2 diabetes. Fixed-dose combination (FDC) therapies have been shown to improve adherence by reducing pill burden, the complexity of treatment regimen, and, potentially, cost. Based on the available evidence regarding the pharmacokinetics and the efficacy and safety profiles of each component drug, the sitagliptin/simvastatin FDC may provide a rational and well-tolerated approach to achieving better adherence to multiple-drug therapy and improved lipid lowering and glycemic control, with consequent reduction in cardiovascular risk, diabetic microvascular disease, and mortality in diabetic patients for whom treatment with both compounds is appropriate.

Keywords: statin, oral antihyperglycemic agent, diabetes, adherence, cardiovascular disease, microvascular disease

Introduction
The International Diabetes Federation estimates that there are approximately 371 million people in the world living with diabetes, of whom half are undiagnosed.1 In the United States and Europe, the prevalence of this disease is 10.5% and 6.7% of the population, respectively.1 The risk of death due to heart disease and stroke is up to four times higher in individuals with diabetes compared to individuals without diabetes.2 Additional serious long-term consequences associated with diabetes include renal failure, retinopathy, and neuropathy.

Dyslipidemia is a major predisposing factor for atherosclerotic cardiovascular disease (CVD) in the general population as well as in diabetic patients. Elevations in low-density lipoprotein cholesterol (LDL-C) have received the greatest attention from the scientific and clinical community, and it is clear that LDL-C level is at least as strong a predictor of coronary heart disease risk in diabetic patients as it is...
in the general population.\textsuperscript{3} In the UK Prospective Diabetes Study, a 57% increased risk of coronary heart disease was reported for every 1 mmol/L increment in LDL-C.\textsuperscript{4,5} While LDL-C is not often greatly increased in diabetic individuals, the presence of diabetes and/or insulin resistance is associated with profound changes in lipid and lipoprotein metabolism, with resultant alterations in particle distribution within lipoprotein classes. This includes increased numbers of small dense LDL particles,\textsuperscript{6} which are believed to be particularly atherogenic due to their increased endothelial permeability, susceptibility to oxidation and glycation, and ability to bind to proteoglycans in the vessel wall.\textsuperscript{7–9} Many diabetic patients also have increased levels of larger apolipoprotein B-containing lipoproteins, including very low-density lipoprotein remnants of intermediate density, as well as reduced levels of high-density lipoprotein cholesterol (HDL-C), both of which are associated with increased CVD risk.\textsuperscript{6,10}

Because of this increased CVD risk, most guidelines that address treatment of dyslipidemia in patients with diabetes (including those from the American Diabetes Association and American College of Cardiology Foundation guidelines, the Task Force on Diabetes and Cardiovascular Diseases of the European Society of Cardiology, and the European Association for the Study of Diabetes) consider diabetes as a CVD “risk equivalent” and recommend intensive treatment of dyslipidemia for the purpose of CVD prevention.\textsuperscript{6,11–13} These treatment guidelines provide goals for lipids and glucose levels. LDL-C has been identified as the primary therapeutic target for patients with hypercholesterolemia, existing CVD, or other CVD risk factors.\textsuperscript{12,14,15} The Canadian Cardiovascular Society guidelines recommend an LDL-C target of \(\leq 77 \text{ mg/dL} (2.0 \text{ mmol/L})\) or \(\geq 50\%\) decrease in LDL-C in all risk categories when pharmacologic intervention is warranted.\textsuperscript{14} However, the US and European guidelines have designated different LDL-C target levels based on CVD risk.\textsuperscript{12,15} Specifically, attaining LDL-C \(\leq 100 \text{ mg/dL} (\leq 2.6 \text{ mmol/dL})\) is recommended for those at high or moderate CVD risk, and the more aggressive target of \(\leq 70 \text{ mg/dL} (\leq 1.8 \text{ mmol/dL})\) is recommended for patients at very high risk for CVD, including those with diabetes and overt CVD.\textsuperscript{12,15,16}

Statins (3-hydroxy 3-methylglutaryl coenzyme A reductase [HMG-CoA reductase] inhibitors) are first-line agents in achieving lipid goals if diet and exercise are not sufficient, since they are generally well tolerated, highly effective for lowering LDL-C, and have been shown to be capable of substantially decreasing cardiovascular mortality and morbidity.\textsuperscript{17–19} A meta-analysis of 14 randomized trials, which included \(-90,000\) subjects, demonstrated that for every 1.0 mmol/L reduction in LDL-C achieved, there was an approximate 20% reduction in coronary heart disease mortality and a 22% reduction in major CVD events.\textsuperscript{20} These results were consistent across baseline LDL-C levels and similar in subjects with and without diabetes.\textsuperscript{20,21}

In addition to lipid management and blood pressure control, glycemic control is a basic component in the management of diabetes and is achieved, in part, via diabetes self-management education, exercise, and improved diet, the latter elements being the cornerstones of treatment for diabetes and high lipid levels. Guidelines for the treatment of diabetes stress the importance of lowering HbA\textsubscript{1c} to a level below 7% in non-pregnant adults in order to reduce the risk of microvascular (retinopathy and nephropathy) and neuropathic complications.\textsuperscript{16,22} Analyses have suggested that lowering HbA\textsubscript{1c} to 6% is associated with further reductions in the risk of microvascular complications, albeit with substantially increased risk of hypoglycemia.\textsuperscript{23,24} According to these guidelines, intensive HbA\textsubscript{1c} lowering beyond 7% may be warranted in selected individuals. In patients with little comorbidity and with long life expectancy, the patient and physician may opt for glycemic targets as close to normal as possible as long as hypoglycemia does not pose a significant problem.\textsuperscript{16}

Even if patients are treated according to the guidelines with proven treatments, the treatments can only be maximally effective if patients are adherent to the treatment regimen. Persistence and adherence to therapy are critical in achieving recommended treatment goals and improving patient outcomes in chronic conditions such as type 2 diabetes mellitus.\textsuperscript{8,11} It has been shown that non-adherence to medication is associated with significantly \((P < 0.001)\) higher all-cause hospitalization and mortality in diabetic patients,\textsuperscript{25} and overall compliance with oral anti-hyperglycemic agents (OHAs) is low in patients with type 2 diabetes.\textsuperscript{26} Several cross-sectional retrospective analyses estimated that adherence to statins tends to be even lower compared with adherence to OHAs in patients who receive them as concomitant therapy (with adherence rates estimated around 52% for statins to 72% for OHAs).\textsuperscript{27–30}

The reasons for differences in adherence between OHAs and statins are complex and not well understood, and several variables play a role in a patient’s compliance with a prescribed treatment plan.\textsuperscript{31} One potential reason for greater adherence to OHAs versus statins may be the difference in associated symptoms. Dyslipidemia is generally
asymptomatic and less likely to be discussed between a patient and health care provider, whereas glycemic abnormalities often have associated symptoms that may cause apprehension.\textsuperscript{11,32,33} Moreover, there may be a perception on the part of the patient that the purpose of statins is to lower cholesterol rather than to reduce cardiovascular risk; since LDL-C and total cholesterol are generally similar in patients with and without type 2 diabetes,\textsuperscript{34} lipid control may not raise the same level of concern as glucose control. Finally, there may be a perceived lack of benefit from statins on the part of the patient,\textsuperscript{35,36} whereas the benefit from OHAs in relation to glucose control may be easily seen by patients as they self-monitor blood glucose.

Fixed-dose combination (FDC) therapies have been shown to improve adherence by reducing costs, pill burden, and the complexity of treatment regimen.\textsuperscript{37–39} A treatment approach with a FDC that includes a statin and an OHA could be used to improve statin compliance in patients with type 2 diabetes. Sitagliptin (Januvia [Merck, Whitehouse Station, NJ, USA]) is a highly selective dipeptidyl peptidase-4 (DPP-4) inhibitor available for the treatment of hyperglycemia in patients with type 2 diabetes,\textsuperscript{40,41} and simvastatin (Zocor\textsuperscript{TM} [Merck]) is an HMG-CoA reductase inhibitor available as an adjunctive treatment to diet for reducing elevated LDL-C and other lipids in patient with primary hypercholesterolemia and is indicated for reducing the risk of cardiovascular mortality.\textsuperscript{42} An FDC tablet of sitagliptin and simvastatin (Juvisync\textsuperscript{TM} [Merck]) has been approved and provides an option for use in patients for whom treatment with both sitagliptin and simvastatin is appropriate.

**Pharmacological profile of Juvisync\textsuperscript{TM}**

The separate pharmacokinetic profiles of sitagliptin and simvastatin have been extensively characterized in healthy subjects, and previously reviewed;\textsuperscript{43,44} however, there is no published literature discussing the pharmacokinetic profile of the FDC tablet of sitagliptin/simvastatin, which is formed by compressing a common sitagliptin blend with a common simvastatin granule to form a bilayer tablet. The FDC tablet is available in doses of (sitagliptin/simvastatin) 100 mg/10 mg, 100 mg/20 mg, and 100 mg/40 mg, as well as dosages for patients with moderate renal insufficiency (50 mg/10 mg, 50 mg/20 mg, and 50 mg/40 mg).

The potential for simvastatin to alter sitagliptin pharmacokinetics was explored in an open-label randomized two-period crossover study in ten healthy men and women, wherein the pharmacokinetics of sitagliptin were compared after administration of a single dose of sitagliptin 100 mg alone or in the presence of steady state simvastatin (on day 5 of a 7-day course of simvastatin 80 mg once daily).\textsuperscript{45} Simvastatin had no clinically relevant effect on sitagliptin: the geometric mean ratio of (sitagliptin + simvastatin)/sitagliptin (90% confidence interval [CI]) of AUC\textsubscript{0–∞} = 1.01 (0.97, 1.05) and C\textsubscript{max} = 1.12 (90% CI 1.00, 1.26). Conversely, the potential for sitagliptin to alter simvastatin pharmacokinetics was explored in an open-label randomized two-period crossover study in 12 healthy men and women, wherein the pharmacokinetics of simvastatin were compared after administration of a single dose of simvastatin 20 mg alone or in the presence of steady state sitagliptin (on day 5 of a 5-day course of sitagliptin 200 mg once daily).\textsuperscript{46} Sitagliptin had no clinically meaningful effect on simvastatin: the geometric mean ratio of (simvastatin lactone + sitagliptin)/simvastatin lactone (90% CI) of AUC\textsubscript{0–last} = 0.85 (0.60, 1.22), C\textsubscript{max} = 0.80 (0.51, 1.26), and for simvastatin acid AUC\textsubscript{0–last} = 1.12 (0.93, 1.35) and C\textsubscript{max} = 1.06 (0.86, 1.32).\textsuperscript{46} Consequently, no dose adjustments are recommended for these drugs when coadministered.\textsuperscript{46}

A demonstration of definitive bioequivalence between the FDC tablet of sitagliptin/simvastatin at two tablet strengths (100 mg/10 mg and 100 mg/80 mg) as compared to coadministration of the corresponding doses of sitagliptin and simvastatin as individual tablets was conducted in two separate open-label randomized two-period single-dose definitive bioequivalence studies presented in this review. Primary endpoints were to compare the pharmacokinetics of sitagliptin (using AUC\textsubscript{0–∞} and C\textsubscript{max}), simvastatin lactone (using AUC\textsubscript{0–last} and C\textsubscript{max}), and simvastatin acid (using AUC\textsubscript{0–last} and C\textsubscript{max}), using [0.80, 1.25] as allowable bounds of variation for each analyte. Both studies enrolled 100 healthy subjects ranging in age from 18 to 55 years (Table 1). The 90% CIs of the observed geometric mean ratios (GMR) (Juvisync\textsuperscript{TM}/[sitagliptin + simvastatin]) for the AUC\textsubscript{0–∞} and C\textsubscript{max} of sitagliptin and the AUC\textsubscript{0–last} and C\textsubscript{max} of simvastatin lactone and simvastatin acid were all within the prespecified bounds in both studies, demonstrating bioequivalence of the FDC tablet and the individual tablets across the assessed dose range (Table 2). Demonstration of bioequivalence for all other manufactured tablet strength combinations, as compared to their separately administered components, was supported through in vitro dissolution data and formulation-proportionality arguments.

The potential for pharmacokinetic interactions between the FDC tablet and digoxin were also investigated as previously conducted pharmacokinetic-interaction studies of single-dose digoxin, administered with and without mul-
tiple doses of sitagliptin or of simvastatin, showed that each medication slightly increased the plasma pharmacokinetics of digoxin relative to digoxin administered alone. Consistent with these earlier investigations, concomitant administration of multiple doses of sitagliptin 100 mg and simvastatin 80 mg with a single 0.5 mg dose of digoxin moderately increased the plasma AUC0–last of digoxin (Table 3), demonstrating that multiple-dose coadministration of sitagliptin/simvastatin 100 mg/80 mg had a roughly additive pharmacokinetic effect on the single-dose pharmacokinetics of digoxin relative to digoxin administration with either sitagliptin or simvastatin alone. However, these effects are not considered clinically important in the context of appropriate monitoring as already recommended in clinical practice with digoxin. Accordingly, patients receiving digoxin concomitantly with JuvisyncTM should be monitored appropriately by their physician as if they were receiving digoxin alone or in combination with either simvastatin or sitagliptin alone.

**Efficacy**

The rationale for using OHAs such as sitagliptin is to improve glucose control, thereby reducing the risk of microvascular disease without inducing hypoglycemia or weight gain in patients with type 2 diabetes. The rationale for using a statin (such as simvastatin) is to improve lipid and lipoprotein levels to within individual therapeutic targets, thereby reducing CV morbidity and mortality. Although no results are available from randomized clinical trials assessing the FDC tablet of sitagliptin/simvastatin, the results from controlled clinical studies of the individual treatments provide information on their respective efficacy that may be extrapolated to infer efficacy and safety of the FDC.

**Sitagliptin**

Four randomized double-blind placebo-controlled trials of 12 to 24 weeks in duration were conducted in type 2 diabetic subjects who were drug-naïve or whose prior treatment with

### Table 1 Subject disposition in bioequivalence studies

<table>
<thead>
<tr>
<th>Patient characteristics</th>
<th>100 mg sitagliptin/10 mg simvastatin</th>
<th>100 mg sitagliptin/80 mg simvastatin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total patients randomized (n)</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Male age (range) in years</td>
<td>41 (19–54)</td>
<td>61 (20–55)</td>
</tr>
<tr>
<td>Female age (range) in years</td>
<td>59 (18–53)</td>
<td>39 (20–55)</td>
</tr>
<tr>
<td>Completed (n)</td>
<td>93</td>
<td>98</td>
</tr>
<tr>
<td>Discontinued (n)</td>
<td>7</td>
<td>2</td>
</tr>
<tr>
<td>Clinical AEs (n)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Laboratory AEs (n)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Withdraw consent (n)</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td>Protocol violation (n)</td>
<td>1</td>
<td>2</td>
</tr>
</tbody>
</table>

**Notes:**

- The PK parameter values following a single-dose administration of study drug were compared using separate linear mixed-effect models appropriate for a two-period crossover design. The linear mixed-effect model contained factors for sequence, period, and treatment as fixed effects, and subject-within-sequence as a random effect. A log transformation was applied to the AUC and Cmax data. Back-transformed summary statistics and inferential results were reported for PK parameter values. The 90% CIs were compared to the prespecified bounds of [0.80, 1.25]. *represents multiplication.

- **Abbreviations:** AUC, area under the curve; Cmax, maximum concentration; CI, confidence interval; GMR, geometric mean ratio; PK, pharmacokinetic.

### Table 2 Statistical comparisons for the plasma PK parameters of sitagliptin, simvastatin, and simvastatin acid after a single-dose administration of FDC sitagliptin/simvastatin 100 mg/10 mg or 100 mg/80 mg tablet, or coadministration of corresponding doses of sitagliptin and simvastatin and individual tablets

<table>
<thead>
<tr>
<th>PK parameter</th>
<th>100 mg/10 mg</th>
<th>100 mg/80 mg</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sitagliptin</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AUC0–last (nM/hr)</td>
<td>1.01 (0.99, 1.02)</td>
<td>0.99 (0.98, 1.00)</td>
</tr>
<tr>
<td>AUC0–last (nM/hr)</td>
<td>1.01 (1.00, 1.03)</td>
<td>0.99 (0.98, 1.00)</td>
</tr>
<tr>
<td>Cmax (nM)</td>
<td>1.03 (0.98, 1.07)</td>
<td>0.98 (0.94, 1.02)</td>
</tr>
<tr>
<td><strong>Simvastatin</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AUC0–last (nM/hr)</td>
<td>1.07 (0.99, 1.16)</td>
<td>0.99 (0.93, 1.05)</td>
</tr>
<tr>
<td>Cmax (nM)</td>
<td>1.13 (1.05, 1.21)</td>
<td>0.98 (0.92, 1.06)</td>
</tr>
<tr>
<td><strong>Simvastatin acid</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AUC0–last (nM/hr)</td>
<td>1.03 (0.96, 1.11)</td>
<td>0.93 (0.87, 0.98)</td>
</tr>
<tr>
<td>Cmax (nM)</td>
<td>1.04 (0.97, 1.12)</td>
<td>0.95 (0.88, 1.02)</td>
</tr>
</tbody>
</table>

**Notes:**

- The PK parameter values following a single-dose administration of study drug were compared using separate linear mixed-effect models appropriate for a two-period crossover design. The linear mixed-effect model contained factors for sequence, period, and treatment as fixed effects, and subject-within-sequence as a random effect. A log transformation was applied to the AUC and Cmax data. Back-transformed summary statistics and inferential results were reported for PK parameter values. The 90% CIs were compared to the prespecified bounds of [0.80, 1.25]. *represents multiplication.

- **Abbreviations:** AUC, area under the curve; Cmax, maximum concentration; CI, confidence interval; GMR, geometric least-squares mean ratio; PK, pharmacokinetic.

### Table 3 Statistical comparisons for the plasma PK parameters after administration of a single dose of digoxin 0.5 mg alone or sitagliptin 100 mg once daily and simvastatin 80 mg once daily for 9 days and a single dose of digoxin 0.5 mg on day 5 in healthy male and female subjects

<table>
<thead>
<tr>
<th>PK parameter</th>
<th>(Digoxin + sitagliptin + simvastatin)/digoxin</th>
</tr>
</thead>
<tbody>
<tr>
<td>AUC0–last (nM/hr)</td>
<td>1.26 (1.13, 1.41)</td>
</tr>
<tr>
<td>Cmax (nM)</td>
<td>1.41 (1.20, 1.66)</td>
</tr>
</tbody>
</table>

**Notes:**

- Back-transformed least-squares mean and CI from linear mixed-effects model performed on natural log-transformed values. The effects and variability due to subjects within sequence (a random effect) and sequence, period, and treatment (fixed effects), as well as the within-subject variability of the PK parameters were estimated using a mixed-effect analysis model appropriate for the two-period study design. A log transformation was applied to the digoxin AUC0–last and Cmax data. A 90% CI was constructed for the difference in least-squares means on the log scale. Exponentiation of the log scale 90% CI provided a 90% CI for the GMR ([(digoxin + sitagliptin + simvastatin)/digoxin alone]). The 90% CIs were compared to the prespecified bounds of [0.80, 1.25]. Summary statistics and comparisons were provided for Cmax of digoxin.

- **Abbreviations:** AUC, area under the curve; Cmax, maximum concentration; CI, confidence interval; GMR: geometric least-squares mean ratio; PK: pharmacokinetic.
OHAs had been discontinued.47–50 These studies demonstrated that treatment with sitagliptin monotherapy resulted in significantly greater reductions from baseline in HbA\textsubscript{1c} (placebo-subtracted HbA\textsubscript{1c} reductions ranged from −0.48% to −0.94%), as well as a significantly greater proportion of subjects achieving HbA\textsubscript{1c} < 7% compared with placebo.47–50 In these studies, the usual clinical dose of sitagliptin was 100 mg daily. Two of the studies also assessed a 200 mg dose47,50 and one of the studies used 50 mg or 100 mg as needed in elderly patients who had moderate renal insufficiency.48 In addition, these same trials showed that sitagliptin treatment, compared with placebo, resulted in significantly greater reductions in fasting plasma glucose (FPG) and 2-hour post-meal glucose, and significantly improved homeostatic model assessments (HOMA-\textbeta)\textsuperscript{47–50} and pro-insulin/insulin ratio.\textsuperscript{47,48,50} Results from a study that was extended from 1 to 2 years indicated that sitagliptin as monotherapy or as initial combination therapy with metformin (500 mg or 1000 mg twice daily) provided substantial and sustained glycemic control for up to 2 years.\textsuperscript{51,52} Generally, the beneficial effect of sitagliptin on HbA\textsubscript{1c} versus placebo was consistent among subgroups defined by age, sex, race, baseline BMI, and prior use of OHAs.\textsuperscript{47–50,53,54} One study specifically assessed type 2 diabetic subjects older than 65 years, and showed significantly greater reductions in HbA\textsubscript{1c} 2-hour post-meal glucose, and FPG, as well as improvements in HOMA-\textbeta over 24 weeks with sitagliptin (100 mg once daily or 50 mg once daily, dependent on renal function) compared with placebo, consistent with reports in younger subjects.46 Moreover, four additional studies confirmed these findings in Asian populations: three in Japanese subjects and one each in Chinese, Indian, and Korean subjects (placebo-subtracted HbA\textsubscript{1c} reductions ranged from −0.96% to −1.4%), as well as showing significant improvements in indices of glycemic control such as FPG, 2-hour post-meal glucose, and HOMA-\textbeta in these populations.\textsuperscript{55–58}

Sitagliptin has also been assessed in subjects inadequately controlled on single or dual OHA therapy or single OHA plus insulin at study entry. Eight clinical trials (sitagliptin 100 mg/day once daily added to metformin ≥1500 mg/day,\textsuperscript{59–61} insulin ≥15 IU/day alone or combined with metformin ≥1500 mg/day,\textsuperscript{62} metformin ≥1500 mg/day plus rosiglitazone ≥4 mg/day,\textsuperscript{63} glimepiride ≥4 mg/day or glimepiride ≥4 mg/day plus metformin ≥1500 mg/day,\textsuperscript{64} ongoing metformin ≥1500 mg/day and pioglitazone ≥30 mg/day,\textsuperscript{65} or pioglitazone 30–45 mg/day)\textsuperscript{66}) demonstrated significantly greater reductions in HbA\textsubscript{1c} (placebo-subtracted HbA\textsubscript{1c} reductions ranged from −0.5% to −1.0%; all \textit{P} < 0.001), as well as a significantly greater proportion of subjects achieving HbA\textsubscript{1c} < 7% and significantly greater improvements in FPG versus placebo (all \textit{P} < 0.001). In the seven studies that assessed 2-hour post-meal glucose, significantly greater reductions were observed with sitagliptin add-on treatment versus placebo (all \textit{P} < 0.001).\textsuperscript{59–65} Several studies also demonstrated significant improvements in HOMA-\textbeta,\textsuperscript{59,61,63,65} pro-insulin,\textsuperscript{66} pro-insulin/insulin ratio,\textsuperscript{63,65} and fasting insulin secretion,\textsuperscript{66} which supports the premise that treatment with sitagliptin improves beta-cell function. When sitagliptin 100 mg add-on was compared with glipizide 5–20 mg/day added on to metformin ≥1500 mg/day, noninferiority in HbA\textsubscript{1c} lowering was demonstrated at the 1-year time point (as noted above\textsuperscript{65}) and no meaningful differences were observed during the 1-year extension period, although no inferential testing was conducted for the results of the extension.\textsuperscript{67} Compared with glipizide over 2 years, greater durability and generally better maintenance of beta-cell function were observed with sitagliptin 100 mg.\textsuperscript{67} In another trial comparing sitagliptin or rosiglitazone with placebo when added on to metformin, similar changes from baseline in HbA\textsubscript{1c} and HOMA-\textbeta, and similar proportions of subjects achieving HbA\textsubscript{1c} < 7% were observed in subjects taking either active treatment regimen.\textsuperscript{68} As in the studies of subjects who were drug-naïve or discontinued from prior treatment, the greater effect of sitagliptin on HbA\textsubscript{1c} versus placebo was consistent among subgroups defined by age, sex, race, baseline BMI, and prior use of OHAs.\textsuperscript{50–64,66,68} In the majority of clinical trials, both add-on and drug-naïve/-washout, there was a significant interaction between baseline HbA\textsubscript{1c} and treatment effect.\textsuperscript{60,61,63,64,66,68,69} Specifically, in subjects with higher baseline HbA\textsubscript{1c}, reductions in HbA\textsubscript{1c} were greater than in subjects with lower baseline HbA\textsubscript{1c}.\textsuperscript{50–53,54,60–61,63,64,66,68,69} Overall, sitagliptin has been shown to improve glycemic control and beta-cell function in type 2 diabetic patients both as a monotherapy and as combination therapy in clinical trials. Most of these trials were relatively short-term, and more long-term studies are needed to confirm the extended effects on glycemic control and beta-cell function.

Simvastatin

The benefits of cholesterol-lowering with simvastatin in the general population have been well-established. The Scandinavian Simvastatin Survival Study (4S) assessed the efficacy of simvastatin 20–40 mg/day versus placebo in 4444 subjects with coronary heart disease for a median duration of 5.4 years.\textsuperscript{17} Simvastatin treatment resulted in mean reductions in total cholesterol, LDL-C, and triglycerides of...
25%, 35%, and 10%, respectively, increased HDL-C by 8%, and demonstrated a 42% reduction in the risk of coronary death compared with placebo (relative risk [RR] = 0.58, 95% CI 0.46–0.73; P = 0.0003). Moreover, simvastatin treatment resulted in a significant reduction in the risk of a major coronary event including coronary death, nonfatal definite or probable myocardial infarction (MI), silent MI, or resuscitated cardiac arrest compared with placebo (RR = 0.66, 95% CI 0.59–0.75; P < 0.00001). These results were consistent in women and in subjects ≥ 60 years.17 The Heart Protection Study, a placebo-controlled double-blind study conducted with 20,532 patients at high risk of developing a major coronary event (ie, subjects with known coronary disease, other occlusive arterial disease, or diabetes), compared treatment with simvastatin 40 mg/day versus placebo with respect to all-cause death and death from coronary heart disease over 5 years of treatment.70 The placebo-adjusted differences between groups in total cholesterol, LDL-C, and triglycerides were −0.8 mmol/L, −1.0 mmol/L, and −0.2 mmol/L, respectively. Simvastatin 40 mg/day significantly reduced all-cause mortality (P < 0.0003) and coronary death rate (18%, P = 0.0005), as well as producing a 38% reduction in the incidence of first nonfatal MI (P < 0.0001), a 25% reduction in first stroke (P < 0.0001), and a 24% reduction in first revascularization procedure (P < 0.0001).70

In addition to clinical outcomes and lipid-lowering in the general population, the benefits of simvastatin have also been assessed specifically in diabetic subjects. In a post hoc analysis of the 4S trial that included subjects with diabetes or impaired fasting glucose, reductions in cholesterol levels and the risk of cardiovascular events were consistent with those observed in the full cohort.71 Diabetic subjects treated with simvastatin experienced a significantly reduced incidence of major coronary events (RR = 0.58; P = 0.001) and revascularizations (RR = 0.52; P = 0.005) compared with placebo, and subjects with impaired fasting glucose also had significantly reduced incidence of major coronary events (RR = 0.62; P = 0.003), revascularization (RR = 0.57; P = 0.009), and total and coronary mortality (RR = 0.57; P = 0.02 and RR = 0.45; P = 0.007, respectively) compared with placebo.72 Analysis of the HPS also assessed simvastatin efficacy in subjects with and without diabetes and showed an effect consistent with the full cohort,72 that is, simvastatin produced highly significant reductions (22%; P < 0.0001) in the first event rate for major coronary events, strokes, and revascularizations in subjects with type 2 diabetes. Furthermore, in type 2 diabetic subjects without occlusive arterial disease or other coronary arterial disease, the reductions were even higher (33%; P < 0.0003). Based on these results, the authors concluded that cholesterol-lowering statin therapy can have a beneficial effect on coronary outcomes in type 2 diabetic patients even if they do not have coronary artery or other occlusive arterial disease.

Safety and tolerability
Sitagliptin

The safety and tolerability profile of sitagliptin was assessed in 10,246 subjects with type 2 diabetes mellitus by pooling results from 19 controlled clinical trials (conducted for up to 2 years), comparing sitagliptin (n = 5429) with either placebo or active comparator (non-exposed group, n = 4817).73 Reports of overall adverse events, serious adverse events, and the number of deaths were similar in the sitagliptin-treated and non-exposed groups, except for a higher incidence of drug-related adverse events in the non-exposed group. This was primarily attributed to the higher rate of hypoglycemia in the non-exposed group (mainly due to use of a sulfonylurea as a comparator agent in two studies). In a separate pooled analysis that removed sulfonylureas and insulin as background or comparator, the incidence rates of hypoglycemia were similar between groups (3.1 versus 3.3 per 100 patient-years in sitagliptin and non-exposed groups, respectively).73 There was a greater number of reports of diarrhea in the non-exposed group, likely due to the use of metformin, and more reports of constipation in the sitagliptin group.73 In addition to hypoglycemia, abdominal pain, diarrhea, gastritis, weight gain, and paresthesia were reported more often in the non-exposed group compared with the sitagliptin group. Constipation was the only drug-related adverse event reported more often in the sitagliptin-treated group.73

Simvastatin

In general, statins have been shown to be safe and well-tolerated in clinical trials. However, important adverse events that are associated with statin use include muscle complaints, ranging from muscle weakness and cramps to myalgia (with and without elevated creatine kinase levels) and rhabdomyolysis, as well as increases in liver enzymes > 3 times the upper limit of normal.74,75 The risk of muscle toxicity may be increased in certain patients, including older individuals (≥ 75 years), those with comorbid conditions and/or taking concomitant medications, and those with impaired hepatic or renal function.74,76 In addition, higher statin doses (eg, simvastatin 80 mg) may increase the risk to a greater extent than moderate doses. In 2011, new dosing recommendations for simvastatin were introduced worldwide based on the dose-response relationship to the risk of myopathy, and the 80 mg dose of simvastatin is limited to patients who have been taking it for 12 months or
more without evidence of myopathy. Although the labeling for every statin reflects a warning for each agent’s potential for myotoxicity, both observational and clinical trial data indicate that the risk of myotoxicity and rhabdomyolysis is low for all marketed statins. Increases in liver transaminases have also been reported with statin use compared with placebo, and these are commonly reversible with discontinuation of treatment or dose reduction.

Recently, there has been concern regarding the effect of statins on glycemia and new-onset type 2 diabetes. Results of the JUPITER trial and a large meta-analysis of statin trials suggested a slight increase in the risk of new onset of type 2 diabetes with the use of statins. Because of these concerns, many statins, including simvastatin, contain a statement regarding this issue in the “warnings and precautions” section of the prescribing information. The Juvisync™ prescribing information also contains a statement regarding this issue in the “warnings and precautions” section. The effects of simvastatin treatment on glucose control in type 2 diabetic patients have been evaluated in clinical trials. In the HPS study, after a 4.6 year follow up (n = 1087), there was no significant difference between treatment groups in the increase in HbA1c concentration (0.15% [standard deviation (SD) = 0.09] versus 0.12% [SD = 0.09], difference 0.03% [SD = 0.13]; P = 0.8), suggesting that simvastatin did not affect diabetes control in this study. Moreover, in this study, simvastatin treatment did not result in any increases in reports of other diabetes-related outcomes, such as hospital admissions for unstable diabetes or laser treatments for retinopathy, in the 5963 participants known to have diabetes. In a smaller double-blind placebo-controlled trial that assessed the efficacy of simvastatin 20–40 mg/day for 6 months in subjects with non-insulin-dependent diabetes mellitus, there were no significant changes in measures of glycemic control, including fasting plasma glucose, insulin, C-Peptide, or HbA1c, with simvastatin treatment. Finally, concerns were raised that statin use could increase the risk of cancers that take longer than 5 years to emerge clinically. Three separate trials with extended follow-up of 15 years (including 5 years of in-trial follow-up), 11.3 years (including 3.3 years of in-trial follow-up), and 11 years (including 5.3 years of in-trial follow-up) showed consistently that incident cancer did not increase in the statin treated subjects compared with placebo groups.

Coadministration and fixed-dose combination

In a pooled subgroup analysis of 19 clinical studies of sitagliptin including 1582 patients whose background therapy included simvastatin, the incidence of adverse events was similar between patients treated with sitagliptin and simvastatin (n = 827) and patients treated with placebo or active comparator and simvastatin (n = 755). Among the patients whose background therapy included simvastatin, 3.3% of the sitagliptin-treated group and 4.2% of controls discontinued due to adverse events. The short-term bioequivalence studies (described above) assessing the pharmacological profile of the FDC sitagliptin/simvastatin also provide some insight into the safety and tolerability profiles. In both studies that assessed bioequivalence, there were no serious adverse experiences or serious laboratory adverse events reported, and no subjects discontinued due to an adverse event. It is important to note that the bioequivalence studies were open label single-dose studies conducted in healthy subjects rather than in patients with diabetes, and the results should therefore be interpreted with caution. The safety results of the bioequivalence studies indicate that the coadministration of sitagliptin and simvastatin was generally well tolerated and without indication of emergent clinical or laboratory safety signals when dosed acutely.

Conclusion

Cardiovascular risk is increased in patients with diabetes, including in those with normal LDL-C levels. Addressing this risk will require statin treatment in most patients. However, underutilization and insufficient intensification of statins and OHA treatment may contribute to a failure to achieve recommended treatment targets. In addition to patient counseling on lifestyle changes and adherence to medication regimens, patients may benefit from use of FDC therapies, which have been shown to increase adherence through reducing pill burden, complexity of treatment regimen, and cost. Based on the available evidence regarding the pharmacokinetics, efficacy, and safety profile of each component drug, the sitagliptin/simvastatin FDC may provide a well-tolerated approach to achieving improved lipid lowering and glycemic control, with consequent reduction in cardiovascular risk, diabetic microvascular disease, and mortality in diabetic patients for whom treatment with both compounds is appropriate.

Acknowledgment

The authors would like to thank Dr Jennifer Rotonda, PhD, of Merck Sharp & Dohme Corp. for assistance with preparation of the manuscript.

Disclosure

All authors are employees of Merck Sharp & Dohme Corp. and may own stock or hold stock options in the company. The authors report no other conflicts of interest in this work.
References


