The clinical application of linagliptin in Asians

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Abstract: Asia has a growing diabetic population. Linagliptin, a member of dipeptidyl peptidase-4 inhibitor class, is unique in its nonlinear pharmacokinetics with the characteristics of rapid attainment of steady state, little accumulation, predominantly nonrenal route of elimination, prolonged terminal half-life, and sustained inhibition of dipeptidyl peptidase-4 enzyme. No clinically relevant difference in pharmacokinetics was observed between Asians and non-Asians. The management of type 2 diabetes is increasingly challenging with the progression of disease, especially with the requirements of minimal hypoglycemia, weight gain, fluid retention, and other adverse effects. Linagliptin was efficacious and well-tolerated in Asian type 2 diabetes patients with or without renal or hepatic dysfunctions, comparable to that in Caucasians. This review will focus on the usage of linagliptin in clinical studies in Asians.

Keywords: linagliptin, Asians, type 2 diabetes

Introduction

In 2014, the International Diabetes Federation (IDF) announced that the prevalence of diabetes was 8.3%, with 387 million people living with diabetes.1 The IDF also predicted that there would be 205 million more people affected worldwide in 2035, which is a rapid growth with greater magnitude than previously expected.2 It is estimated that >60% of the whole population with diabetes will have an Asian derivation, as it remains the world’s most densely populated region.3 Such a tremendous number puts a huge financial burden, owing to direct health care expenditure and impairment of productivity, on developing regions.

Asians with the same age, sex, and BMI, particularly those of South Asian lineage, have a higher body fat percentage and are more prone to central obesity and insulin resistance (IR) than their western counterparts. Furthermore, insufficiency of the compensatory insulin secretion capacity, which could not increase proportionately with the severity of IR, is another characteristic of Asian type 2 diabetic population.3

Another characteristic of Asian diabetic patients is the higher risk of renal complications when compared with their Caucasian counterparts.4 Many oral antihyperglycemic agents need dose adjustments or to be avoided in patients with diabetic nephropathy, even under periodic renal function examinations.5 Most patients with diabetic nephropathy have to finally turn to exogenous insulin therapy, despite its adverse effects including increased rates of hypoglycemia due to impaired renal function, exacerbated fluid retention, and weight gain.5 New and more effective treatments are under development.

Dipeptidyl peptidase-4 (DPP-4) inhibitors suppress the enzymatic degradation of incretin hormones, including glucagon-like peptide-1 (GLP-1) and glucose-dependent insulino-tropic polypeptide (GIP), which could promote the biosynthesis of insulin and further stimulate insulin release glucose-dependently in addition to oral glucose load, and this phenomenon was absent with intravenous glucose infusion.7 When...
administered at pharmacological doses, GLP-1 also has other non-insulinotropic effects, including inhibition of postprandial glucagon excursions, suppression of gastric emptying and intestinal mobility, induction of satiety, and weight loss. Moreover, GLP-1 exhibits a protecting/preserving effect of β-cells in animal experiments. Thus, DPP-4 inhibitors play a key role in the maintenance of glucose homeostasis through the potentiation of the action of GLP-1 and GIP. Incretin effects are impaired in patients with type 2 diabetes mellitus (T2DM), despite a similar incretin hormone response to healthy controls after increasing oral glucose loads, emphasizing the necessity for the supplementation of exogenous incretins or an enhanced action of endogenous incretins.

Linagliptin, based on a xanthine scaffold structure, not only shares many properties with other members of DPP-4 inhibitor class, such as low risk of hypoglycemia and weight neutrality, but also has a special pharmacokinetic (PK) profile that is clinically relevant. Unlike other DPP-4 inhibitors, linagliptin is predominantly excreted unchanged in feces, with no necessity of dose adjustment in the case of renal impairment since renal excretion only makes a minor contribution to the overall elimination (primarily nonrenal-clearance pathway). Extensive binding with plasma protein and a long terminal half-life make once-daily oral administration possible. Coadministration with some other antidiabetic and cardiovascular drugs results in low potential of drug–drug interaction. Considering the unique characteristics of linagliptin, we will here review the updated publications about the usage of linagliptin in Asians.

**Pharmacokinetics and pharmacodynamics**

In a Phase II, randomized, double-blind, placebo-controlled study, 72 Japanese T2DM patients were assigned to receive placebo or linagliptin 0.5 mg, 2.5 mg, or 10 mg once daily for consecutive 28 days according to the proportion of 1:1:1:1. Linagliptin was rapidly absorbed with a median 

max,ss

of –1.5 hours across all dose ranges. At steady state, neither 

AUC, nor 

max,ss

parameters of systemic linagliptin exposure did increase dose-proportionally across all dose ranges, and this was a reflection of the unique nonlinear PK profile of linagliptin. Linagliptin was widely distributed in the body, as the geometric mean (gMean) of apparent volume of distribution (V_d,F,ss) was from 4,090 L for the 0.5 mg dose to 21,200 L for the 10 mg dose at steady state. The terminal half-life was long enough (223–260 hours) for sustained suppression of DPP-4. In contrast, the accumulation half-life was short (10–38.5 hours), resulting in rapid attainment of steady state. Steady-state AUC ranged from 2.9-fold for 0.5 mg dose to 1.2-fold for 10 mg dose greater than after a single dose, indicating little accumulation. Apparent clearance (CL/F,ss) and renal clearance (CL,R,ss) after oral administration increased with dose. Even so, renal excretion only played a minor part in the overall elimination of linagliptin, with steady-state urinary excretion not exceeding 7% of the administered dose. These PK parameters were comparable to that of linagliptin in healthy Japanese and Chinese subjects and Caucasian T2DM patients as shown in Table 1. Consistently, the prescribing information indicates no dose adjustments given the fact that race has no clinically relevant influence on the PK of linagliptin, regardless of Caucasian, Hispanic, Blacks, and Asians.

Plasma protein binding was extensive and in a concentration-dependent manner. An in vitro experiment had shown that the protein-binding percentage of linagliptin decreased with increasing concentration (from ~99% at a concentration <1 nM to 70%–80% at a concentration >100 nM), as a result of the high affinity but easily saturated binding capacity to plasma DPP-4 at therapeutic concentrations. Based on the nonlinear PK and concentration dependency of plasma protein binding, researchers had developed a two-compartment model, with linagliptin distributed in the central and peripheral compartments. In a previous study, the 28-day treatment of multiple-dose linagliptin produced maximal inhibition of DPP-4 by 66.7%, 89.6%, and 92.9% for 0.5 mg, 2.5 mg, and 10 mg dose groups, respectively. DPP-4 inhibition at 24-hour postdose was 46%, 78%, and 90% for 0.5 mg, 2.5 mg, and 10 mg dose groups, respectively. Typically, DPP-4 inhibition ≥80% >24 hours is therapeutically required to achieve a maximal hypoglycemic effect. However, in this study, there were no data about the PK/pharmacodynamics (PD) of linagliptin 5 mg once daily in Japanese T2DM patients. With these experimental data, researchers simulated the plasma DPP-4 inhibition of linagliptin 5 mg once daily in a well-established and carefully verified target-mediated drug disposition model, suggesting that 5 mg linagliptin once daily resulted in 84.2% DPP-4 inhibition at 24 hours after the last dose, which was therapeutically meaningful. Similarly, data from 56 healthy male Japanese subjects participating in a 12-day Phase I, double-blind, placebo-controlled study indicated that after multiple doses, ≥80% DPP-4 inhibition after 24-hour dosing interval was achieved in 33%, 100%, 100% of subjects in the linagliptin 2.5 mg, 5 mg, and 10 mg once-daily group, with maximal DPP-4 inhibition being 92%–93% for all
Table 1 Pharmacokinetics of linagliptin in Asians compared with Caucasians

<table>
<thead>
<tr>
<th>Subjects</th>
<th>Regimen</th>
<th>T_{max} (h)</th>
<th>C_{max} (nmol/L)</th>
<th>AUC (nmol ⋅ h/L)</th>
<th>Terminal t_{1/2} (h)</th>
<th>Accumulation t_{1/2} (h)</th>
<th>RA_{AUC}</th>
<th>V/F (L)</th>
<th>CL/F (mL/min)</th>
<th>CL_{R} (mL/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Healthy Chinese(^\text{15})</td>
<td>5 mg SD</td>
<td>1.75</td>
<td>10.4</td>
<td>150</td>
<td>82.4</td>
<td>–</td>
<td>–</td>
<td>1.910</td>
<td>268</td>
<td>22.3</td>
</tr>
<tr>
<td></td>
<td>5 mg MD for 6 d</td>
<td>1.5</td>
<td>14.1</td>
<td>204</td>
<td>103</td>
<td>11.5</td>
<td>1.35</td>
<td>7.730</td>
<td>866</td>
<td>68.1</td>
</tr>
<tr>
<td>Healthy Japanese(^\text{14})</td>
<td>1 mg SD</td>
<td>1.77</td>
<td>4.27</td>
<td>62</td>
<td>104</td>
<td>–</td>
<td>–</td>
<td>1.260</td>
<td>140</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>2.5 mg SD</td>
<td>2.0(^\text{a})</td>
<td>5.92</td>
<td>108</td>
<td>96.9</td>
<td>–</td>
<td>–</td>
<td>1.430</td>
<td>171</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>5 mg SD</td>
<td>6(^\text{a})</td>
<td>9</td>
<td>159</td>
<td>105</td>
<td>–</td>
<td>–</td>
<td>2.090</td>
<td>231</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>10 mg SD</td>
<td>1.5(^\text{a})</td>
<td>23.1</td>
<td>294</td>
<td>113</td>
<td>–</td>
<td>1.31</td>
<td>3.060</td>
<td>314</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>2.5 mg MD for 12 d</td>
<td>3.75(^\text{a})</td>
<td>7.79</td>
<td>133</td>
<td>142</td>
<td>10–15</td>
<td>1.28</td>
<td>8.180</td>
<td>664</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>5 mg MD for 12 d</td>
<td>2.25(^\text{a})</td>
<td>12</td>
<td>193</td>
<td>143</td>
<td>10–15</td>
<td>1.14</td>
<td>11.300</td>
<td>913</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>10 mg MD for 12 d</td>
<td>4(^\text{a})</td>
<td>21.8</td>
<td>285</td>
<td>175</td>
<td>10–15</td>
<td>1.14</td>
<td>18.700</td>
<td>1,240</td>
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<tr>
<td>Japanese T2DM patients(^\text{13})</td>
<td>0.5 mg MD for 28 d</td>
<td>1.5(^\text{a})</td>
<td>5.02</td>
<td>89.4</td>
<td>240</td>
<td>38.5</td>
<td>2.88</td>
<td>4.090</td>
<td>197</td>
<td>4.5</td>
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<tr>
<td></td>
<td>2.5 mg MD for 28 d</td>
<td>1.5(^\text{a})</td>
<td>11</td>
<td>164</td>
<td>223</td>
<td>10.7</td>
<td>1.27</td>
<td>10.400</td>
<td>537</td>
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<tr>
<td></td>
<td>10 mg MD for 28 d</td>
<td>1.25(^\text{a})</td>
<td>44</td>
<td>373</td>
<td>260</td>
<td>10</td>
<td>1.16</td>
<td>21.200</td>
<td>945</td>
<td>65</td>
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<tr>
<td>Healthy Caucasians(^\text{23})</td>
<td>5 mg SD</td>
<td>2.05(^\text{a})</td>
<td>4.4</td>
<td>75.3</td>
<td>79.9</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Caucasian T2DM patients(^\text{16})</td>
<td>5 mg SD</td>
<td>1.47(^\text{a})</td>
<td>5.71</td>
<td>100</td>
<td>69.7</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>1 mg MD for 12 d</td>
<td>1.48(^\text{a})</td>
<td>4.53</td>
<td>81.7</td>
<td>121</td>
<td>23.9</td>
<td>2.03</td>
<td>–</td>
<td>–</td>
<td>14.0</td>
</tr>
<tr>
<td></td>
<td>2.5 mg MD for 12 d</td>
<td>1.42(^\text{a})</td>
<td>6.58</td>
<td>117</td>
<td>113</td>
<td>12.5</td>
<td>1.37</td>
<td>–</td>
<td>–</td>
<td>23.1</td>
</tr>
<tr>
<td></td>
<td>5 mg MD for 12 d</td>
<td>1.53(^\text{a})</td>
<td>11.1</td>
<td>158</td>
<td>131</td>
<td>11.4</td>
<td>1.33</td>
<td>–</td>
<td>–</td>
<td>70</td>
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<tr>
<td></td>
<td>10 mg MD for 12 d</td>
<td>1.34(^\text{a})</td>
<td>13.6</td>
<td>190</td>
<td>130</td>
<td>8.59</td>
<td>1.18</td>
<td>–</td>
<td>–</td>
<td>59.5</td>
</tr>
</tbody>
</table>

Note: \(^\text{a}\) Median value. \(T_{\text{max}}\) values are geometric mean (gMean) unless specified.

Abbreviations: \(T_{\text{max}}\), time to reach \(C_{\text{max}}\); \(C_{\text{max}}\), maximum (peak) plasma drug concentration; \(RA_{AUC}\), accumulation ratio calculated from area under the plasma concentration–time curve at steady state; \(V/F\), apparent volume of distribution; \(CL/F\), apparent clearance; \(CL_{R}\), renal clearance; SD, single dose; MD, multiple dose; d, days; h, hours; T2DM, type 2 diabetes mellitus.
Linagliptin is the most potent DPP-4 inhibitor currently available in the market. Under the identical in vitro experimental conditions, linagliptin produced an IC\textsubscript{50} of approximately 1 nM, compared with sitagliptin of 19 nM, alogliptin of 24 nM, saxagliptin of 50 nM, and vildagliptin of 60 nM. As a competitive inhibitor, linagliptin had a K\textsubscript{i} of 1 nM. Moreover, linagliptin reversibly but tightly binds DPP-4 enzyme, with approximately tenfold slower dissociation rate from DPP-4 than another DPP-4 competitive inhibitor, vildagliptin, resulting in prolonged DPP-4 inhibition even if the free fraction had been cleared from the circulation.\textsuperscript{22} Both in healthy and in type 2 diabetic Japanese patients, 50% and 80% inhibition of maximal DPP-4 inhibition occurred at a concentration range of 2–4 nmol/L and 4–6 nmol/L, respectively, comparable to the data in Caucasians.\textsuperscript{13,14,23} Besides its potency, linagliptin is a highly selective DPP-4 inhibitor, being \( \approx 10,000 \)-fold more selective for DPP-4 compared to DPP-8 and DPP-9 in vitro. The selectivity, to some extent, ensures the safety of the treatment, given the detrimental effect of DPP-8 and DPP-9 inhibition on multiorgans and immune system observed in animals.\textsuperscript{22}

With its primarily nonrenal elimination route and potency, which means easily saturable binding to its target DPP-4 enzyme at low doses, it seems that linagliptin is unlikely to accumulate even in T2DM patients with renal impairment. In a 24-week, multicenter, Phase III study, nearly half of participants were Asians. No statistical significance was reported in the mean linagliptin trough concentrations over time between patients with normal renal function (estimated glomerular filtration rate [eGFR] \( \geq 90 \) mL/min/1.73 m\(^2\)) and those with mildly (eGFR 60 to \(< 90 \) mL/min/1.73 m\(^2\)) or moderately (eGFR 30 to \(< 60 \) mL/min/1.73 m\(^2\)) impaired renal function: 8.0±7.3 nmol/L, 8.0±7.6 nmol/L, and 6.6±1.8 nmol/L, respectively.\textsuperscript{24}

**Efficacy**

**Monotherapy**

In a multicenter, randomized, double-blind, placebo-controlled, Phase III study comparing linagliptin (5 mg once daily, n=336) with placebo (n=167), 46.1% of participants with T2DM had an Asian derivation. A total of 24 weeks of intervention resulted in an adjusted mean change in HbA\textsubscript{1c} of −0.69% (\( P<0.0001 \)), favoring linagliptin over placebo, with no significant difference between Asian and Caucasian, the two main population recruited.\textsuperscript{24}

Similarly, in a randomized, double-blind, parallel-group, Phase IIb/III study carried out in 561 Japanese T2DM patients, linagliptin 5 mg and 10 mg yielded placebo-corrected adjusted mean reduction in HbA\textsubscript{1c} of −0.87% and −0.88%, respectively, at week 12. Furthermore, linagliptin monotherapy also showed superiority in terms of glucose control when compared with another active comparator, voglibose 0.2 mg tid, with adjusted mean treatment difference in HbA\textsubscript{1c} of −0.32% (\( P=0.0003 \)) and −0.39% (\( P<0.0001 \)) at week 26 for linagliptin 5 mg and 10 mg once daily. Reduction in proinsulin/insulin ratio reached statistical significance in both linagliptin 5 mg and 10 mg group. Linagliptin 5 mg also showed a trend of nonsignificance to improve indexes of β-cell function, including fasting insulin, disposition index (DI), and homeostasis model assessment (HOMA) of IR.\textsuperscript{21}

**Linagliptin as add-on therapy to one oral antidiabetic drug**

A 52-week, multicenter, open-label study evaluates the efficacy of linagliptin 5 mg once daily as add-on therapy to one background oral antidiabetic drug (OAD), including biguanide, glinide, glitazone, sulphonylurea (SU), and α-glucosidase inhibitor (A-GI), in 618 Japanese patients with T2DM inadequately controlled. Out of these, patients on SU (2:1) or A-GI (1:1) were randomized to linagliptin 5 mg qd or metformin (twice or thrice daily, \( \leq 2,250 \) mg/day) (randomized group). Subjects on other background therapies receive linagliptin 5 mg qd (nonrandomized group). Results showed that linagliptin therapy led to significant reduction in mean HbA\textsubscript{1c}, level of −0.7% to 0.9% in nonrandomized group and that the hypoglycemic effect was comparable between linagliptin and metformin as add-on therapy to SU or A-GI, while once-daily regimen of linagliptin was more convenient.\textsuperscript{26}

In a 2-year, randomized (1:1 in linagliptin: glimepiride), double-blind, parallel-group, Phase III study carried out in a large population of 1,551 T2DM patients inadequately controlled, including 12% Asians, linagliptin 5 mg once daily exhibited noninferiority to glimepiride 1–4 mg once daily as add-on therapy to metformin, with change in adjusted mean HbA\textsubscript{1c} being similar for linagliptin (−0.16%) and glimepiride (−0.36%). There was no ethnic difference in HbA\textsubscript{1c} changes in each treatment group. Of note, linagliptin added to metformin was associated with significantly lower hypoglycemia rates, decreased body weight, and fewer cardiovascular events.\textsuperscript{27} Initial combination of linagliptin with metformin also substantially improved the glycemic control of patients with previously uncontrolled T2DM (including 32.5% Asians), and no interaction of significance between linagliptin and metformin was observed.\textsuperscript{28}
Linagliptin as add-on therapy to basal insulin with or without OADs

In a multicenter, multinational, randomized (1:1 to linagliptin vs placebo), double-blind, Phase III study of linagliptin versus placebo as add-on therapy to basal insulin alone or in combination with one to two OADs (metformin and/or pioglitazone) in previously inadequately controlled patients with T2DM, 12.2% of the participants were Asians. The basal insulin remained unchanged for the first 24 weeks, and during this period, adjusted mean HbA1c changed by −0.58% for linagliptin and 0.07% for placebo (treatment difference −0.65%, \(P<0.0001\)). The placebo-corrected change in fasting plasma glucose (FPG) was −0.6 mmol/L (\(P<0.0001\)). After week 24, titration of basal insulin was allowed for both groups to achieve a target FPG of 6.1 mmol/L, which was more in line with the clinical practice. At week 52, the mean change in FPG of linagliptin-treated group was not significantly different from the placebo group, while the adjusted mean change in HbA1c for linagliptin and placebo was −0.48% and 0.05%, respectively, leading to a treatment difference of −0.53% (\(P<0.0001\)), which was maintained for up to 76 weeks. Of note, subgroup analysis found that variables such as renal function and age did not interact with treatment, while significant interaction (\(P<0.1\)) was observed between treatment and other variables including race (\(P=0.0603\)), which suggested that Asians were the greatest benefited population (placebo-adjusted mean change in HbA1c was greatest for patients from Asian [−1%] and of Asian origin [−0.93%]).

Linagliptin as add-on therapy to metformin plus pioglitazone

A multicenter, randomized (2:1 to linagliptin vs placebo), double-blind, Phase III study carried out in 271 T2DM patients with inadequate glycemic control on metformin and pioglitazone (Asians accounted for 69%), compared linagliptin vs placebo as add-on therapy to background regimen. After 24-week intervention, linagliptin significantly improved glycemic control, with a placebo-adjusted mean change in HbA1c of −0.57% (\(P<0.0001\)). The treatment difference between the groups was significant (\(P<0.0001\)) at each evaluated time point and did not vary over time significantly. Subanalysis of HbA1c changes by country revealed that India and Philippines, the two countries representing Asia in this study, achieved a significant reduction in HbA1c, with a placebo-adjusted change from baseline of −0.9% (\(P<0.0001\)) and −0.8% (\(P=0.014\)), respectively. Linagliptin was also superior in reducing FPG and in the attainment of an HbA1c target of 7% or 6.5%. Moreover, HOMA-β was 27% greater from baseline for linagliptin group vs placebo group (\(P=0.0055\)), indicating that linagliptin may have a protective effect on β-cells.30

Linagliptin as add-on therapy to metformin plus sulfonylurea

In a prespecified Chinese patient subanalysis of a multinational, randomized, placebo-controlled clinical trial, linagliptin in triple combination with existing background regimen of metformin and SU (n=144) led to significant improvement compared with placebo group (n=48), consistent with the efficacy profile of the parent study.6,31 Placebo-corrected adjusted mean change in HbA1c at week 24 for linagliptin was −0.68% from a mean baseline of 8.1%, comparable to −0.62% reduction in the parent study. Placebo-corrected adjusted mean decrease in FPG at week 24 from baseline was −1 mmol/L for linagliptin-treated group and −0.7 mmol/L for the overall population.6,31 Moreover, significantly more patients achieved a target HbA1c <7% in the linagliptin group (29.8% vs 10.4%, \(P=0.01\)). The assessments of β-cell function and IR (HOMA-β and HOMA-IR) showed a trend of improvements, but they did not show statistical significance. However, treatment difference in DI (adjusted mean change from baseline was 3.71 mmol/L vs −2.83 mmol/L for linagliptin vs placebo, respectively) was statistically significant (\(P=0.0007\)), suggestive of improved β-cell function.

Efficacy data were extracted from four multiethnic, randomized, double-blind, placebo-controlled Phase III studies of 24-week duration for pooled analysis of 5 mg linagliptin once daily for Asian T2DM patients with uncontrolled glycemia (743 Asians in linagliptin group vs 286 Asians in placebo group) as monotherapy, as add-on therapy to metformin, as add-on therapy to metformin plus SU, and as initial therapy in combination with pioglitazone. Despite its post hoc nature, the analysis still provided evidence that linagliptin was efficacious in Asians, with a treatment difference at week 24 from baseline in HbA1c of −0.79% (\(P<0.0001\)), favoring linagliptin compared with placebo.32 The superiority of linagliptin over placebo was also reflected in significantly reducing FPG (placebo-corrected mean change was −1 mmol/L, \(P<0.0001\)) and postprandial glucose (PPG) (placebo-corrected mean change was −3.2 mmol/L, \(P=0.0002\)).32 These results showed that the efficacy of linagliptin in Asians was in consistence with that of the overall multiethnic population in the four parent studies.
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Safety in patients with renal impairment

There is no direct design to confirm the renal safety of linagliptin. A pooled analysis, consisting of 1,102 patients from Asian studies, showed that linagliptin 5 mg once daily was efficacious and well-tolerated. Despite one-third of subjects having mild (eGFR 60 to 90 mL/min) to-moderate (eGFR 30 to < 60 mL/min) renal impairment, no dose adjustment was made in accordance with ASAS and AACE. Linagliptin 5 mg once daily was efficacious and well-tolerated in patients previously on OADs or drug-naïve. Studies of linagliptin exclusively in Asian T2DM patients are listed in Table 2.

Table 2 Studies of linagliptin exclusively in Asian T2DM patients

<table>
<thead>
<tr>
<th>Studies</th>
<th>Patients</th>
<th>Race</th>
<th>Intervention</th>
<th>Study duration</th>
<th>Major outcomes</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Horie et al13</td>
<td>72 patients on lifestyle intervention or 1–2 OADs</td>
<td>Japanese</td>
<td>Linagliptin of different doses: 0.5 mg, 2.5 mg, 10 mg daily and placebo</td>
<td>4 wks</td>
<td>1) Change of HbA1c at 4 wks</td>
<td>1) −0.31% (P &lt; 0.01), −0.2% (NS), −0.44% (P &lt; 0.001) (0.5 mg, 2.5 mg, 10 mg linagliptin) vs 0.4% (placebo)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>2) Change of FPG</td>
<td>2) −11.5 mg/dL, −13.6 mg/dL, −25 mg/dL (0.5 mg, 2.5 mg, 10 mg linagliptin) vs −3.2 mg/dL (placebo), P &lt; 0.05</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>3) Change of GLP-1</td>
<td>3) 3.4 mmol/L (NS), 4.1 mmol/L (NS), 7.2 mmol/L (P &lt; 0.001) (0.5 mg, 2.5 mg, 10 mg linagliptin) vs 2.5 mmol/L (placebo)</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>4) Adverse events</td>
<td>4) 10.5%, 27.8%, 22.2% (0.5 mg, 2.5 mg, 10 mg linagliptin) vs 35.3% (placebo) (NS)</td>
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<td>Kawamori et al22</td>
<td>561 patients previously on OADs or drug-naïve</td>
<td>Japanese</td>
<td>Linagliptin 5 mg and 10 mg daily vs placebo vs voglibose 0.2 mg tid</td>
<td>12 wks (linagliptin vs placebo) 26 wks (linagliptin vs voglibose)</td>
<td>1) Change of HbA1c at 12 wks</td>
<td>1) −0.24%, −0.25% (5 mg, 10 mg linagliptin) vs 0.63% (placebo), P &lt; 0.0001</td>
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<td>2) Change of HbA1c at 26 wks</td>
<td>2) −0.13%, −0.19% (5 mg, 10 mg linagliptin) vs 0.19% (voglibose)</td>
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<td>3) Drug-related adverse event</td>
<td>3) 9.4% linagliptin 5 mg, 8.8% linagliptin 10 mg, 10% placebo at 12 wks, 11.3% linagliptin 5 mg, 10.6% linagliptin 10 mg, 18.5% voglibose at 26 wks</td>
</tr>
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the prescribing information, which indicates no necessity of dose adjusting for renal impairment.\textsuperscript{17,32} Studies carried out exclusively in Japanese patients also showed that the linagliptin was efficacious and safe regardless of baseline renal function\textsuperscript{25,26} and that kidney function maintained and even improved for linagliptin-treated patients.\textsuperscript{26} A retrospective study in 145 Japanese patients in real clinical practice demonstrated the consistence of the efficacy of 5 mg linagliptin once daily >6 m for subjects with varying degrees of renal function, with no association of reduction of HbA\textsubscript{1c} level and baseline renal function.\textsuperscript{37} A multiciracial study including Asians found that renal function had no clinically relevant impact on the long-term exposure of linagliptin in patients. Geometric mean trough $C_{\text{ss}}$ in patients with normal, mildly, moderately, and severely impaired renal function was 5.93 nmol/L, 6.07 nmol/L, 7.34 nmol/L, and 8.13 nmol/L, respectively, with a maximal increase of 37.1\% in trough systemic exposure.\textsuperscript{39} Since a single dose of 600 mg was well-tolerated in healthy Caucasian volunteers,\textsuperscript{22} the 37.1\% increase in trough exposure of multiple dosing 5 mg once daily did not seem to have clinical relevance. Moreover, a pooled analysis including Asians revealed that linagliptin 5 mg qd of 18–24 weeks was efficacious and well-tolerated for T2DM patients with renal and cardiovascular risk factors of microalbuminuria (urine albumin-to-creatinine ratio of 30–300 mg/g) and hypertension, with renal function remaining stable.\textsuperscript{38} Furthermore, 24-week 5 mg linagliptin once-daily monotherapy was shown retrospectively efficacious and well-tolerated for Indian patients with new-onset diabetes after kidney transplant, with no impact on immune-suppressant levels and eGFR.\textsuperscript{40}

### Safety in patients with hepatic impairment

Pharmacological study in Caucasians concluded that hepatic functions, regardless of mild (Child–Pugh class A), moderate (Child–Pugh class B), and severe (Child–Pugh class C) hepatic impairment, had no clinically significant impact on the exposure of single or multiple dosing of linagliptin 5 mg once daily; therefore, no dose adjustment was required in patients with impaired hepatic functions.\textsuperscript{41} However, in the multinational studies including Asian participants or clinical studies carried out exclusively in Japanese subjects mentioned earlier, patients with hepatic impairment, defined as serum levels of either alanine transaminase, aspartate transaminase, or alkaline phosphate greater than three times the upper limit of normal, were excluded.\textsuperscript{6,24–26,29–31,35,36,43} Nonetheless, race had no influence on the PD of linagliptin, indicating the applicability of the earlier conclusion to Asians.
In addition, linagliptin as monotherapy or additional therapy had no clinically meaningful impact on clinical laboratory tests, physical examination, and vital signs.

Discussion

Worldwide, 77% of people affected with diabetes are now living in low- and middle-income countries, with People’s Republic of China and India being the “top 2” countries that have the largest number of population with diabetes in 2013 and the same for the estimation in 2030. Asian diabetic population is also continuing to rise sharply, driven by rapid nutrition transition associated with socioeconomic development, reduced physical activity related to sedentary lifestyle, genetic susceptibility, and other contributing factors. Asian diet structure is unique in its refined rice and wheat basis, both with high glycemic index and dietary glycemic load values, leading to higher proportion of postchallenge hyperglycemia. Postprandial hyperglycemia was found to be associated with endothelial dysfunction, an early marker of cardiovascular complications. DPP-4 inhibitor class could effectively improve glucose homeostasis and is particularly effective in reducing PPG excursions in a glucose-dependent manner. As a xanthine-based DPP-4 inhibitor, linagliptin was proposed to have an antioxidative stress action in vitro at least partly via its inhibition of xanthine oxidase activity, which may bring about a protective effect on endothelia. Furthermore, not only protection against oxidative stress but also improvement of cardiomyocyte ultrastructure was observed in animal models. Correspondingly, it was reported that linagliptin may have cardiovascular effects.

In contrast with Caucasians, decline in β-cell function occurred early and may be the primary abnormality for Asian population at risk of or with T2DM. Linagliptin had shown protective effect of β-cells, in part attributable to amelioration of glucos- and lipotoxicity in Asians. In addition, animal and ex vivo experiments demonstrated that linagliptin could reduce lymphocyte insulinitis, restore proliferation, and inhibit apoptosis of β-cells via suppression of oxidative stress and proinflammatory cytokine responses, which may also be a part of the underlying mechanisms contributing to the β-cell-protecting effect. Consistently, it has been proposed that DPP-4 inhibitors were particularly efficacious for Asians compared with other ethnic groups. Moreover, preclinical experiments had demonstrated the immunomodulatory effects of DPP-4 inhibitors through both GLP-1-dependent and -independent pathways, in favor of the usage of DPP-4 inhibitors in type 2 diabetes since immune dysfunctions were also present in T2DM. Therefore, the β-cell-preserving effect of DPP-4 inhibitors was another promising prospect for the treatment of T2DM. This is an area requiring further investigation since the hypothesis was based on animal or in vitro studies and biomarkers.

As a potent DPP-4 inhibitor, the HbA1c-lowering efficacy of linagliptin 5 mg qd was comparable to sitagliptin 100 mg qd, vildagliptin 50 mg bid, and saxagliptin 5 mg qd in Asians as monotherapy or add-on therapy to metformin, despite the lowest bioavailability of ~30% as compared with sitagliptin 87%, vildagliptin 85%, and saxagliptin 50%. However, there is still a scarcity of head-to-head comparisons of all DPP-4 inhibitors in Asian diabetic patients, since an identical design in a more homogenous population will provide more compelling evidence.

Renal insufficiency results in reduced clearance of hypoglycemic agents and impaired kidney glyconeogenesis. In addition, diabetic patients with renal dysfunctions are often elderly and concomitantly complicated with diabetic neuropathy and/or cognitive disorder, and these factors act together to put the patients in a greater risk of hypoglycemia. Furthermore, renal failure continues to increase despite strict glucose and blood pressure management strategy, including renin–angiotensin system blocking with a role of albuminuria lowering and renoprotection. Thus, novel treatments aimed to maintain and even improve kidney function are in urgent need. Different from other DPP-4 inhibitors, linagliptin is excreted primarily through a nonrenal pathway in unchanged form, together with the PK property of little accumulation, making dose adjustment unnecessary even in the case of renal insufficiency. Race has no impact on PK and PD of linagliptin in diabetic patients with or without nephropathy. Clinical studies confirmed that renal impairment had no clinically relevant impact on PK and PD of linagliptin in patients with diabetes including Asians. Linagliptin was efficacious, safe, and well-tolerated in Asian patients with T2DM and various degrees of renal impairment, without the AEs of increased hypoglycemia, fluid retention, weight gain, or deterioration of renal functions. It was indicated that linagliptin had an additional albuminuria-lowering effect, on the basis of preexisting renin–angiotensin–aldosterone system inhibitors, in T2DM patients with albuminuria (urinary albumin-to-creatinine ratio of 30–3,000 mg/g), independent of alterations in plasma glucose and blood pressure. Preclinical experiments had demonstrated that animals treated with linagliptin manifested a mitigation of kidney fibrosis, associated with the suppression of
endothelial-to-mesenchymal transition, without an alteration of glucose level. Supportive evidence included an in vitro study that showed that linagliptin inhibited the activation and release of TGF-β, followed by suppression of related profibrotic downstream pathways. Moreover, significant improvement of albuminuria, reduction of advanced glycation end products-receptor (AGE-RAGE)-induced oxidative stress and intercellular adhesion molecule-1 (ICAM-1) mRNA level in the kidney, and amelioration of glomerular lymphocyte infiltration were observed in linagliptin-treated rats. However, whether or not it is the same situation in human still needs further exploration. And studies carried out exclusively in Asian patients with varying degrees of renal dysfunction or in comparison with other ethnic group aimed at comparative PK and PD are needed.

Suppression of the enzymatic degradation of incretins by DPP-4 inhibitors could potentiate the physiological action of GLP-1 and GIP. Of note, unlike GLP-1, the suppressive effect of DPP-4 inhibitors on postprandial glucagon secretion is quite weak and of short duration. Moreover, DPP-4 inhibitors have no significant impact on GI mobility and are weight neutral, indicating that DPP-4 inhibitors do not regulate glucose homeostasis via exactly the same mechanisms as native incretins. In addition, the subsets of immune cells regulated by GLP-1 and DPP-4 inhibitors were not uniformly identical. It was also speculated that the increase in GLP-1 concentration caused by DPP-4 inhibition was insufficient to explain the hypoglycemic effect, and possibly by the same token, DPP-4 inhibitors had no significant impact on GI mobility and body weight. More researches are needed for the specific mechanisms.

With the progression of the disease, the patients fail to achieve the glycemic control target on monotherapy (usually metformin) need to switch to dual therapy and even triple therapy of OADs (with metformin being the cornerstone) before the initiation of insulin treatment. Incretin mimetic and DPP-4 inhibitors have similar antidiabetic efficacy and a broader safety and tolerability profile when compared with other OADs such as SU and thiazolidinediones. Furthermore, linagliptin was demonstrated to be efficacious and safe in patients for whom metformin was contraindicated owing to renal impairment or intolerability.

Conclusion

In conclusion, similar to that in Caucasians, linagliptin has a nonlinear PK with the characteristics of rapid attainment of steady state, little accumulation, primarily nonrenal route of elimination, long-terminal half-life, and sustained inhibition of DPP-4 enzyme in the Asians. Linagliptin was efficacious and well-tolerated in Asian T2DM patients, comparable to that in Caucasians. However, most conclusions are derived from pooled data of multiracial studies or post hoc comparison of data in different races. Therefore, more researches carried out exclusively in Asian patients or for head-to-head comparison with other ethnic groups are needed.

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Disclosure

The authors report no conflicts of interest in this work.

References


