Comparison of Pharmacokinetic and Pharmacodynamic Effects of Two Hydrofluoroalkane Formulations of Salmeterol

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Abstract

Introduction: To compare the Pharmacokinetic (PK) and Pharmacodynamic (PD) effects of two Hydrofluoroalkane (HFA) formulations of salmeterol xinafoate (Test HFA formulation, Cipla Ltd., India; Reference HFA formulation, Allen and Hanburys, UK) administered using pressurized metered dose inhalers.

Methods: Three separate randomized, crossover, PK studies and one PD study comparing the efficacy and safety of the two HFA formulations of salmeterol xinafoate (25 μg per actuation) in healthy subjects were conducted. The PK assessments of the two formulations were done without charcoal blockade, with charcoal blockade, and with a Volumatic spacer device using a single dose. A PD study was also conducted to evaluate the systemic exposure of the two formulations using three different doses (50 μg, 150 μg and 300 μg).

Results: In the PK study without charcoal, the 90% CI for the difference between the two formulations for AUC0-12h was within the bioequivalence limits of 80-125%; however, Cmax marginally exceeded the upper bioequivalence limit to 136%. In the PK study with charcoal, the 90% CI for the difference between the two formulations for Cmax was within the bioequivalence limits of 80-125%; however, AUC0-12h marginally exceeded the upper bioequivalence limit to 128%. The impact of marginally higher systemic exposure was therefore further evaluated in the PD study. The PD study confirmed there were no greater systemic safety effects of the test formulation on the primary PD endpoints such as heart rate and serum potassium as well as on other safety PD endpoints such as blood glucose and QTc interval. The PK study with spacer demonstrated bioequivalence between the test and reference formulations. Both formulations were safe and well tolerated.

Conclusion: The test HFA formulation of salmeterol was therapeutically equivalent to the reference HFA formulation of salmeterol when used with and without a spacer.

Keywords: Pharmacokinetics; Salmeterol; Metered dose inhalers

Introduction

According to the World Health Organization (WHO), 235 million people suffer from asthma and 65 million people suffer from moderate to severe Chronic Obstructive Pulmonary Disease (COPD) [1,2]. In 2009, 3388 people died of asthma; the age-adjusted death rate was 1.1 per 100,000 people [3]. More than 3 million people died of COPD in 2005, which corresponds to 5% of all deaths globally. Asthma costs the European economy approximately 17.7 billion Euros while COPD accounts for 38.6 billion Euros every year [4].

Bronchodilators are prescribed either on as-needed basis or on regular basis to prevent or reduce symptoms of asthma and COPD. Long-acting inhaled β2-agonists such as salmeterol are convenient and more effective in producing maintained symptom relief than short-acting β2-agonists [5,6]. Salmeterol significantly improves Forced Expiratory Volume in 1 s (FEV1) and lung volumes, dyspnea, quality of life, and exacerbation rate; and reduces the rate of hospitalization when used in combination with a corticosteroid [7-12].

The reference Hydrofluoroalkane (HFA) formulation, Serevent Evohaler (Allen and Hanburys, United Kingdom) contains salmeterol xinafoate. Cipla Limited, India has developed an alternative formulation of salmeterol xinafoate (test HFA formulation) as a cost-effective alternative for treatment of asthma and COPD. Both formulations are administered using Pressurized Metered Dose Inhalers (pMDIs).

A vital element of a therapeutic equivalence study using a clinical endpoint is demonstration of dose-response relationship, in order to...
confirm that a lack of difference in the clinical endpoint is truly because test and reference formulations generate a similar level of effect, and not because the study lacks sensitivity in detecting differences [13]. For salmeterol, the dose-response relationship is flat when assessment is done using either bronchodilation or bronchospasm models [14-17]. Hence, though different doses of salmeterol show marked differences in Area under the Concentration-Time Curve (AUC) and maximum plasma concentration (Cmax) in a Pharmacokinetic (PK) study, there is no significant difference observed in an efficacy study. PK studies are therefore the most sensitive in vivo studies which provide information about the total amount of drug deposited in the lungs, pulmonary residence time, and systemic exposure; and are able to detect differences between two formulations [18,19].

There is limited published data describing the extent of first pass metabolism of inhaled salmeterol [20,21]. The differences between salmeterol given with and without charcoal blockade suggest that 28% to 36% of the systemic response to salmeterol administered from an MDI is due to drug absorbed from the gastrointestinal tract. This indicates salmeterol is subjected to only partial first pass metabolism and the swallowed fraction contributes to 28% to 36% of the total systemic bioavailability [22]. Therefore, for salmeterol, area under the plasma concentration curve from administration to last observed concentration at time t (AUC0-t) represents bioavailability from both the lung and swallowed fractions of the drug but in the presence of oral charcoal it reflects absorption from the lung only. The absorption of salmeterol in the lung is very quick (Tmax ≤ 5 min); therefore, Cmax reflects early bioavailability from the lung, as it excludes the later component of gut bioavailability from the swallowed fraction [23]. Significant differences in pharmacodynamic (PD) systemic effects (changes in heart rate, blood glucose and serum potassium) are seen between 150 µg and 300 µg and between 100 µg and 400 µg of salmeterol [20,22].

This review discusses 3 PK studies and 1 pharmacodynamic safety that were conducted to evaluate equivalence between two formulations of salmeterol.

Materials and Methods

Study design and subjects

All the four studies were conducted according to the declaration of Helsinki, International Conference on Harmonisation of Good Clinical Practice guidelines, European guideline, and relevant national laws and regulations. All subjects provided written informed consent for participation in the studies [24-26].

The PK studies were open-label, randomized, two-treatment, two-period, two-sequence, single-dose, crossover, phase 1 comparable bioavailability studies in healthy volunteers under fasting conditions. The pulmonary deposition of salmeterol was assessed, by administering activated charcoal to block the gut absorption, in 24 healthy volunteers following administration of a single dose of 200 µg (8 × 25 µg/puff) of test and reference HFA formulation of salmeterol (PK study with charcoal). The systemic exposure of salmeterol was assessed in 24 healthy volunteers following administration of a single dose of 200 µg (8 × 25 µg/puff) of test and reference HFA formulation of salmeterol (PK study without charcoal). The systemic exposure to salmeterol with a Volumatic spacer was assessed in 24 healthy volunteers following administration of a single dose of 100 µg (4 × 25 µg/ actuation) of the test and reference formulations of salmeterol (PK study with spacer). A safety PD dose-response study was conducted to evaluate the potential impact of higher PK levels on pharmacological effects in 21 healthy volunteers following administration of a single doses of 50 µg (2 × 25 µg/actuation), 150 µg (6 × 25 µg/actuation) and 300 µg (12 × 25 µg/ actuation) of the test and reference formulations of salmeterol. The PD study was a single-center, randomized, double-blind, placebo controlled, seven-way, crossover study.

The inclusion criteria for the PK and PD studies were as follows: healthy men aged between 18 and 45 years; Body Mass Index (BMI) between 18 and 25 kg/m² (18.5-30 kg/m² for PK study with spacer; 18-26 kg/m² for PD study); FEV1 >80% of that predicted by the European Community for Coal and Steel (ECCS) formulae (confirmed by spirometry test); no medical history of significant diseases; normal findings as determined by laboratory parameters, physical examination, and vital signs; negative result for breath alcohol test and test for drugs of abuse; and able to perform the inhalation technique correctly.

Subjects were excluded from the PK and PD studies if they had known history of hypersensitivity to salmeterol xinafoate, any component of the product, or related class of drug; had history of chronic bronchitis, emphysema, asthma or any other lung disease of clinical significance; had recent upper or lower respiratory tract infection; had consumed drugs that induce/inhibit the hepatic microsomal enzymes two months prior to dosing; and had ingested any herbal product, prescribed or non-prescribed drug four weeks prior to dosing and throughout the study.

Study products

In all the PK and PD studies, subjects fasted overnight for at least 10 h. The study inhalers were shaken well and primed before using for the first time. In the PK study with spacer, the spacers were also primed from inside to reduce the static charges.

In the PK studies with and without charcoal, subjects self-administered single dose of either test or reference product (salmeterol xinafoate HFA 25 µg/actuation × 8 puffs) as per the randomization sequence in both the periods, with a gap of at least 30 s between each puff. Additionally, in the PK study with charcoal, 50 ml (approximately 5 g) activated charcoal suspension was given 2 min prior to the first puff and 2 min after last puff, followed by 100 ml (approximately 10 g) activated charcoal suspension at 1, 2, and 3 h post dose, according to the methods of Borgstrom and Nilsson and validated for salmeterol by Bennett et al. [22]. In the PK study with spacer, after an overnight fasting of at least 10 h, subjects self-administered a single dose of either test or reference product (salmeterol xinafoate HFA 25 µg/actuation × 4 puffs) with the help of a Volumatic spacer. Deep breathing technique was used for inhalation through the spacer device as it has a lesser variability as compared to tidal breathing method [27].

In the PD study, subjects were randomized to receive salmeterol xinafoate HFA and placebo on 7 separate days as follows: test product (2 puffs=50 µg)+reference placebo, test product (6 puffs=150 µg)+reference placebo, test product (12 puffs=300 µg)+reference placebo, test placebo+reference product (2 puffs=50 µg), test placebo+reference product (6 puffs=150 µg), test placebo+reference product (12 puffs=300 µg), and test placebo+reference placebo.

Pharmacokinetic analysis

In all the three PK studies, the pre-dose samples were taken within 1 h prior to dosing; and salmeterol plasma levels were analyzed over a 24 h period. At each time point, 5 mL of blood sample was collected in a test tube containing an anticoagulant.

For the PK studies, the primary PK variables were Cmax and AUC0ₜ. Analysis of plasma samples for concentrations of salmeterol was done...
using a validated liquid chromatography-mass spectrometry (LC MS)/MS method. The Lower Limit of Quantification (LLOQ) of this method for the estimation of salmeterol concentrations in plasma was 5 pg/mL in the PK studies with and without charcoal and 2 pg/mL in the PK study with spacer. A non-compartmental method was used to calculate the PK parameters using drug concentrations versus time profile.

**Pharmacodynamic analysis**

Approximately 12 h after dosing, subjects blood sample collection and safety evaluations were carried out. Heart rates were measured by pulse oximeter for the first 9 min after dosing at an interval of 1 min. Heart rate, QTc interval, blood pressure and venous sampling for assay of serum potassium and plasma glucose was assessed at 10 min, 20 min, 30 min, 40 min, 60 min, 90 min, 2 h, 3 h, 4 h, 6 h, 8 h, 10 h, and 12 h post dose. Tremor assessment was done at 30 min pre dose and at 30 min, 1, 2, 4, and 6 h post dose.

The primary endpoints were Maximum Heart Rate (MxHR) over 12 h and Minimum Serum Potassium level (MnSP) over 4 h at the lower dose of 50 μg. The secondary endpoints were MxHR (0-12 h) and MnSP (0-4 h) at the higher doses (150 μg and 300 μg); and maximum systolic blood pressure (0-12 h), minimum diastolic blood pressure (0-12 h), maximum QTc interval (0-12 h), and maximum plasma glucose levels (0-12 h) for all doses.

**Safety analysis**

For all the four pharmacology studies, safety was evaluated by monitoring of Adverse Events (AEs), Serious Adverse Events (SAEs), physical examination, blood pressure, pulse rate, serum potassium, blood glucose, tremor assessment, well-being assessment, and ECG.

**Statistical analysis**

**Pharmacokinetic studies:** All the PK studies were designed with a sample size of 24 subjects to obtain minimum of 12 evaluable subjects to obtain reasonable data for appropriate comparison.

Analysis of covariance (ANOVA), two one-sided tests for relative bioavailability of test and reference products was compared for Cmax and AUC0-t, except in the PK study with charcoal (being the first study), and in which considering the rapid absorption of salmeterol from the lung vascular bed, a slightly wider bioequivalence limit of 75% to 133% was proposed for Cmax.

**Pharmacodynamic study:** In the PD study, assuming no difference between the two treatments, 14 subjects were considered sufficient to allow equivalence to be detected if the 95% Confidence Interval (CI) for the mean difference between the two treatments is within ± 10 bpm for heart rate and ± 0.33 mmol/L for serum potassium concentration. Accordingly to account for the dropouts, 21 subjects were enrolled in the study.

The equality or proximity of heart rate and serum potassium level at baseline over periods was analyzed using ANOVA. The treatment difference was calculated from Least Square Mean (LSM), and the 95% CIs were determined using the estimate option in mixed procedure in SAS Software.

In addition to the above analysis, additional relative potency analysis was also conducted for the primary endpoints in the PD study. The relative dose potency of the two salmeterol HFA formulations was calculated as the horizontal distance between parallel lines approximating the dose-response curves. The statistical analysis of the data included an ANOVA associated with the statistical model containing the factors: sequence, subject nested within subject, formulation, dose, formulation x dose interaction, subject (sequence) x dose interaction as factors in the model. Using the ANOVA model, the linear slope of the log dose-response curve was estimated for both the formulations. 95% CIs for the relative potency were computed according to Fieller’s theorem. Equivalence was to be concluded if the 90% CI calculated for the log (relative potency) was within 0.67 - 1.50 [28].

**Results**

**Study population**

Table 1 shows the baseline demographics of the study populations. In the PK studies with charcoal and spacer, all 24 subjects were randomized and analyzed for final bioequivalence analysis. In the PK study without charcoal, 24 subjects were randomized, and 20 were analyzed for final bioequivalence analysis (4 subjects were withdrawn due to inadequate inhalation technique). In the PD Study, 21 were randomized, and 16 completed the study and were analyzed for primary PD analysis. In the PD study, 2 subjects dropped out due to personal reason, 2 due to AEs and 1 due to inadequate inhalation technique. The baseline characteristic of the healthy volunteers across the PK and PD studies were similar which thereby enables cross comparison between the studies.

**Comparison of systemic exposure between two formulations of salmeterol (upper 90% CI limit of the difference was 136%; Table 2; Figure 1):** The 90% CI of the difference for AUC0-12h was within the bioequivalence limits of 80% to 125%; however, the Cmax of the test formulation marginally exceeded than that of the reference formulation (upper 90% CI limit of the difference was 136%; Table 2; Figure 1). The marginally higher levels in Cmax with the test formulation had no impact on the PD variables such as heart rate, serum potassium, blood glucose, and QTc interval as compared to the reference formulation when evaluated at predefined intervals over 24 h in the same study.

**Comparison of pulmonary deposition between two formulations of salmeterol (PK study with charcoal):** The 90% CI of the difference for Cmax was within the bioequivalence limits of 80% to 125%; however, AUC0-12h of the test formulation was marginally exceeded than that of the reference formulation (upper 90% CI limit of the difference was 128%; Table 2, Figure 2). The difference in AUC observed for the test

<table>
<thead>
<tr>
<th>Parameters</th>
<th>PK Study with charcoal (N=24)</th>
<th>PK study without charcoal (N=20)</th>
<th>PK study with spacer (N=24)</th>
<th>PD Study (N=21)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>26.42 ± 4.12</td>
<td>25.90 ± 6.69</td>
<td>25.96 ± 5.24</td>
<td>26.7 ± 4.1</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>63.37 ± 8.30</td>
<td>61.92 ± 7.43</td>
<td>62.15 ± 6.67</td>
<td>60.81 ± 6.66</td>
</tr>
<tr>
<td>Height (cm)</td>
<td>169.39 ± 7.28</td>
<td>168.53 ± 6.07</td>
<td>167.75 ± 5.43</td>
<td>166.8 ± 5.4</td>
</tr>
<tr>
<td>BMI (kg/m2)</td>
<td>22.06 ± 2.28</td>
<td>21.76 ± 2.22</td>
<td>22.06 ± 1.76</td>
<td>21.84 ± 1.95</td>
</tr>
</tbody>
</table>

Note: All values are in mean ± SD.

**Table 1:** Demographics characteristics
product is unlikely to result in differential pharmacological effects, since the systemic effects of salmeterol (such as increase in heart rate, serum potassium) is likely to correlate with peak concentrations and early absorption due to rapid appearance of salmeterol in blood. The marginally higher levels in AUC with the test formulation had no impact on the PD variables such as heart rate, serum potassium, blood glucose, and QTc interval as compared to the reference formulation evaluated at predefined intervals over 24 h in the same study.

Comparability in systemic exposure with spacer device using two formulations of salmeterol: The 90% CIs of the difference for both AUC0-t and Cmax were within the bioequivalence limits of 80% to 125% (Table 2 and Figure 3) with the volumatic spacing device. The test formulation showed an increase of around 40% and the reference formulation showed an increase of around 54% with the spacer device in peak plasma concentration as compared to the conventional actuator. The increase in lung deposition with the spacer device is consistent with the published studies with Volumatic spacer device [29,30].

Comparison of systemic PD effects of the two formulations of salmeterol: At each of the doses, the 95% CI for the treatment difference was within the clinically relevant limits of ±10 bpm for increase in heart rate and ±0.33 nmol/L for decrease in serum potassium (Table 3; Figures 4 and 5). The treatments were also comparable for the other safety variables such as blood glucose and QTc interval.

The relative potency was also calculated for heart rate and serum potassium using 150 µg and 300 µg doses as these doses were on the steep part of the dose response curve. The 90% CI for the relative potency was well within limits of 0.67-1.5 for MxHR (0-12 h) and MnSP (0-4 h) (Table 4).

### Table 2: Pharmacokinetic parameters.

<table>
<thead>
<tr>
<th>PK Parameters</th>
<th>Test (T)</th>
<th>Reference (R)</th>
<th>% (T/R)*</th>
<th>90% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>PK Study with charcoal (N=24)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AUC0-t (pg·h/mL)</td>
<td>1230.16</td>
<td>1088.09</td>
<td>113.06</td>
<td>99.58-128.35</td>
</tr>
<tr>
<td>Cmax (pg/mL)</td>
<td>891.01</td>
<td>807.07</td>
<td>110.40</td>
<td>99.33-122.71</td>
</tr>
<tr>
<td>AUC0-30min (pg·h/mL)</td>
<td>223.02</td>
<td>199.36</td>
<td>111.87</td>
<td>(100.99-123.91)</td>
</tr>
<tr>
<td>Tmax (h)</td>
<td>0.05 (0.05-0.08)</td>
<td>0.05 (0.05-0.08)</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

| PK study without charcoal (N=20) |                |               |          |                 |
| AUC0-t (pg·h/mL)      | 1502.27        | 1453.13       | 103.38   | 92.07-116.09    |
| Cmax (pg/mL)          | 972.98         | 880.01        | 110.56   | 89.70-136.29    |
| AUC0-30min (pg·h/mL)  | 222.18         | 214.61        | 103.53   | 85.13-125.90    |
| Tmax (h)              | 0.05 (0.03-0.08) | 0.05 (0.03-0.08) | -      | -               |

| PK study with spacer (N=24) |                |               |          |                 |
| AUC0-t (pg·h/mL)      | 1271.42        | 1402.13       | 90.68    | 83.87-98.03     |
| Cmax (pg/mL)          | 1391.64        | 1471.96       | 94.54    | 87.11-102.61    |
| AUC0-30min (pg·h/mL)  | 327.76         | 355.84        | 92.11    | 85.50-99.23     |
| Tmax (h)              | 0.05 (0.03-0.12) | 0.05 (0.03-0.12) | -      | -               |

All values are in geometric mean except Tmax which is in median (range). *(%) T/R is ratio of TestGeoLSM/RefGeoLSM.
was within 5 min which indicates that all subjects had performed the studies. The pulmonary absorption peak observed in all PK studies prior to the start of the study with an in-check dial, placebo more appropriately.

The PK data presented in this paper was 5 to 12 times below what has been used in published studies which enabled complete characterization of the PK profile till 24 h even with 200 µg and 100 µg doses [20,31].

It is known that pMDIs have no flow rate dependency; therefore, healthy subjects were selected for the PK and PD studies. In addition, healthy cohort is the most sensitive cohort to detect differences arising due to the device/formulation that are not confounded by factors (e.g. airway caliber changes) which can cause PK/PD differences that are independent of differences in the device/formulation.

The salmeterol concentration reported in the PK study without charcoal is consistent to findings in the published literature in healthy subjects. The C max obtained was 972 pg/mL and 880 pg/mL with 200 µg of the test and reference formulations, respectively, which is consistent with 700.8 pg/mL which was obtained with 150 µg of salmeterol. The AUC values reported with 200 µg doses of the test and reference formulations are 1502.27 and 1453.13 pg/mL.h, respectively, which are far greater than 558.9 pg/mL.h reported with 150 µg of salmeterol as we are able to characterize the complete plasma profile of salmeterol at the dose used in the PK studies [20]. As expected, higher AUC levels were observed with spacer for both the formulations.

The differences between salmeterol given with and without charcoal indicate that approximately 25% of the systemic response to salmeterol administration is due to drug absorbed from the gastrointestinal tract. This is in accordance with the study by Bennett and Tattersfield wherein 28% to 36% of salmeterol systemic response was due to drug absorbed from the gastrointestinal tract [22]. The exponential decline, single plasma peaks within 5 min post administration, and lack of additional secondary peaks indicates an effective blockage of gastrointestinal absorption by charcoal. Although the with and without charcoal comparisons have been derived from two separate studies, both the PK studies were conducted under the same standardized set up such as same dose, same centre, same design, similar sampling duration, and similar washout periods; subjects studied in both studies were healthy and had similar demographic profile.

The PK studies demonstrated that the test formulation had a marginally higher plasma concentration compared with the reference formulation. However, it should be noted that no significant difference was seen in systemic β2-mediated adverse effects (maximum heart rate and maximal fall in potassium); even at 6 times the recommended single dose in the assay-sensitive PD study. Likewise, the test formulation had no pronounced effects on other β2 agonist PD effects such as QTc interval, maximum systolic blood pressure, and maximum plasma glucose concentration. Considering that the systemic exposure of the test product was marginally greater in the PK studies, the PD study was therefore a good alternative clinical study model to evaluate impact of greater systemic exposure and additionally evaluate relative potency by satisfying the requirement of the OIP guideline in terms of assay sensitivity.

Dose-response curves of the systemic drug effects are equally important for establishing full therapeutic equivalence. The ratio of doses giving equivalent effects (potency ratio) indicates relative potency of two formulations [32,33]. In a previous study, three doses of formoterol (24, 48, and 96 µg) and salmeterol (100, 200 and 400 µg)
were evaluated to estimate relative potency between the two drugs [33]. Relative potency was adequately estimated in this study as both drugs had dose-related effects on the systemic parameters. The relative dose potency for systemic effect such as maximum heart rate was 4.1 (95% CI: 3.0 to 5.6) over 4 h. This indicates that formoterol is 4.1 times more potent than salmeterol. It is interesting to note that the 4-fold difference in the recommended doses of formoterol and salmeterol (12 μg formoterol vs. 50 μg salmeterol twice daily) are similar in their relative dose potency ratio of 1:4 for systemic endpoints. Therefore, in absence of a sensitive efficacy clinical model relative potency of a product can be adequately evaluated through a PD safety model.

The PD model utilized in the PD safety study presented in this paper is a well-established model used by Bennett and Tattersfield and subsequently elaborated by Guhan et al. [33] and satisfies the requirement of assay sensitivity. Therefore, the PD study reconfirms that the test product was equipotent to the reference product.

The data presented in this paper raise some interesting points with respect to the procedure for establishing therapeutic equivalence between generic and innovator products. A product may be acceptable to patients and clinicians if there is no clinical difference between the two formulations, even if bioequivalence is not established. Therefore it is important to evaluate difference in PD data in conjunction with PD data to understand the relevance of its impact on the efficacy and safety.

### Conclusion

Overall, it can be concluded that the test HFA formulation of salmeterol is therapeutically equivalent to the reference HFA formulation of salmeterol when used with and without the spacer device. Both test and reference products were safe and well tolerated.

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### Conflict of Interest

The authors have indicated that they have no other conflicts of interest regarding the content of the article.

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15. GSK Study SLGH05 (2005) GSK data.


