Effect of Fluconazole on Indinavir Pharmacokinetics in Human Immunodeficiency Virus-Infected Patients

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To evaluate a potential pharmacokinetic interaction of coadministration of fluconazole and indinavir, a human immunodeficiency virus (HIV) protease inhibitor, 13 patients were enrolled in a multiple-dose, three-period, placebo-controlled, crossover study. Patients were randomly assigned to receive indinavir at 1,000 mg every 8 h for 7 1/2 days (with fluconazole placebo), fluconazole at 400 mg once daily for 8 days (with indinavir placebo), and indinavir with fluconazole in combination. The pharmacokinetics of both drugs were measured on day 8 of each treatment period. The peak concentration in plasma (Cmax) and the time to reach Cmax were obtained by inspection, and the area under curve (AUC) was calculated for indinavir and fluconazole for each treatment period in which the respective drugs were administered. There was a marginally (P = 0.08) statistically significant decrease in the AUC from 0 to 8 h (AUC0–8) for indinavir when it was administered with fluconazole. However, the magnitudes of the decreases in Cmax and the concentration at 8 h postdosing (C8) were not as great as the decrease in AUC0–8. Although the 90% confidence interval for the geometric mean ratio was within the hypothesized limits, the clinical significance is not clear. Indinavir coadministration with fluconazole had no statistically (P > 0.5) or clinically significant effect on the Cmax and C8 of indinavir. Fluconazole coadministration with indinavir had no statistically or clinically significant effect on the pharmacokinetics of fluconazole. One patient was discontinued because of mild to moderate abdominal pain and diarrhea while on indinavir and fluconazole in combination. No serious adverse experience according to the results of laboratory tests was noted. Total bilirubin levels in serum were mildly increased in most patients treated with indinavir. This was not clinically significant and was not affected by the coadministration of fluconazole. Although the values of the pharmacokinetic parameters for indinavir decrease in the presence of fluconazole, indinavir and fluconazole can be administered concomitantly to HIV-infected patients without adjustment of the dose of either drug, and both drugs are generally well tolerated.

Human immunodeficiency virus (HIV) protease inhibitors are a new class of antiretroviral drugs with a high in vitro antiviral potency and a favorable toxicity profile. Recent clinical trials have demonstrated very promising results in terms of viral load, CD4-cell count, morbidity, and mortality, particularly in patients with advanced HIV infection.

Although these agents demonstrate antiretroviral effects exceeding those of the nucleoside analogs, as measured by CD4-cell count and plasma viremia levels, their long-term clinical benefit and their usefulness in patients with higher CD4-cell counts remain to be determined. Nevertheless, HIV protease inhibitors are increasingly used in the treatment of HIV-infected patients, particularly in those with advanced HIV disease and low CD4-cell counts. These patients are generally exposed to multiple drug interventions including other antiretroviral agents, drugs used as primary prophylaxis for and treatment of opportunistic infections, cancer chemotherapy, and symptomatic therapies.

Indinavir is an orally bioavailable protease inhibitor that has shown a significant antiviral in vivo effect alone at the indicated dosage of 800 mg every 8 h (q8h) and in combination with zidovudine and lamivudine, with a 1- to 2-log reduction in the number of HIV type 1 (HIV-1) RNA copies in plasma and increases in the CD4-cell number of up to 80 to 140 after 48 weeks of treatment (5).

Fluconazole is the most widely used azole antifungal compound in HIV-infected patients. This implies that coadministration of indinavir and fluconazole could be a frequent situation in the clinical setting.

The major role of the P-450 isozyme CYP3A4 in the metabolism of indinavir suggests that coadministration with drugs such as fluconazole which inhibit P-450 enzymes might alter the pharmacokinetics of indinavir (1, 2, 7). The objectives of this study were to determine the effect of coadministration of fluconazole and indinavir on the pharmacokinetic profile of indinavir in plasma and to evaluate the safety and tolerability of coadministration. The effect of indinavir on the pharmacokinetic profile of fluconazole in plasma was also assessed.

MATERIALS AND METHODS

Patients. HIV-positive patients of both sexes (provided that, for females, a serum pregnancy test was negative and barrier contraception was used) between the ages of 18 and 60 years, with a CD4-cell count of greater than 50 cells/mm3, and with no active AIDS-defining opportunistic infection were evaluated for the trial, provided that their weight was above 45.5 kg.

They were excluded in the case of a history of hepatic disease, a positive test for hepatitis B virus surface antigen or hepatitis C virus antibodies, or any elevation of serum aspartate aminotransferase, alanine aminotransferase, or bilirubin levels during the previous 3 months or a history of threefold or greater elevations in the levels of these components in the past. Other biological exclusion criteria included a serum creatinine level above 1.5 mg/dl, a granulocyte count below 1,000/mm3, or a hemoglobin level below 9.5 g/dl. No concomitant medication with the exception of acetaminophen (paracetamol) and prophylaxis (223-227 Vol. 42, No. 2 Antimicrobial Agents and Chemotherapy, Feb. 1998 © 1998 American Society for Microbiology)
with co-trimoxazole, aerosolized pentamidine, or topical antifungals agents was allowed. Daily intakes of greater than six 12-oz. portions of caffeine-containing beverages, more than two drinks of alcohol (wine, beer, or spirits), or more than 20 cigarettes were prohibited. This protocol was approved by the Ethical Review Committee of Centre Hospitalier Universitaire Saint-Pierre, and informed consent was obtained.

Study design and procedures. The trial was designed as a multiple-dose, randomized, three-period, crossover study. Patients were randomly assigned to treatment sequences according to a two-balanced 3 by 3 Latin Square design consisting of active indinavir with fluconazole placebo (treatment A), indinavir placebo with active fluconazole (treatment B), and active indinavir with active fluconazole (treatment C). Thus, data for six sequences of treatment were obtained: DBC, DCA, BAC, and CBA. Indinavir was administered at a dosage of 1,000 mg qd for 7½ days on an empty stomach (2 h following or 1 h prior to a meal). At the time that the trial was designed, this was the highest dose shown to be well tolerated. Fluconazole was administered at a dosage of 400 mg once daily (qd) for 8 days. There was at least a 7-day washout period between each final dose of one treatment and the first dose of the subsequent treatment. Blood and urine for laboratory tests for drug safety were obtained prior to the administration of dose 1 and 4 h following the administration of the final dose of each treatment (day 8). Additionally, liver function tests (total and direct bilirubin, aspartate aminotransferase, alanine aminotransferase, and alkaline phosphatase levels) were determined on days 3 and 5 of each treatment.

Physical examinations were performed prior to the administration of the first dose of each treatment (day 1) and between 0 and 8 h following the administration of the final dose of each treatment (day 8). Electrocardiograms were performed prior to the administration of the first dose (day 1) of the first treatment, as well as 1 h following the administration of the final dose of each treatment (day 8). Vital signs were measured and frequently scheduled times on the first and last day of dosing. Patients were asked to maintain a diary on a card to record the time of administration of all doses and to note any adverse experiences during each treatment. A poststudy evaluation for safety consisting of laboratory tests, physical examination, and electrocardiogram was performed 24 h following the final treatment.

Pharmacokinetics. Blood was drawn for determination of the concentration of indinavir in plasma at 0, 0.5, 1, 1.5, 2, 3, 4, 6, and 8 h following the administration of the final dose of each treatment (day 8). Blood was drawn for determination of the concentration of fluconazole in plasma prior to the administration of the morning dose on days 3, 5, 6, 7, and 8 during each treatment as well as following the administration of the final dose of each treatment (day 8) at 1, 2, 4, 8, 12, 24, and 48 h postdosing. The blood drawing schedule was identical during each treatment to preserve the binding of the sequence of treatments, but plasma was actually analyzed for indinavir and/or fluconazole levels only for the treatments when the active drug was administered. On day 8 urine was collected from 0 to 8 h for assay of indinavir levels. A blood sample and a urine sample as blanks for assays for indinavir levels were obtained prior to administration of the first dose of the first treatment only.

The concentrations of indinavir in plasma were determined by the high-pressure liquid chromatography technique (10). We used a modified version of the high-pressure liquid chromatography assay reference procedure in which the assay limit of quantitation was 25 ng/ml (40.7 nM) instead of the originally described limit of 5 ng/ml (8 nM). The range of the linear standard curve was 25 to 5,000 ng/ml. The interday coefficient of variation was 6% for 75 ng/ml and described limit of 5 ng/ml (8 nM). The range of the linear standard curve was 25 to 5,000 ng/ml and 6.8% for 55.28 g/ml.

The peak concentration in plasma (Cmax) and the time to reach Cmax (tmax) were obtained by inspection, and the areas under the curve (AUCs) were calculated for indinavir and fluconazole for each treatment in which the respective drugs were administered. AUCs for both indinavir and fluconazole were calculated over the interval from 0 to 8 h (AUC0–8) by the modified trapezoidal rule. The areas under the curve for combination therapy were calculated by using the taking of the mean for each observed value. Plasma was sampled using the modified trapezoidal rule and geometric mean ratios with 90% confidence interval of 0.50 to 2.0.

RESULTS

Patients. Thirteen HIV-seropositive patients participated in the study. The demographic characteristics of the group are as follows. Of the 13 patients 11 were males, 2 were females, 11 were Caucasian, and 2 were black. The mean ± standard deviation (SD) age was 39 ± 10.4 years. The mean ± SD weight was 74.1 ± 10.7 kg. Two patients were discontinued from the study due to clinical adverse experiences. One patient was discontinued from the study while receiving both active drugs and was not replaced. One patient was discontinued from the study on day 2 and was replaced. Data for all 11 patients who completed the study are included in the pharmacokinetic analysis. Data for all 13 patients are included in the safety analysis.

Pharmacokinetics. (i) Effect of fluconazole on indinavir. The geometric means, geometric mean ratios, and 90% confidence intervals for AUC0–8, Cmax, and C0 for indinavir administered alone and in combination with fluconazole are summarized in Table 1. Figures 1, 2, and 3 illustrate the individual AUCs, Cmax, and geometric mean ratios with 90% confidence intervals for indinavir, respectively. Hypothesis 1 states that the AUC0–8, Cmax, and C0 of indinavir after 1 week of coadministration with fluconazole would not be substantially altered compared to those observed after 1 week of coadministration with placebo (geometric mean ratios of values for

<table>
<thead>
<tr>
<th>Parameter</th>
<th>AUC0–8 (nM h)</th>
<th>Cmax (nM)</th>
<th>C0 (nM)</th>
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</thead>
<tbody>
<tr>
<td>Geometric mean without fluconazole</td>
<td>39.1423</td>
<td>17.2163</td>
<td>235.2</td>
</tr>
<tr>
<td>Geometric mean with fluconazole</td>
<td>29.8314</td>
<td>15.0185</td>
<td>210.6</td>
</tr>
<tr>
<td>Geometric mean ratio for combination therapy/monotherapy</td>
<td>0.76</td>
<td>0.87</td>
<td>0.90</td>
</tr>
</tbody>
</table>

90% Confidence interval 0.59–0.98 0.72–1.05 0.72–1.12

*Results are based on data for 11 subjects.*
combination therapy to those for monotherapy, no less than 0.50 and no more than 2.0). The geometric mean AUC_{0–8} values for indinavir administered alone and indinavir administered in combination with fluconazole were 39,142.3 and 29,831.4 nM · h, respectively (Table 1; Fig. 1). The geometric mean ratio of the values with combination therapy to the values with monotherapy was 0.76. This ratio was marginally statistically significant different from 1.0 (P = 0.08). The geometric mean AUC_{0–8} for indinavir did not differ by more than 41% between combination therapy and monotherapy. This is shown by the 90% confidence interval for the geometric mean ratio of 0.59 to 0.98. This was within the hypothesized interval of 0.50 to 2.0. The geometric mean C_{max} of indinavir alone and in combination with fluconazole were 17,216.3 and 15,018.5 nM, respectively. The geometric mean ratio of the C_{max} with combination therapy to the C_{max} with monotherapy was 0.87. The 90% confidence interval for the geometric mean ratio is 0.72 to 1.05 (Table 1; Fig. 2). This was within the interval of 0.50 to 2.0. The geometric mean C_{max} of indinavir administered alone and indinavir administered in combination with fluconazole were 235.2 and 210.6 nM, respectively (Table 1; Fig. 3). The geometric mean ratio of the C_{g} with therapy combination to that with monotherapy was 0.90. The geometric mean C_{g} of indinavir did not differ by more than 28% between combination therapy and monotherapy. This is shown by the 90% confidence interval for the geometric mean ratio of 0.72 to 1.12. This was within the hypothesized interval of 0.50 to 2.0. The individual C_{g} of indinavir when it was coadministered with fluconazole ranged from 92.1 to 332.8 nM. The range is within the 95% confidence interval of 25 to 100 nM. Thus, there was a marginally statistically significant decrease in the AUC_{0–8} for indinavir when it was administered with fluconazole. Although the 90% confidence interval for the geometric mean ratio was within the hypothesized limits, the clinical significance is not clear. There was no statistical or clinical significant effect of coadministration of indinavir with fluconazole on C_{max} or C_{g}. In summary, there is no evidence of a clinically significant pharmacokinetic effect of the coadministration of indinavir and fluconazole on indinavir.

(ii) Effect of indinavir on fluconazole. The pharmacokinetic parameters analyzed for fluconazole, which was administered at a dosage of 400 mg q.d. included AUC_{0–8} and C_{0} (or the trough concentration 24 h after dosing on day 7). There were two limitations to the study design. First, in order to evaluate the effect of indinavir on fluconazole over the 24-h dosing interval, all three indinavir doses on day 8 must have been administered; this was not required according to the protocol. Therefore, the AUC_{0–8} and C_{0} rather than AUC_{0–24} and C_{24} were calculated on day 8 for fluconazole. Second, fluconazole was administered for 8 days. However, trough concentrations indicated that the fluconazole concentrations did not reach steady state on day 8. The geometric mean AUC_{0–8} and C_{0} for fluconazole administered alone and in combination with indinavir are summarized in Table 2. The C_{max} and T_{max} for fluconazole were calculated but were not analyzed statistically. The T_{max} values for fluconazole administered alone ranged from 0 to 24 h, and the T_{max} values for fluconazole coadministered with indinavir ranged from 1 to 12 h. The arithmetic mean C_{max} values for fluconazole administered alone and coadministered with indinavir were 20.79 and 20.17 μg/ml, respectively. The half-life of fluconazole is long, and thus concentrations in plasma are relatively constant throughout the

<table>
<thead>
<tr>
<th>Parameter</th>
<th>AUC_{0–8} (μg · h/ml)</th>
<th>C_{0} (μg/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Geometric mean without indinavir</td>
<td>145.3</td>
<td>13.5</td>
</tr>
<tr>
<td>Geometric mean with indinavir</td>
<td>146.0</td>
<td>13.5</td>
</tr>
<tr>
<td>Geometric mean for combination therapy/monotherapy</td>
<td>1.00</td>
<td>1.00</td>
</tr>
<tr>
<td>90% Confidence interval</td>
<td>0.96–1.05</td>
<td>0.84–1.20</td>
</tr>
</tbody>
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* Results are based on data for 11 subjects.
fluconazole dosing interval on day 8. For a drug with this type of concentration profile in plasma (with no clear peak) after the administration of multiple doses, the parameters \(T_{\text{max}}\) and \(C_{\text{max}}\) are of limited value. The geometric mean \(\text{AUC}_{0-8}\) values for fluconazole administered alone and in combination with indinavir were 145.3 and 146.0 µg · h/ml, respectively. The geometric mean ratio of the \(\text{AUC}_{0-8}\) with combination therapy to that with monotherapy was 1.00 (Table 2; Fig. 4). The geometric mean \(\text{AUC}_{0-8}\) for fluconazole did not differ by more than 5% between combination therapy and monotherapy. This is shown by the 90% confidence interval for the geometric mean ratio of 0.96 to 1.05. This was within the interval of 0.50 to 2.0. The geometric mean \(C_{\text{ss}}\) of fluconazole administered alone and in combination with indinavir were 13.5 and 13.5 µg/ml, respectively (Table 2; Fig. 5). The geometric mean ratio of the \(C_{\text{ss}}\) with combination therapy to the \(C_{\text{ss}}\) with monotherapy was 1.00. The geometric mean \(C_{\text{ss}}\) of fluconazole did not differ by more than 20% between combination therapy and monotherapy. This is shown by the 90% confidence interval for the geometric mean ratio of 0.84 to 1.20. This was within the interval of 0.50 to 2.0. Collectively, the pharmacokinetic data indicate that the coadministration of fluconazole with indinavir had no statistically or clinically significant effect on the pharmacokinetics of fluconazole.

**Safety.** Data for all 13 patients were included in the safety evaluation. Thirteen patients had clinical adverse experiences. The most common clinical adverse experiences were diarrhea, nausea, abdominal pain, acid regurgitation, headache, taste disturbances, hot flushes, and eye accommodation disorder during treatment with either indinavir alone, indinavir and fluconazole, or fluconazole alone. Eight patients had adverse experiences which were judged to be related to one of the study drugs. Among patients receiving the combination of indinavir and fluconazole, one patient had taste disturbance and one patient had taste loss, and another patient receiving indinavir alone reported taste loss. These were judged to be possibly related to study drugs. Four patients experienced nausea which was judged to be possibly drug related. One patient receiving indinavir alone and one patient receiving fluconazole alone had nausea. One patient receiving indinavir and fluconazole in combination had nausea. Two patients receiving indinavir and fluconazole in combination while not receiving a study drug had nausea. All of these adverse experiences were mild or moderate.

Two patients were discontinued from the study due to clinical adverse experiences. One patient had a serious clinical adverse experience judged to be definitely not related to study drug. He received indinavir at 1,000 mg q8h plus fluconazole at 400 mg on day 1 while complaining of cough and sputum production. On day 2 he was hospitalized for a lung disorder and was discontinued from the study while off of drug. The other patient experienced nausea, taste disturbance, hot flushes, abdominal pain, and diarrhea while receiving both indinavir at 1,000 mg q8h and fluconazole at 400 mg q.d. He was discontinued on day 5 of the second treatment while receiving indinavir alone. These clinical adverse experiences were judged to be possibly related to study drug.

All 13 patients were included in the safety evaluation performed with data from laboratory tests. Three of 13 patients had adverse experiences according to the results of laboratory tests. One patient had pyuria on day 8 following the administration of fluconazole at 400 mg q.d. and on day 26 while off of either study drug. One patient had pyuria predosing on day 1 and had activated PT increased on day 25 following the administration of indinavir at 1,000 mg q8h and fluconazole at 400 q.d. The third patient had hyperkalemia on day 8 following the administration of indinavir at 1,000 mg q8h and fluconazole at 400 mg q.d. No patient had an adverse experience that was judged on the basis of laboratory test data to be related to study drug. No adverse experience was serious on the basis of laboratory test data. No patient was discontinued from the study due to an adverse experience on the basis of laboratory test data. Total bilirubin levels in serum were increased in some patients. The highest bilirubin level was 1.9 mg/dl. No clinically significant deviations in the evaluation of the data from tests for clinical safety such as physical examination, vital signs, or electrocardiogram were found.

**DISCUSSION**

Multiple drugs are commonly prescribed to HIV-infected patients, and the potential for drug interactions leading to either adverse events or reduced efficacy is great. It is thus particularly important to carefully evaluate potential interactions between drugs whose metabolic pathways are similar and which will commonly be administered concomitantly. This is the case for HIV protease inhibitors andazole derivatives, in particular, indinavir and fluconazole, whose coadministration could lead to increased plasma indinavir levels through inhibition of the CYP3A4 isozyme.

Several dosing regimens for indinavir have been investigated to date. At the time that this study was designed a progressive dose-response from 200 mg every 6 h (q6h) (0.8 g/day) to 600...
mg q6h (2.4 g/day) had been demonstrated. In view of a potential risk of the development of resistance with dosages of less than 2.4 g/day, a study aimed at determining the maximum antiretroviral response possible with indinavir monotherapy was initiated. Three dosage regimens were investigated: 800 mg q8h (2.4 g/day), 1,000 mg q8h (3.0 g/day), and 800 mg q6h (3.2 g/day). At the time that the indinavir-fluconazole study started, the dosage of 1,000 mg q8h had been shown to be well tolerated and was selected for use in the trial. Following completion of the comparative trial with the three dosage regimens, the lower dosage (800 mg q8h) was selected for clinical development. Indinavir dosages above 2.4 g/day did not appear to exert a greater antiviral effect (8). The dosage of fluconazole chosen for the study, 400 mg q.d., is the highest dosage indicated for the treatment of fungal infections.

There was a decrease in the AUC0–8, Cmax, and Cmax for indinavir when it was coadministered with fluconazole. However, the magnitudes of the decreases in Cmax and C8 are not as great as the decrease observed for AUC0–8. Due to the nonlinear pharmacokinetics of indinavir, the AUC0–8 of indinavir when it is coadministered with fluconazole is anticipated to be reduced, but it is greater than the AUC0–8 seen with indinavir given at 600 mg q8h. The mechanism for the decrease in the AUC0–8 could be due to induction of P-450 metabolism by fluconazole. Fluconazole appears to induce some mammalian P-450 enzymes, and thus, in theory, fluconazole has the potential to increase as well as decrease the clearance of P-450-metabolized drugs (3).

However, the effects of fluconazole on the pharmacokinetics of other drugs (such as phenytoin, cyclosporine, and anticoagulants) appear to be mediated predominantly by P-450 inhibition, not P-450 induction. In this study, no increase in the AUC0–8 for indinavir consistent with P-450 inhibition was observed. In contrast, coadministration of indinavir with ketoconazole does result in an increase in plasma indinavir concentrations (6). This is consistent with the greater potency of ketoconazole versus that of fluconazole as a P-450 inhibitor (7). Another possibility for the apparent effect of fluconazole on indinavir pharmacokinetics is interference with absorption, but there is no known precedent for such an effect of fluconazole. Given that the effect of fluconazole on indinavir did not quite reach statistical significance, this result could well have been a chance occurrence. In any event, the clinical significance of this small decrease in the AUC0–8 for indinavir is unclear. However, this does not warrant a dose adjustment. The AUC0–8 and C8 values for fluconazole when it was and those for fluconazole when it was administered alone in combination with indinavir did not differ significantly. This suggests that indinavir, a potential CYP3A4 inhibitor, did not affect the metabolism of fluconazole. This is consistent with the minor role of hepatic metabolism in the clearance of fluconazole. The Tmax and Cmax values for fluconazole administered alone and coadministered with indinavir are of limited value. The half-life of fluconazole is long, and thus, concentrations in plasma are relatively constant, with no clear Cmax or Tmax on day 8 (4).

A review of reports of adverse experiences and tabulated summaries of selected laboratory values indicate that indinavir and fluconazole administered alone or together were generally well tolerated. Two patients were discontinued from the study due to clinical adverse experiences. One patient had a serious clinical adverse experience, judged not to be drug related, following day 1 of treatment with indinavir plus fluconazole. He was hospitalized for a lung disorder. The other patient was discontinued from the study due to nausea, taste disturbance, hot flashes, abdominal pain, and diarrhea while receiving both indinavir and fluconazole. These clinical adverse experiences were judged to be possibly drug related. No patient had an adverse experience that was judged on the basis of data from laboratory tests to be related to one of the study drugs. No adverse experience was serious on the basis of data from laboratory tests. No patient was discontinued from the study due to an adverse experience on the basis of data from laboratory tests.

In summary, indinavir and fluconazole may be administered concomitantly to HIV-seropositive patients without adjustment of the dose of either drug. The two drugs given concurrently generally appear to be well tolerated.

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