Effect of a Fat- and Calcium-Rich Breakfast on Pharmacokinetics of Fleroxacin Administered in Single and Multiple Doses

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The effect of a fat- and liquid-calcium-rich meal on the pharmacokinetics of single and multiple doses of fleroxacin in 20 healthy men and women was investigated in a randomized crossover fashion. Fleroxacin was administered as 400 mg daily for 3 days and as a single 400 mg dose. Concurrent administration of fleroxacin with food resulted in a statistically significant (P ≤ 0.05) decrease in the area under the curve (13.9% for multiple-dose administration, 10% for single-dose administration) and in the peak concentration (25.9% for multiple-dose administration, 27% for single-dose administration) and a lengthening of the time to peak (more than doubled for single- and multiple-dose phases). In addition, by using an equivalence criteria of 80 to 125%, the two one-sided tests procedure indicated that the mean areas under the curves for fleroxacin administered in a fed and a fasted state were statistically bioequivalent (P ≤ 0.05) for both the single- and multiple-dose regimens. Although a meal high in fat and containing liquid calcium reduces the peak concentration by approximately 25%, a minimal effect on bioavailability is seen with concomitant food administration. In addition, multiple-dose bioavailability studies appear to give similar information to single-dose studies while representing the clinical setting more closely.

Fleroxacin is a new fluoroquinolone anti-infective agent which is structurally related to nalidixic acid but has a broader spectrum of activity and favorable pharmacokinetic properties. This agent is a bactericidal drug whose mechanism of action is through the inhibition of an essential bacterial enzyme, DNA gyrase, which is needed for DNA replication.

Fleroxacin has extensive activity against gram-negative bacteria. This high level of antimicrobial activity is combined with excellent penetration in the body fluids and tissues (2, 6). After oral administration of 400 mg, peak concentrations (C_{max}) of 3 to 7 µg/ml have been reported (6, 14). The elimination half-life (t_{1/2}) for fleroxacin ranges from 7.9 to 13 h, the volume of distribution (V_{d}) ranges from 1.2 to 1.7 liters/kg, and total body clearance (CL) ranges from 5.9 to 10.6 liters/h (6, 14). The drug is approximately 23% protein bound. The oral absorption of fleroxacin is characterized by nearly complete bioavailability. From 45 to 70% of the drug is renally excreted as unchanged fleroxacin.

The interaction of fluoroquinolones with food has been studied extensively. Results of pharmacokinetic studies of orally administered ciprofloxacin, lomefloxacin, enoxacin, pefloxacin, and ofloxacin have indicated variation in the pharmacokinetics of some of these agents when administered with food (4, 7, 9, 10, 18, 20). Data for ciprofloxacin suggest that administration concurrently with food containing calcium in a liquid form results in an approximately 35% decrease in the peak concentration (C_{max}) and area under the curve (AUC) (13). Similar results have been noted for norfloxacin (8). Studies conducted after administration of single oral doses of fleroxacin have shown that food does not affect the pharmacokinetic parameters of this drug in a clinically significant fashion (6, 11, 17, 21). However, no studies have been conducted after administration of multiple-dose regimens. Multiple-dose regimens mimic the manner in which antibiotics are used in the clinical setting more closely than single doses do. In addition, shorter sampling periods are needed, thus decreasing costs associated with specimen collection and assay.

The purpose of the present study was to determine the influence of a fat- and calcium-rich breakfast on pharmacokinetics of single and multiple doses of fleroxacin. The calcium source for this study was a liquid source (whole milk) which has been shown to cause a clinically and statistically significant reduction in the absorption of oral ciprofloxacin (13).

MATERIALS AND METHODS

This study was approved by the Institutional Review Board of The Mary Imogene Bassett Hospital. Written informed consent was obtained from the subjects before their enrollment in the trial. A total of 20 subjects (10 men and 10 premenopausal women) participated in the trial. This trial was an open-label, randomized, four-period crossover study of the effect of food on the rate and extent of absorption of fleroxacin given in single and multiple doses.

Before randomization, all subjects underwent an initial history and physical examination, including laboratory testing, to ascertain the presence of normal renal and hepatic function. No subject was enrolled if creatinine clearance was 30 ml/min or below as determined by the method of Cockcroft and Gault (3). Subjects were excluded if they had clinical or laboratory evidence of impairment of hepatic function (alanine or asparate transaminase or lactate dehydrogenase at or above twice the upper limit of normal range, or total bilirubin ≥ 2 mg/100 ml). Subjects were ineligible if they had known or suspected hypersensitivity to nalidixic acid, ciprofloxacin, or other quinolones; documented history of recurrent seizures or a seizure

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disorder; current diagnosis of drug or alcohol abuse or dependence; or gastrointestinal disorders that might affect drug absorption. All subjects were 18 years of age or older and within 30% of ideal body weight according to the 1983 Metropolitan Life Insurance Company tables. Women underwent pregnancy testing during the enrollment period and before initiation of treatment for each phase of the study.

On day 1 of the study, subjects were randomly assigned to the order in which they would receive four dosing regimens of fleroxacin. After the subjects received one of the four designated phases, they were allowed a minimum 7-day washout period and then crossed over to receive one of the other phases. Each subject participated in all four phases.

Fleroxacin dosing regimens were as follows: (i) fleroxacin (400 mg) once daily for 3 days (multiple dose) in the fasted state, (ii) fleroxacin (400 mg) once daily for 3 days (multiple dose) in the fed state immediately following a standard breakfast, (iii) fleroxacin (400 mg) single dose in the fasted state, and (iv) fleroxacin (400 mg) single dose in the fed state immediately following a standard breakfast. All doses were given between 0700 and 0900 h. All subjects had no food or liquid for at least 12 h before the fleroxacin dose.

The standard breakfast used during the fed phases of the study consisted of 240 ml of whole milk, two eggs scrambled in three teaspoonsful of margarine, two strips of bacon, one slice of white toast with one teaspoonful of jam and one tablespoonful of jelly, and 120 ml of orange juice. This breakfast provided 770 calories total, 58% fat (50 g), 29% carbohydrate (55 g), 13% protein (25 g), and 393 m of calcium in a liquid form. Subjects consumed the breakfast over a 15-min period, and at the completion of the breakfast the fleroxacin dose was immediately administered with 120 ml of water. The drug was given (by one of the investigators, L.S.) on each of the three study days following the standard breakfast. Volunteers did not eat for 4 h after the breakfast and/or the fleroxacin dose. No alcohol was consumed for 3 days before and during each study phase.

During the multiple-dose phases, blood samples for fleroxacin assay were collected before the first dose (baseline), before the third dose, and at 0.25, 0.5, 0.75, 1, 1.5, 2, 3, 4, 6, 8, 10, 12, 14, and 24 h after fleroxacin administration. During the single-dose study phases, blood samples were collected as above and additionally at 36, 48, 60, and 72 h after fleroxacin administration.

Baseline urine samples (30 ml) were collected before the first dose during each study period. During the study phases, following bladder emptying, a 24-h urine sample was collected for fleroxacin analysis in the multiple-dose phases and a 72-h urine sample was collected in 24-h aliquots for fleroxacin determination in the single-dose phases. Baseline blood and urine samples were used as blank samples to ensure lack of assay interference for each subject.

Serum and urine fleroxacin concentrations were determined by using a specific high-pressure liquid chromatography assay. Briefly, to 250 μl of serum or urine was added 20 ml of 30-μg/ml pipemidic acid and 250 μl of 25% Na2SO4 (plasma) or 0.5 M phosphate buffer (pH 7.5; urine). After the tube was vortexed, 3.5 ml of chloroform was added and the tube was shaken at low speed for 10 min and centrifuged at 1,155 × g for 10 min. The top layer was then aspirated, leaving a 3-ml organic layer. To this organic layer, 200 μl of 1 N NaOH (plasma) or 1/15 M Na2PO4 (pH 12.5; urine) was added. (This mixture was shaken fast for 20 min and then centrifuged for 10 min at 1,155 × g. An injection of 150 μl was made into a Alltech Nucleosil C18, 10-μm column by using a mobile phase of 13% acetonitrile-6% methanol (19% organic) and 0.01 M phosphate buffer-0.005 M tetrabutyl ammonium hydrogen sulfate (81% aqueous) at 1 ml/min. An ABI-980 fluorescence detector at 274 nm was used.

Interday variability was 2.68 and 4.03% for 0.2- and 3.0-μg/ml concentrations, respectively. Interday variabilities for these concentrations were 4.76 and 2.28%. Inter- and intraday coefficients of variation for urine control samples were 2.06 and 1.57% and 6.08 and 3.94%, respectively. The assay had a lower limit of sensitivity of 0.05 μg/ml and a range of 0.05 to 5.0 μg/ml.

Fleroxacin data were analyzed by a noncompartmental analysis technique (5). Peak concentration (Cmax) and time to peak concentration (Tmax) were determined by graphic observation. The area under the curve (AUC) (AUC<sub>24-h</sub> for multiple dose; AUC<sub>single dose</sub> for single dose) was determined by the trapezoidal rule. For fasted phases, bioavailability was set at 1. Relative bioavailability for the fed phases was determined by calculating the ratio for each subject and then calculating the mean AUC<sub>fed</sub>/mean AUC<sub>fasted</sub> for both the single- and multiple-dose phases. Total body clearance (CL) for the single-dose and multiple-dose fed and fasted phases was determined as (dose × relative bioavailability)/AUC. Elimination rate constant (β) was determined by first graphically examining the terminal portion of the concentration-in-serum-versus-time curve and then using a generalized least-squares regression technique. The half-life (t1/2) was calculated as 0.693/β.

Statistical analyses were performed by using SAS version 6.07 (14). The analysis of variance (ANOVA) for a four-period crossover design was used to analyze AUC, Cmax, T<sub>max</sub>, CL, and β. If the overall test for differences was statistically significant, pairwise differences between regimens were tested by constructing linear contrasts. The linear contrasts were constructed within the context of the ANOVA model for the crossover design. A paired Student t test was applied to analyze the percent fleroxacin excreted unchanged in the urine (F) for the fed and fasted phases in studies of single-dose (72-h collection) and multiple-dose (24-h collection) administration, respectively. By using a bioequivalence criteria of 80 to 125%, a two one-sided tests procedure was performed to determine bioequivalence (16). A significant P value (<0.05) for this test indicates that the mean AUCs for the fed and fasted phases are bioequivalent. Westlake's 90% confidence interval for the relative bioavailability of single- and multiple-dose regimens was calculated by using log transformation of AUC data (22). Data are expressed as mean ± standard deviation. A P value of <0.05 was considered statistically significant.

RESULTS

A total of 20 subjects, 10 men and 10 premenopausal women, were enrolled in this protocol. Seven subjects were smokers (two men, five women). The mean age of the subjects was 38.3 ± 6.5 years, and their mean weight was 74.0 ± 13.1 kg. No subject had a baseline urine or serum sample positive for fleroxacin before any study phase.

Table 1 and Fig. 1 and 2 illustrate the effect of a fat- and calcium-rich breakfast on pharmacokinetic parameters of single- and multiple-dose fleroxacin administration. There were statistically significant (P < 0.05) differences between the fed and fasted phases for Cmax (reduced 25%) and T<sub>max</sub> in both single- and multiple-dose regimens. For F<sub>F</sub> (0 to 24 h), there was no statistically significant difference (P > 0.05) between the fed and fasted phases in the multiple-dose regimens. For F<sub>F</sub> (0 to 72 h), there was a statistically significant difference (P ≤ 0.05) between the fed and fasted phases in the single-dose regimens. For CL and β, there were no statistically significant
TABLE 1. Fleroxacin pharmacokinetics (mean ± standard deviation) for the fed and fasted phases in single- and multiple-dose regimens

<table>
<thead>
<tr>
<th>Phase</th>
<th>AUC (μg · h/ml)*</th>
<th>C_{max} (μg/ml)*</th>
<th>T_{max} (h)*</th>
<th>CL/F (ml/min/1.73 m²)*</th>
<th>β (h⁻¹)*</th>
<th>t_{1/2} (h)*</th>
<th>F (%)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>MD-Fast</td>
<td>96.5 ± 20.6#</td>
<td>8.5 ± 1.6</td>
<td>1.6 ± 0.8</td>
<td>67.6 ± 13.5@</td>
<td>0.078 ± 0.021@</td>
<td>8.9</td>
<td>50.7 ± 12.2</td>
</tr>
<tr>
<td>MD-Fed</td>
<td>83.1 ± 12.3#</td>
<td>6.4 ± 1.1*</td>
<td>3.5 ± 2.0*</td>
<td>76.9 ± 9.8*</td>
<td>0.072 ± 0.016*</td>
<td>9.7</td>
<td>52.7 ± 13.4</td>
</tr>
<tr>
<td>SD-Fast</td>
<td>83.3 ± 14.3</td>
<td>6.3 ± 1.0</td>
<td>1.6 ± 1.1</td>
<td>74.5 ± 11.8</td>
<td>0.065 ± 0.010</td>
<td>10.7</td>
<td>59.2 ± 13.4</td>
</tr>
<tr>
<td>SD-Fed</td>
<td>77.7 ± 13.3#</td>
<td>4.6 ± 1.0*</td>
<td>3.6 ± 2.6*</td>
<td>82.7 ± 13.4*</td>
<td>0.065 ± 0.011*</td>
<td>10.6</td>
<td>51.0 ± 11.6**</td>
</tr>
</tbody>
</table>

* MD-Fast, multiple-dose fasted phase; SD-Fast, single-dose fasted phase; MD-Fed, multiple-dose fed phase; SD-Fed, single-dose fed phase.

* Symbols: *, P ≤ 0.05 versus corresponding fasting phase, using ANOVA; **, P ≤ 0.05 versus corresponding fasted phase, using a paired t test; @, P ≤ 0.05 versus corresponding single-dose phase, using ANOVA; #, P ≤ 0.05 versus corresponding fasting phase, using the two one-sided tests procedure, where statistical significance indicates equivalence within the range ± 20% (see text for Westlake’s 90% confidence intervals for relative bioavailabilities).

Values in this column are harmonic means.

DISCUSSION

The fluoroquinolone antibiotics possess broad-spectrum antimicrobial activity and appear to have pharmacokinetic properties which make them desirable agents for oral use. However, recent data have called into question whether a food-drug interaction is present with certain fluoroquinolones. Although initial studies with drugs such as ciprofloxacin showed no effect of concurrent food ingestion on pharmacokinetics and bioavailability (4, 9, 13), recent investigations involving difference tests have shown that meals containing liquid calcium sources may cause significant reductions of both C_{max} and AUC for ciprofloxacin (13). This inhibitory effect on the absorption of quinolones when administered with meals containing liquid-calcium sources has also been shown for norfloxacin (13). Limited data on ofloxacin (12) (a small number of subjects) do not suggest substantial changes in AUC or urinary excretion when the drug was administered with liquid-calcium-containing products.

Previous pharmacokinetic studies of fleroxacin when administered in the fed state have been performed only with single-dose designs. Wachler et al. (21) administered 400-mg single doses of fleroxacin with food and reported a 17% reduction in...
C_{\text{max}}, an approximate doubling of T_{\text{max}}, and a reduction in the percent urinary excretion of fleroxacin by 12% (no significant differences). Nakashima et al. (11) reported a 21% reduction in the AUC (P \leq 0.05) after administration of a 200-mg single dose of fleroxacin with food. Neither of these studies used a liquid-calcium source for their standardized meal. Seelmann et al. reported no significant effect of a fat-rich meal on the absorption characteristics of fleroxacin (17).

The current study illustrates a number of important findings. First, although food resulted in statistical differences in the effect on fleroxacin absorption parameters after single-dose and multiple-dose administration, there was no clinical difference between results of studies of single-dose or multiple-dose administration in terms of a possible food effect. Studies of multiple-dose administration provide a more relevant pharmacokinetic profile than studies of single-dose administration since multiple-dose pharmacokinetics mimic the clinical situation of treating patients (19). Second, it is apparent that administration of a meal high in fat and liquid calcium results in a reduced C_{\text{max}} and longer T_{\text{max}} for fleroxacin. However, a meal containing high fat and a liquid-calcium source did not significantly reduce the relative bioavailability of fleroxacin in either the single-dose (91.6% ± 16.5% relative bioavailability) or multiple-dose (88.2% ± 13.4% relative bioavailability) phases. In addition, urinary excretion of fleroxacin showed no significant change in the multiple-dose study for the fed and fasted state. Finally, the analysis of these data suggests that use of difference testing by ANOVA rather than equivalence testing (two one-sided procedure), the accepted standard (16), can give opposing interpretation of the data. The current study had sufficient power to detect differences and bioequivalence between 80 and 125%. The use of equivalence testing rather than difference testing is recommended by the Food and Drug Administration Division of Bioequivalence (1).

No significant reduction in bioequivalence is seen when fleroxacin is administered single- or multiple-dose regimens concomitantly with a meal rich in fat and calcium. Although administration with food resulted in prolongation of T_{\text{max}} and reduction of C_{\text{max}} by approximately 25%, these differences are of questionable clinical significance. Fleroxacin can be administered concurrently with food and still maintain its bioequivalence with respect to administration on an empty stomach. Studies of multiple-dose administration appear to give similar information to studies of single-dose administration. However, studies of multiple-dose administration may be less costly (owing to fewer sampling periods) and may more closely mimic the clinical situation in which these agents are used.

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REFERENCES


