Key Factors in Effective Patient-Tailored Dosing of Fluoroquinolones in Urological Infections: Interindividual Pharmacokinetic and Pharmacodynamic Variability

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Abstract: Fluoroquinolones (FQs) are a critical group of antimicrobials prescribed in urological infections as they have a broad antimicrobial spectrum of activity and a favorable tissue penetration at the site of infection. However, their clinical practice is not problem-free of treatment failure, risk of emergence of resistance, and rare but important adverse effects. Due to their critical role in clinical improvement, understanding the dose-response relation is necessary to optimize the effectiveness of FQs therapy, as it is essential to select the right antibiotic at the right dose for the right duration in urological infections. The aim of this study was to review the published literature about interindividual variability in pharmacological processes that can be responsible for the clinical response after empiric dose for the most commonly prescribed urological FQs: ciprofloxacin, levofloxacin, and moxifloxacin. Interindividual pharmacokinetic (PK) variability, particularly in elimination, may contribute to treatment failure. Clearance related to creatinine clearance should be specifically considered for ciprofloxacin and levofloxacin. Likewise, today, undesired interregional variability in FQs antimicrobial activity against certain microorganisms exists. FQs pharmacology, patient-specific characteristics, and the identity of the local infecting organism are key factors in determining clinical outcomes in FQs use.

Keywords: fluoroquinolone; interindividual variability; pharmacokinetic; pharmacodynamic

1. Introduction

Fluoroquinolones (FQs) are considered a critically important antimicrobial class to human medicine [1]. They have a broad spectrum of activity against numerous Gram (+) and Gram (-) bacteria and exhibit favorable pharmacokinetic properties that facilitate adequate drug disposition at the site of infection. Due to these properties, FQs have been used to treat different types of systemic infections both in the outpatient and inpatient setting [2,3], and also a variety of urological infections, such as pyelonephritis, urethritis, and bacterial prostatitis [4]. Especially in the latter, their role is essential to saving lives. However, two clear problems currently exist in clinical practice in the management of urological infections with FQs: the emergence of uropathogen resistance [5,6] and the wide
The objective of the current review is to identify sources of interindividual variability in PK/PD properties of the most commonly FQs used in urological practice—CIP, LEV and MOX. Optimally dosing FQs is dependent on several factors, such as pathophysiological characteristics of the patient, the infecting organism, the site of infection, and the pharmacokinetic/pharmacodynamic (PK/PD) properties of the drug (Figure 1). PK properties include the factors affecting drug absorption, distribution, metabolism, and elimination, which determine its concentration in the body. Physiopathological factors associated with the patient may be responsible for the interindividual variability of the PK processes. PD describe the mechanism by which the drug exerts its antimicrobial effect [14,15]. Interregional and time-dependent differences in MIC distribution values are responsible for the PD variability of the antibiotic. Careful consideration of the factors affecting PK/PD should allow for selection of the most appropriate antibiotic when treating urolological infections and establishing the dosage with a better risk/benefit ratio in terms of efficacy, safety, and development of resistances [8,13,16,17]. The interindvidual variability in PK/PD processes is likely to be one of the main contributors to the variability in the antibacterial dose-exposure response relation [18,19].

Figure 1. Pharmacokinetic/Pharmacodynamic factors affecting the dose-antimicrobial response relation. Cmax: peak plasma concentration, AUC: area under the plasma drug concentration-time curve from time, MIC: minimum inhibitory concentration of an antibiotic against a bacterial pathogen.

The most commonly used FQs in urological practice in the United States and Europe are ciprofloxacin (CIP), levofloxacin (LEV), and moxifloxacin (MOX) [2,3,10]. The recommended doses are 250 to 500 mg orally or 400 mg intravenously for CIP, 500 to 750 mg orally or intravenously for LEV, and 400 mg orally or intravenously for MOX [2–4]. In general, these dosing regimens allow adequate drug distribution at the site of infection to achieve sufficient antibiotic exposure and treatment efficacy. However, the use of standard empirical doses may lead to unnecessary overexposure and a higher incidence of adverse effects in some patients and, consequently, non-compliance or discontinuation of drug treatment. In contrast, some patients could experience underexposure, risking treatment failure and the development of bacterial resistance [11–13].

Array of reported adverse effects [7,8]. Consequently, their use has been limited to strictly necessary situations in clinical practice [7–9].

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MOX—which could influence the antibacterial dose-exposure response relation. Evaluation of their effect on FQs clinical outcomes would allow the development of patient-tailored dosing strategies, leading to reduced treatment failure, especially in critical infections as prostatitis.

2. Concerns about the Clinical Efficacy of Fluoroquinolone Dosing: The Role of PK/PD Index as a Tool

Selecting a dose to guarantee antimicrobial efficacy while minimizing the risk of resistance emergence with minimum adverse effects should be based on the FQ’s PK properties, PD properties, and the probability to attain the PK/PD index associated with clinical efficacy with the administered dose [14,16,20,21]. PK properties refer to drugs’ absorption, distribution, metabolism, and excretion, while PD properties are related to the potential activity of an antibiotic against a bacterial pathogen, measured as the minimum inhibitory concentration (MIC), and postantibiotic effect. The PK/PD target is the key value associated with antimicrobial efficacy and varies according to a chosen endpoint such as stasis, maximal kill, or resistance suppression for preclinical studies, and microbiological or clinical cure for clinical studies [15,16,22]. To attain a favorable PK/PD target for FQ, pathogens must have adequate antimicrobial exposure based on their MIC. This exposure, measured as peak plasma concentration (Cmax) and area under the plasma drug concentration-time curve from time 0–24 h ($AUC_{0–24h}$), is dependent on the dose used and the PK properties of the drug. Only the unbound drug concentrations are microbiologically active, and therefore, the PK/PD index should be based on free drug concentrations [21].

A substantial number of studies have been performed to identify the PK/PD index relation associated with the bactericidal activity and clinical efficacy of FQs. Antimicrobial activity of FQs exhibits concentration-dependent killing along with prolonged persistent effects. Multiple clinical and preclinical data suggest that the ratio of free $AUC_{0–24h}$ (f$AUC_{0–24h}$) to MIC (f$AUC_{0–24h}$/MIC) is the best PK/PD index to link antimicrobial disposition, the MIC value, and the clinical efficacy of FQs [15,23,24]. While the ratio Cmax/MIC outweighs the ratio f$AUC_{0–24h}$/MIC as an indicator of resistance suppression, fewer studies exist with the aim to identify drivers for resistance suppression [11].

Based on the above concepts, Monte Carlo simulations (MCS) can be run to computationally estimate the likelihood of a given drug dose to attain a predefined value of a PK/PD target previously defined for urological FQs [19,21]. The probability of target attainment (PTA)—defined as the probability that a specific value of the PK/PD index associated with the efficacy of the antibiotic is achieved at a certain MIC [20]—can therefore be calculated for different MIC values against a variety of pathogens. Thus, PK/PD indexes can be used as a tool to select dosing regimens with PTA >90% in the studied population, increasing the probability of selecting clinically successful treatments, identifying clinical breakpoints, and preventing the emergence of resistance [14,19].

The use of these metrics is therefore essential to streamline FQs treatment and adjust dosing regimens in clinical practice. In addition, the individual status of the patient and the suspected infecting organism should be accounted for in dose decision-making. In an exposure–response model based on clinical data from patients with community-acquired pneumonia associated with S. pneumoniae, Gram (+) microorganism, and treated with a 500 mg oral dose daily of LEV, the probability of successful clinical response was 95% in patients who achieved a target of f$AUC_{0–24h}$/MIC > 33.8, and was 67% in patients who did not achieve that target [23]. For infections caused by Gram (-) bacteria, the threshold of the ratio $AUC_{0–24h}$/MIC was found to be a significant breakpoint for probabilities of both clinical and microbiological cures, but the required value for treatment success was higher than that required for Gram (+) bacterial infections. In hospitalized patients treated with a 400 mg intravenous dose of CIP, at $AUC_{0–24h}$/MIC > 125, the probabilities of clinical and microbiological cure were 80% and 82%, respectively, for Gram (-) bacteria [25]. Similar results were obtained for patients with nosocomial pneumonia and Gram (-) isolates treated with a 750 mg intravenous dose of LEV [24]. The United States Committee on Antimicrobial
Susceptibility Testing (USCAST) report provides an extensive integrative evaluation of the in vitro susceptibility testing, PK/PD breakpoints (i.e., AUC$_{0-24h}$/MIC ratio targets for Enterococcus spp., Escherichia coli, and other Enterobacteriaceae spp., Pseudomonas aeruginosa, and Streptococcus aureus), and clinical breakpoints by infection type [26]. However, similar studies in patients with urological infections are still scarce.

Published PK/PD studies have raised concerns about patient clinical outcomes with the use of FQs, particularly with regards to treatment failure after empiric dosing related to variability in PK and bacteria susceptibility, as MIC value. Rattanaumpawan et al. [27] examined the impact of PD variability associated with MIC value on FQs’ clinical outcomes in adult female patients with complicated urinary tract infections caused by E. coli. Treatment failure rates of 0.8% and 6.9% were observed when they compared isolates with low and high MIC values, respectively. Peloquin et al. [28] found that resistance to CIP occurred in patients after treating nosocomial lower respiratory tract infections caused by Gram (-) bacteria, such as P. aeruginosa. The authors explained that the Cmax/MIC ratio was far greater with isolates that were eradicated than with those that persisted, attributing variability in MIC values as the determining factor in treatment resolution. In patients with nosocomial pneumonia, measuring CIP and LEV concentrations, determining pathogen MIC, and subsequently performing dose adjustments significantly improved the probability of successful clinical outcomes and pathogen eradication [29]. For patients diagnosed with Mycoplasma genitalium, a pathogen transmitted through sexual contact, the increased use of 400 mg of MOX caused an emergence of cases with treatment failure. The MIC of MOX in the mutant strains increased 4-fold as compared with that of the parent strain [30].

Noreddin et al. [31] determined that age plays an important role in explaining the interindividual variability in the PK of hospitalized patients with community-acquired pneumonia treated with LEV. This factor influenced the ability to achieve the target attainment related with clinical response. Variations in clinical response to S. pneumoniae were observed when comparing elderly and younger patients. Elderly patients showed higher AUC$_{0-24h}$ values leading to a higher AUC$_{0-24h}$/MIC ratio and improved bacteriological outcome compared to younger adults. In another study analyzing CIP use in hospitalized patients with urinary tract, abdominal, and various other infections produced by Gram (-) microorganisms, 21–75% of the patients did not achieve the efficacy target of AUC$_{0-24h}$/MIC ratio ≥125 with MICs of 0.25 and 0.5 mg/L, respectively. The AUC$_{0-24h}$ achieved with standard dosing was related to the high interindividual variability of CIP clearance, associated with age and serum creatinine. This pharmacokinetic variable and the elevated MIC values observed in this study highlight the need for individualizing dosing regimens to maximize efficacy, minimize adverse effects, and prevent the appearance of resistance [32].

Ongoing efforts are still need to identify optimal FQs dosing strategies to achieve the efficacy target of AUC$_{0-24h}$/MIC during urological clinical use due to the critical role that the management of these drugs plays in this type of infections [2,12,21]. Interindividual variability in FQs PK and urological microorganism MIC needs to be known and considered during dose adjustment in patients, as its variability can negatively influence the probability of reaching the efficacy PK/PD index and achievement of successful clinical outcomes [33,34].

3. Interindividual Pharmacokinetic Variability and Their Causes

Antimicrobial exposure related to drug disposition, given as AUC$_{0-24h}$ and Cmax, is subject to interindividual variability in PK properties, such as drug absorption, distribution, and elimination. Tables 1–3 summarize the PK parameters and their important interindividual variability for CIP, LEVO, and MOX, respectively, in different patient populations extracted from literature. It is important to consider that the estimation of individual PK parameters for each patient allows the estimation of individual exposures to the drug; therefore, population PK studies are essential to achieve this objective [18,19,33,34].
the following sections, the factors identified as potential sources of variability in the PK processes of FQs are discussed (absorption, distribution, and elimination).

### 3.1. Absorption Process: Role of Food

Absorption refers to the amount of drug reaching the bloodstream from the site of administration. FQs are well absorbed after oral administration with bioavailability (F) values of 70% for CIP [35], 99% for LEV [36], and 86% for MOX [37]. Concomitant oral administration of antacids containing multivalent cations, such as calcium, aluminum, or magnesium, calcium or iron supplements, and sucralfate, decrease FQs absorption due to the formation of insoluble quinolone-multivalent cation chelates in the gastrointestinal tract. For example, for CIP, F decreases to 15% with concomitant aluminum and magnesium antacid use within 5 to 10 min of drug administration [38]. Similar effects have been reported for milk, other dairy products, and supplements containing multivalent cations. The extent of the interaction diminishes when the interacting drug is administered at least 2 to 4 h before or 6 to 8 h after the FQs [39]. Multivalent cations present in food, supplements, or other drug products can lead to clinically relevant interactions with FQs, contributing to variability in drug absorption, reducing the overall exposure, and increasing the risk for therapeutic failure. Conversely, food not containing multivalent cations is not expected to modify FQs absorption [39,40].

According to the Food and Drug Administration’s Biopharmaceutics Classification System (BCS), CIP is categorized as class III, though this is somewhat controversial, with some authors classifying CIP as class II/IV. Unlike LEV or MOX, CIP presents a pH-dependent solubility. It is highly soluble at an acidic pH, however, at an intestinal pH of 6.8 to 7.5, its solubility is much lower [41]. Any meals or beverages able to significantly affect the pH may thus affect CIP oral bioavailability [42].

### 3.2. Distribution Process: Role of Patient’s Pathophysiological Characteristics

After entering systemic circulation, the drug must distribute throughout the body via the bloodstream to the tissues. The extent of drug distribution depends on a variety of factors, including the physicochemical properties of the drug, the rate of blood flow to the tissue, and the ability of the drug to bind to plasma proteins and tissue. Given that only unbound or free drugs can access the site of infection [43,44], the influence of plasma protein binding on the distribution of FQs was evaluated. The percent of plasma protein binding is low for FQs (30% for CIP [45], 31% for LEV [46], and 48% for MOX) [37]. Moreover, it has not been established that variability in plasma protein binding has any significant direct or indirect impact on the therapeutic effectiveness of FQs [47]. Regarding the tissue distribution, the physicochemical properties of FQs permit rapid penetration into extravascular and intracellular sites, with a rapid equilibrium established between compartments. CIP, LEV, and MOX are widely distributed throughout the body and reach high concentrations in a variety of tissues, such as the urinary tract (e.g., urine, prostate) [48,49], and other areas such as the lungs, paranasal sinuses, inflamed lesions, and bones [50,51].

Specifically, FQs are effective in the treatment of many types of urological infections and other systemic infections due to their ability to achieve high concentrations in tissues and body fluids and their wide antibacterial spectrum. However, several complex mechanisms are involved in the penetration of special tissues, such as prostate. In addition to passive diffusion [52], conditioned by the drug’s acidic or alkaline nature, its pKa, and the pH of prostatic fluid, there is evidence of the involvement of efflux transporters—primarily P-glycoprotein—on FQs tissue penetration (Figure 2).

The results of Zimmermann et al. strongly support the role of efflux transporters on the prostatic tissue penetration of LEV [53], but not of CIP [54]. Due to this complexity, interindividual variability in drug penetration into tissues could result in variability of concentrations at the site of infection and condition the effectiveness of the treatment or affect to the emergence of bacterial resistance. Whole body physiologically based pharmacokinetic
(PBPK) models provide a valuable tool to incorporate drug disposition characteristics—including the role of transporters—and predict unbound tissue distribution in different organs. The application of PBPK modeling has increased over the past decade to improve the mechanistic understanding of drug PK and support dosing recommendations [55]. PBPK models can also incorporate relevant disease-specific changes in the physiology, allowing the prediction of drug PK under different chronic conditions, as for example renal or hepatic disease, heart failure, or obesity [56–58].

![Figure 2](image-url) Distribution mechanisms of fluoroquinolone from the blood to the prostate gland. F: unbound fluoroquinolone; FP: protein-bound fluoroquinolone.

Distribution studies use central blood/plasma concentrations as a surrogate for tissue distribution as they are easily accessible to measure. Volume of distribution (Vd) is the PK parameter that represents the degree to which a drug is able to distribute throughout the body to the tissues [59]. Body weight and its changes in obese patients, age, and pathological condition of patients can explain the interindividual variability of distribution seen with FQs to a certain degree. As can be observed in Table 1, when CIP was infused to a group of obese, the Vd was found to be 23% larger in the obese group than in the non-obese group. However, when the Vd was adjusted for the total body weight, the obese exhibited lower Vd/kg than the non-obese subjects. These findings indicate that CIP is not highly distributed into adipose tissue [60]. Additional population PK analysis studies conducted in elderly patients [61–63] and in adult patients with septic shock [64] revealed that total body weight is a significant covariate on the Vd of CIP. No significant changes in Vd have been found in patients with hepatic or renal impairment [65–67]. A high variability in this parameter was observed in patients with critical illness, but no covariates were associated with this variability due the complex of the pathology [62,64,68,69]. For LEV, patient-
specific factors such as age, sex, and race [70], but not obesity—even considering obese patients and severely morbidly obese [71,72]—contributed to variability in Vd (Table 2). The Vd of MOX, however, was not significantly affected by age or sex [73], but was found to be correlated with lean body weight for both normal weight [74] and obese patients [75], as has been published in the articles referenced in Table 3.

3.3. Elimination Process: Role of Renal and Hepatic Function

FQs are eliminated from the body via two main mechanisms: biotransformation—or hepatic metabolism—and renal excretion. Once the antibiotic reaches the systemic circulation, these elimination processes function to decrease the blood concentration of FQs and consequently decrease antibiotic exposure at the site of infection [23,34]. Clearance (CL) is defined as the volume of body fluid, usually plasma, from which the drug is completely removed per unit of time. This PK parameter reflects the rate of drug elimination from the body and is proportional to the blood concentration of the drug. For every drug, each organ of elimination has its own clearance (e.g., hepatic clearance or renal clearance). The total body clearance of the drug is therefore the sum of the clearances from all eliminating organs (CL = CL\text{Renal} + CL\text{Hepatic} + CL\text{Other}) [59]. Clearance is the factor determining the average concentration of FQs after continuous intravenous infusion. After oral administration, however, the elimination process is determined by both the clearance and absorption process and underlying bioavailability (CL/F).

Different factors contribute to the interindividual variability in the CL of the urological FQs under review. First, Table 1 summarizes the changes in PK parameters related to CIP elimination processes (T\text{1/2} and CL) and different patient subpopulations. For CIP, several mechanisms and factors may contribute to the interindividual variability in CL. Non-renal mechanisms of elimination—mainly hepatic metabolism—account for approximately one-third of CIP elimination. Four metabolites of CIP—desethyleneciprofloxacin, sulfo-ciprofloxacin, oxo-ciprofloxacin, and N-acetylciprofloxacin—have been recovered in the urine and feces. Due to changes in chemical structure, these metabolites have some antibacterial activity, but less than that of the parent compound [46]. Approximately 15% of a 100 mg intravenous dose of CIP is excreted in the feces, likely due to elimination directly through the intestinal mucosa and biliary excretion. The remaining two-thirds of the CIP dose is eliminated via the kidneys, due to a combination of glomerular filtration and tubular secretion [76]. As a result of undergoing CL through both non-renal and renal pathways, CIP has a relatively short half-life when compared to other FQs and requires twice daily dosing [39,46]. Population PK modeling has been used in several studies to estimate the effect of individual PK parameter values in a variety of patient populations and bacterial infections [61,62,64,68,69,77–80], showed in Table 1. Factors affecting renal and hepatic function could also be responsible for the interindividual variability in the CL of CIP, and their effect may be difficult to predict. Hepatic dysfunction appears to have minimal effect on the elimination of CIP, with no changes in CL found in chronic cirrhotic patients [65]. Creatinine clearance (CL\text{CR}), however, has been identified in multiple population PK studies as one of the main covariates responsible for interindividual variability in the systemic CL of CIP. In patients with varying degrees of renal dysfunction, CIP CL has been shown to decrease as CL\text{CR} decreases [66,67,77]. Consequently, age-related decline in renal function could also lead to a reduction in CIP elimination in older adults [61–63]. In addition, an increase in CL has been reported in obese patients when compared to patients of normal weight, which could be related to the increase in glomerular filtration and tubular secretion known to occur in obese adults [60]. Lastly, critically ill patients exhibit higher interindividual variability in CL associated with pathophysiological changes driven by altered renal function [62,64,68,69,80]. Non-renal mechanisms, such as biliary clearance, may effectively compensate for the reduction in renal CL in these patients, and could further contribute to the increase in interindividual variability [64,80].

Approximately 83% of LEV is excreted in the urine as an unchanged drug, indicating that it primarily undergoes renal elimination [36,39]. Similarly to CIP, population PK
modeling has been used in several studies with LEV to estimate the effect of individual PK parameter values in a variety of patient populations and bacterial infections [70,81–93] (Table 2). In several studies, CL_{CR} [71,82–85,91–93], age [70], and race [70] were found to be covariates that influenced the CL of LEV. In hospitalized elderly patients with varying degrees of renal function, CL_{CR} was again shown to be the main covariate associated with interindividual variability in LEV CL [90]. A prospective population PK study conducted in patients with bone and joint infections demonstrated that age and glomerular filtration rate were covariates related to interindividual variability of CL/F [81]. Critical illness was not a significant variable in altering LEV CL per se, with altered renal function being the determining factor [77,93–96]. Obesity may be another factor affecting the interindividual variability of LEV PK. However, most studies with LEV have been performed in normal weight patients, and only a few published studies performed in overweight and obese patients. One such study reported a higher CL of LEV in morbidly obese patients and suggested that CL was related to CL_{CR} estimated by the Cockcroft–Gault equation and ideal body weight [72].

MOX primarily undergoes hepatic metabolism and fecal excretion. Despite the large percentage of metabolism by the liver, moxifloxacin does not appear to be transformed by the cytochrome P450 (CYP) isoenzyme system, making it less susceptible to drug–drug interactions. Moxifloxacin has two metabolites, M1 (sulpho-compound) and M2 (glucuronide) [96–98]. Total clearance is modified only by lean body weight in healthy adults [37,73]. As shown Table 3, the PK after a single and multiple intravenous doses of MOX differed only marginally in patients with severe hepatic impairment compared to healthy patients, and demonstrated no accumulation [99]. Only 20% of MOX is excreted unchanged by the kidneys, conditioned by the processes of glomerular filtration and tubular reabsorption. As a result, renal impairment has little clinically relevant effect on the PK of MOX, including CL, and does not require dose adjustments [96]. MOX PK in critically ill patients with acute renal failure undergoing extended daily dialysis are similar to healthy patients without renal impairment [96,100,101]. Other patient-specific factors such as age [74], race [102], and obesity [75] have not been shown to be responsible for the interindividual variability in the CL of MOX.

FQs exhibit dose-independent PK, meaning that F, CL, and Vd are constant over the range of doses encountered clinically [39]. Several pathophysiological factors, that may be present in patients with urological infections, could influence interindividual variability in FQs PK and potentially affect clinical response and outcomes. Given their large Vd and ability to accumulate in tissues, interindividual variability in the Vd of FQs could affect the degree that FQs are able to penetrate the site of infection. Additional tissue distribution studies could therefore help to better understand variability in the Vd of FQs, especially associated with patients’ pathophysiological characteristics. However, the importance of CL is far more evident [103]. Given FQs’ concentration-dependent antibacterial activity, understanding the interindividual variability of CL after intravenous administration and F variability after oral administration is crucial to ensuring adequate antibiotic exposure–AUC–is achieved and maintained when treating urological infections. CL, especially for CIP and LEV, decreases fundamentally with decrease in renal function. This decrease in CL translates to a higher AUC in patients and, as a result, a higher probability of experiencing concentration-dependent adverse effects [104,105]. Another important aspect to consider is the impact of drug–drug interactions (DDIs) on drug exposure. PBPK modeling and simulation can be used as a tool to determine the impact of disease-related physiological changes and DDIs on the systemic exposure of FQs, and the possible need of dose adjustment in specific diseases and/or due to co-medications [106]. As an example, alterations in blood flow to main organs and decrease in clearance observed in chronic kidney disease or chronic heart failure can be incorporated in the model to predict changes in the ADME properties of FQs. In addition, mechanistic modeling can be used to explore possible disease effects, test hypotheses, and generate supporting evidences when not enough clinical data are available [107].
Table 1. Steady-state pharmacokinetic parameters for ciprofloxacin in patients with several physiopathology conditions after intravenous or oral administration (values expressed as mean (standard deviation)).

| Ciprofloxacin | Study Characteristic | Vd (L) | Cl (L/h) | T<sub>1/2</sub> (h) | Reference |
|---------------|----------------------|--------|----------|---------------------|-----------|
| Healthy, non-obese | 200 mg Infusion IV 21–30 years | 199.1 (34.2) | 26.8 (5.7) | 4.2 (0.8) | Plaisance et al., 1987 [35] |
| | 200 mg Infusion IV 21–30 years | 219.0 (35.8) | 44.6 (7.2) | 4.0 (0.3) | Allard et al., 1993 [60] |
| | 146.0 (27.4) | 25.2 (5.8) | 4.4 (0.9) | Drusano et al., 1986 [77] |
| | 750 mg Oral | 256.0 (80.0) | 29.5 (5.9) | 5.2 (0.7) | Plaisance et al., 1987 [35] |
| | 46–68 years | 217.0 (45.0) | 50.4 (14.4) | 3.7 (0.4) | Drusano et al., 1986 [77] |
| Healthy, obese | 400 mg Infusion IV 29 ± 7 years BMI = 36 ± 4 kg/m² | 269.1 (51.6) | 53.8 (9.5) | 4.3 (0.6) | Allard et al., 1993 [60] |
| Patients with cirrhosis | 750 mg Oral 52 ± 6 years | 218.1 (45.4) | 45.9 (14.1) | 3.7 (0.4) | Frost et al., 1989 [65] |
| Patients with renal disease | 200 mg Infusion IV 22–62 years | 191.7 (35.4) | 26.8 (5.7) | 4.3 (0.8) | Drusano et al., 1987 [66] |
| | CL<sub>CR</sub> ≥ 100 mL/min | 243.0 (97.1) | 26.3 (10.3) | 6.1 (1.6) |
| | CL<sub>CR</sub> = 86–60 mL/min | 183.2 (47.7) | 15.0 (3.8) | 7.7 (1.2) |
| | CL<sub>CR</sub> = 11–57 mL/min | 210.2 (70.8) | 15.4 (4.3) | 8.5 (3.3) |
| | CL<sub>CR</sub> = 0 mL/min | 158.0 (46.5) | 70.4 (48.9) | 3.5 (1.2) | Gasser et al., 1987 [67] |
| | 750 mg Oral 48–90 years | 113.8 (34.2) | 29.4 (6.4) | 6.3 (3.2) |
| Elderly patients | 200 mg Infusion IV 78 ± 11 years | 100.8 (37.8) | 16.6 (6.8) | 5.8 (2.4) | Hirata et al., 1989 [63] |
| | 200 mg Infusion IV 73 ± 11 years | 61.0–118.0 | 18.4 (4.5) | ND | Cios et al., 2014 [61] |
| Acutely ill patients | 200–400 mg Infusion IV 24–91 years | 111.0 (33.0) | 17.0 (6.6) | ND | Forrest et al., 1993 [25] |
| | CL<sub>CR</sub> = 63 ± 30 mL/min | 255.0 (51.0) | 25.4 (67.8) | ND | Abdulla et al., 2020 [68] |
| | 400 mg Infusion IV 23–79 years | 107.5 (21) | 18.6 (18.7) | ND | Li et al., 2019 [80] |
### Table 1. Cont.

| Study Characteristic | Vd (L) | Cl (L/h) | T_{1/2} (h) | Reference |
|----------------------|--------|----------|-------------|-----------|
| **Ciprofloxacin**    |        |          |             |           |
| 400–600 mg Infusion IV 24–89 years | ND | 15.2 (42.9) | ND | Roberts et al., 2019 [64] |
| Cl\textsubscript{CR} = 7–204 mL/min | | | | |
| 400 mg Infusion IV 55–77 years | 160 (51.2) | 10.7 (46.9) | ND | Roger et al., 2016 [79] |
| 200–400 mg Infusion IV 30–87 years | ND | 20.3 (51.2) | ND | Gieling et al., 2020 [69] |
| GFR = 23–208 mL/min | | | | |

Vd: volume of distribution in steady state; Cl: systemic clearance; BMI: body mass index calculated as: body weight [in kilograms]/height\textsuperscript{2} [in meters]; GFR: Glomerular filtration rate (mL/min) by MDRD (MDRD: Modification of Diet in Renal Disease Study Group developed a four-variable formula to estimate the GFR); Cl\textsubscript{Cr}: creatinine clearance; T\textsubscript{1/2}: elimination half-life; D: unpublished data. \textsuperscript{1} value of apparent volume of distribution in steady state (Vd/F) and apparent clearance (Cl/F), respectively.

### Table 2. Steady-state pharmacokinetic parameters for levofloxacin in patients with several physiopathology conditions after intravenous or oral administration (values expressed as mean (standard deviation) or mean (range) when standard deviation is not published).

| Study Characteristic | Vd (L) | Cl (L/h) | T_{1/2} (h) | Reference |
|----------------------|--------|----------|-------------|-----------|
| **Levofloxacin**     |        |          |             |           |
| Healthy young volunteers | | | | |
| 500 mg Oral 22–36 years | 90.6 (11.9) \textsuperscript{1} | 9.5 (1.7) \textsuperscript{1} | 7.0 (0.8) | Chien et al., 1997 [36] |
| Cl\textsubscript{CR} = 90–117 mL/min | | | | |
| Healthy elderly volunteers | | | | |
| 500 mg Oral 66–75 years | 70.8 (8.4) \textsuperscript{1} | 7.3 (1.9) \textsuperscript{1} | 7.6 (2.0) | Chien et al., 1997 [36] |
| Cl\textsubscript{CR} = 47–80 mL/min | | | | |
| Patients with respiratory, urinary, and other infections | | | | |
| 250–500 mg Infusion IV 47 ± 18 years | ND | 9.3 (4.3) | ND | Preston et al., 1998 [70] |
| Cl\textsubscript{CR} = 86 ± 31 mL/min | | | | |
| Patients adults with pulmonary tuberculosis | | | | |
| 1000 mg Oral 30–54 years | (33.5–114.5) \textsuperscript{1} | 7.6 (1.5–19.2) \textsuperscript{1} | ND | Peloquin et al., 2008 [82] |
| Cl\textsubscript{CR} = 51–125 mL/min | | | | |
| Patients with bone and joint infections | | | | |
| 750 mg Oral 57 ± 20 years | 90.6 \textsuperscript{1} (0.06) | 6.10 (0.17) \textsuperscript{1} | ND | Eloy et al., 2020 [81] |
| BW = 72 ± 16 kg Cl\textsubscript{CR} = 120 ± 74 mL/min | | | | |
| Obese patients | | | | |
| 750 mg Infusion IV 18–55 years | 83.8 (21.6) | 9.8 (4.2) | 5.9 (3.5) | Cook et al., 2011 [71] |
| BMI (kg/m\textsuperscript{2}) = 49.3 ± 20.7 Cl\textsubscript{CR} = 140 ± 64 mL/min | | | | |
| Acutely hospitalized older patients with several degrees of renal function | | | | |
| 125–750 mg Oral 81 ± 28 years | ND | 2.53 (1.46) \textsuperscript{1} | ND | Cojutti et al., 2017 [90] |
| Cl\textsubscript{CR} = 18–50 mL/min | | | | |
Table 2. Cont.

| Study Characteristic | Vd (L) | Cl (L/h) | T_{1/2} (h) | Reference |
|----------------------|--------|----------|-------------|-----------|
| **Intensive Care Unit** |        |          |             |           |
| Acute renal failure 500 mg Infusion IV 33–62 years | 114.0 (74–155) | 3.1 (2.9–3.2) | 34.5 (21.2–47.7) | Czock et al., 2006 [96] |
| Acute renal failure 33–62 years | 82.8 (50.0) | 2.5 (0.9) | 21.8 (5.5) | Hansen et al., 2001 [95] |
| Critical illness in continuous hemodiafiltration 500 mg Infusion IV 59 ± 6 years | ND | 3.6 (0.4) | ND | Wada et al., 2015 [91] |
| Continuous veno-venous hemofiltration 500 mg Infusion IV 23–70 years | ND | 1.8–3.6 | ND | Malone et al., 2001 [93] |
| Continuous veno-venous hemofiltration 500 mg Infusion IV 68 ± 5 years | 105.7 (36.4) | 3.26 (1.4) | 28.0 (4.5) | Guenter et al., 2002 [94] |

Vd: volume of distribution in steady state; Cl: systemic clearance; BW: body weight (kg); BMI: body weight [in kilograms]/height² [in meters]; ClCr = creatinine clearance; T_{1/2}: elimination half-life. ND: unpublished data.

1 value of apparent volume of distribution in steady state (Vd/F) and apparent clearance (Cl/F), respectively.

Table 3. Steady-state pharmacokinetic parameters for moxifloxacin in patients with several pathophysiologic conditions after intravenous or oral drug administration. Values expressed as mean (standard deviation) or mean (range) when standard deviation is not published.

| Moxifloxacin | Study with | Vd (L) | Cl (L/h) | T_{1/2} (h) | Reference |
|--------------|------------|--------|----------|-------------|-----------|
| **Healthy volunteers** |           |        |          |             |           |
| 200 mg Oral 33 ± 5 years | 222.0 (1.2) | 13.1 (0.1) | 11.8 (1.2) | Stass et al., 1998 [73] |
| 400 mg Oral 18–46 years | 175.9 (19.4) | 101.0 (2.1) | ND | Grosjean et al., 2012 [74] |
| **Morbidly obese patients** |           |        |          |             |           |
| BMI > 40 kg/m² 400 mg Infusion IV 41 ± 12 years | 165.0 (30.0) | 9.6 (2) | 12.2 (2.2) | Keess et al., 2011 [75] |
| Hospitalized severe liver insufficiency with pneumonia or spontaneous bacterial peritonitis 400 mg Infusion IV 40–78 years | 154.1 (118.5–216.1) | 8.8 (6.4–10.5) | 10.4 (8.5–16.0) | Barth et al., 2008 [99] |
| Outpatients with pneumonia receiving hemodialysis 400 mg Oral 47–78 years | ND | 6.5 (1.9) | ND | Tokimatsu et al., 2017 [101] |
| **Critical ill patients receiving continuous hemodiafiltration** |           |        |          |             |           |
| 400 mg IV infusion 25–76 years | 266 | 15.7 | 12.3 | Czock et al., 2006 [96] |
Table 3. Cont.

| Study with Vd (L) Cl (L/h) T_{1/2} (h) Reference |
|------------------------------------------------|
| Intensive care unit with COPD 2 400 mg Infusion IV 70 ± 10 years | 115.0 (40.0) 8.85 (2.6) 9.7 (3.7) Sionidou et al., 2019 [100] |

Vd: Volume of distribution in steady state; Cl: Systemic clearance; T_{1/2}: elimination half-life; IV: Intravenous administration; BMI: Body mass index, defined as body weight [in kilograms]/height^2 [in meters]; ND: Unpublished data. 1 Value of apparent volume of distribution in steady state (Vd/F) and apparent clearance (Cl/F), respectively. 2 COPD: Chronic Obstructive Pulmonary Disease.

4. Antibacterial Activity of FQs: Interregional Variability in Pharmacodynamic Properties

When FQs reach the site of infection at an appropriate concentration and remain there for sufficient time, they interact with the microorganism, resulting in an antibacterial effect (Figure 1) [108]. The antibacterial effect is related to the specific spectrum of activity of each FQ. CIP has the most potent activity against Gram (-) bacteria—including Enterobacteriaceae and P. aeruginosa—and atypical bacteria, with little to no activity against Gram (+) bacteria. LEV and MOX retain activity against Gram (-) and atypical bacteria similar to CIP, but expand coverage to include certain Gram (+) bacteria, such as S. pneumoniae [46]. In addition, MOX is active against anaerobic bacteria. The broad spectrum of activity makes FQs highly effective against a wide variety of acute and chronic bacterial infections, including urological infections [2–4,109].

MIC is the most relevant PD parameter to define the potential inhibitory activity of an antimicrobial against a microorganism. Antibiotic susceptibility rates for bacterial pathogens are available through the European Committee on Antimicrobial Susceptibility Testing (EUCAST) guidelines (European Society of Clinical Microbiology and Infectious Diseases) [110], the American Clinical and Laboratory Standards Institute (CLSI) guidelines [111], and local databases.

MIC distribution could vary according to whether the microorganisms are sensitive, intermediate, or resistant to the antibiotic [11,13]. Resistance to FQs can occur through various mutational mechanisms, including alterations in the target enzymes, DNA gyrase and topoisomerase IV, or in the permeability of the cytoplasmic membrane and expression of efflux pumps and proteins [46]. Global surveillance studies demonstrated that FQs resistance rates increased in the past years in almost all bacterial species [112]. This has led to the development of many antimicrobial surveillance programs, such as the SENTRY Antimicrobial Surveillance Program [113], the Center for Disease Control and Prevention’s National Healthcare Safety Network (CDC NHSN) [114], and the European Antimicrobial Resistance Surveillance Network (EARS-Net) [115], which are essential in the fight against emerging resistance. These programs provide information on microorganism frequency and distribution and antimicrobial resistance trends in different geographical regions and in nosocomial and community-acquired infections using information from medical centers worldwide by antimicrobial susceptibility testing in a central laboratory. This information has the potential to guide therapeutic approaches for serious infections and may have value in the prevention and control of infection [114]. Due to the ongoing emergence of antibiotic resistance, it is critical to take into account the local MIC values and susceptibility patterns for different microorganisms, and their time-dependent evolution, as they are responsible for PD variability of FQs [11,13]. An increase in the use of FQs for the treatment of infections caused by P. aeruginosa has led to reductions in susceptibility rates by agent and by geographical region, with consequences in microbiological and clinical outcomes. Rates of resistance of P. aeruginosa strains to CIP ranged from 23.2% in North America, 29.7% in Europe, 17.8% in Asian-Pacific, and 40.3% in Latin America between 1997 and 2016 [116]. E. coli is responsible for causing multiple urological infections, and the development of
FQs-resistant strains could have a significant impact on clinical efficacy and outcomes in the treatment of these infections [117–119]. Resistance rates of Neisseria gonorrhoeae to FQs are highly variable, with rates in Asia as high as 40% to 100%, whereas resistance rates in Europe and North America range from <10% in rural areas to >30% in established sexual networks [112]. Higher rates of FQs resistance are expected in intensive care units (ICUs) due to the multiple factors, including frequent use of broad-spectrum antibiotics, multitude of invasive procedures, and increased likelihood of multidrug resistant pathogen transmission. For CIP and LEV, rates of resistance to E. coli, Klebsiella spp., Enterobacter spp., and P. aeruginosa were shown to be 35%, 12%, 9%, and 32%, respectively, in the United States, and 24%, 25%, 24%, and 39%, respectively, in Europe [119–121].

Differences in the MIC distribution for a specific bacterial pathogen could have a key role in the clinical and microbiological response after a standard empiric dose [14,16,17]. Since the selection of an antimicrobial therapy and its dose should be guided based on local susceptibility and resistance patterns, there is a critical need for determination of current antibacterial resistance rates and their time-dependent evolution at a local scale for urological infections [17,119]. However, in addition to the local susceptibility profile, the PK/PD analysis that allows to estimate the CFR should also be considered to optimize the antimicrobial dosing selections for clinical decision making. In fact, susceptibility data alone are not always useful for detecting changes in the likelihood of treatment success [122,123].

FQs susceptibility rates can vary widely for different bacterial pathogens, which can affect the ability to achieve the PK/PD indices necessary for clinical and microbiological cure after a drug dose, without ignoring the individual patient’s PK [114]. Several MCS analyses demonstrated that a 400 mg intravenous dose of CIP given every 12 h to critically ill patients achieved a PTA > 90% for an AUC/MIC ≥ 125 for isolates with an MIC of 0.25 mg/L. However, the PTA decreased to 50% and 10% as the MICs increased to 0.5 mg/L and 1 mg/L, respectively, for Gram (-) bacteria [25]. Another study demonstrated FQs treatment failure rates of 0.8% for E. coli isolates with an MIC of ≤0.12 mg/L compared to 6.9% for isolates with an MIC of >0.12 mg/L to ≤2 mg/L in adult female patients with complicated urinary tract infections [27].

Considering FQs’ interindividual pharmacological variability, developing urological patient-tailored effective dosing strategies in order to improve microbiological and clinical outcome may be necessary [13,19,124], as is proposed in Figure 3.

**Figure 3.** Model-based approach to select FQ dosing regimen based on % probability of target attainment. Scheme of the steps needed to apply PK/PD analysis and Monte Carlo simulation in clinical setting. PK: Pharmacokinetics; PD: Pharmacodynamics; CL: Clearance; F: Biodisponibility; AUCPL: area under the plasma drug concentration-time curve from time 0–24 h; MIC: minimum inhibitory concentration; PTA: Probability of target attainment.
5. Conclusions

Due to a broad antimicrobial spectrum of activity and a favorable tissue penetration at the site of infection, fluoroquinolones (FQs) are a critical group of antimicrobials prescribed in urological infections, especially in prostatitis where they are life-saving [2]. However, urological FQs, including CIP, LEV, and MOX, present an important interindividual variability in PK associated with patient-specific characteristics. Thus, differences in interregional microorganism frequency, distribution, and resistance patterns could be encountered in clinical practice. This review highlights the need to take into account FQs' interindividual pharmacological variability to develop urological patient-tailored effective dosing strategies in order to improve microbiological and clinical outcomes, prevent the emergence of resistance, and minimize the incidence of adverse effects.

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