

Quinolone-Resistant *Haemophilus influenzae*: Determination of Mutant Selection Window for Ciprofloxacin, Garenoxacin, Levofloxacin, and Moxifloxacin

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Stepwise selection of ciprofloxacin-resistant *Haemophilus influenzae* mutants produced first-, second-, third-, and fourth-step substitutions in GyrA (S84Y), ParC (S84R), GyrA (D88N), and ParC (E88K), respectively. Successive mutations raised the mutant selection window. The wild-type selection window for garenoxacin, levofloxacin, and moxifloxacin was also measured.

Fluoroquinolone resistance in highly susceptible, gram-negative organisms is thought to arise in a stepwise manner through nucleotide sequence changes occurring largely at mutational hot spots (quinolone resistance-determining regions [QRDRs]) of genes encoding gyrase (*gyrA* and *gyrB*) and DNA topoisomerase IV (*parC* and *parE*) (7). Mutants at each step are enriched when fluoroquinolone concentrations are within a specific range called the mutant selection window (19). The lower boundary of the window is approximated by MIC₍₉₉₎, the minimal concentration that blocks growth of 99% of cells in a culture. The upper boundary is the MIC of the least susceptible, next-step mutant. Above this concentration two resistance mutations must be acquired concurrently for growth. Since this occurs rarely, the upper boundary of the window is called the mutant prevention concentration (MPC). To characterize stepwise development of fluoroquinolone resistance in vitro, we determined QRDR alterations and changes in fluoroquinolone susceptibility [MIC₍₉₉₎ and MPC] with *Haemophilus influenzae*.

Strain ATCC 49247 was grown on chocolate II agar (Becton Dickinson and Co., Cockeysville, Md.) or as liquid cultures in *Haemophilus* test medium broth (HTM broth; Becton Dickinson and Co., Sparks, Md.) at 37°C in 5% CO₂. Garenoxacin was obtained from Bristol-Myers Squibb (Wallingford, Conn.), moxifloxacin and ciprofloxacin were from Bayer Corp. (West Haven, Conn.), and levofloxacin was from R. W. Johnson Pharmaceutical Research Institute (Spring House, Pa.). Fluoroquinolone susceptibility [MIC₍₉₉₎] was determined by counting colonies following plating of serial dilutions on fluoroquinolone-containing agar (Fig. 1A and 2). MPC, which estimates the MIC of resistant mutant subpopulations, was defined as the fluoroquinolone concentration at which no colony was recovered when more than 10¹⁰ cells were applied to agar plates (Fig. 1A and 2). Plates were screened for colonies every 24 h during incubation for 96 to 120 h to assure that colony number had stabilized. Colonies obtained at high fluoroquinolone concentration were composed of resistant mutants, as confirmed by regrowth on drug-free agar followed by transfer and growth

on agar containing the fluoroquinolone concentration used for selection. Duplicate measurements gave similar results. Nucleotide sequence changes associated with loss of susceptibility were identified from regions of chromosomal DNA amplified by PCR as described previously (12).

Stepwise accumulation of fluoroquinolone resistance mutations was attained by sequential growth of laboratory strain ATCC 49247 on ciprofloxacin-containing agar. Application of wild-type cells to agar (Fig. 1A) allowed selection of a GyrA Ser-84-to-Tyr variant (Fig. 1A). One of these, strain KD2308, served as starter culture for a second round of selection. Colony recovery dropped sharply as ciprofloxacin concentration increased (Fig. 1A). Colonies recovered at concentrations indicated in Fig. 1A contained ParC variants in which Ser-84 changed to Arg. When the second-step *gyrA parC* mutant, strain KD2322, was applied to ciprofloxacin-containing agar, increasing drug concentration caused colony recovery to drop sharply, pass through an inflection, and drop a second time (Fig. 1A). Mutants recovered at concentrations indicated in Fig. 1A were GyrA variants in which Asp-88 changed to Asn. A third-step mutant, strain KD2364, was then used to select fourth-step mutants at ciprofloxacin concentrations indicated in Fig. 1A. Colonies obtained at the concentrations indicated in Fig. 1A contained a mutation that changed ParC position 88 from Glu to Lys. No mutation was observed in the QRDR of *gyrB* or *parE* (not shown). Each successive mutation raised the boundaries of the selection window (Fig. 1B). Each successive mutation also affected the composition of next-step mutant subpopulations, as indicated by the shapes of the recovery curves for the mutant (Fig. 1A).

When strain ATCC 49247 was applied to agar plates containing garenoxacin, levofloxacin, or moxifloxacin, colony recovery dropped with drug concentration as described for ciprofloxacin (Fig. 2). Two resistant mutants, recovered from agar plates containing high concentrations of each compound, had single nucleotide changes causing amino acid substitutions at position 84 or 88 of the GyrA protein (Table 1). Quinolones differed in selection of mutant subpopulations, as indicated by differences in shapes of mutant selection curves (Fig. 2).

GyrA and ParC changes have been observed with clinical isolates of *H. influenzae* (2–5, 8, 9, 12, 14, 17). GyrA mutants

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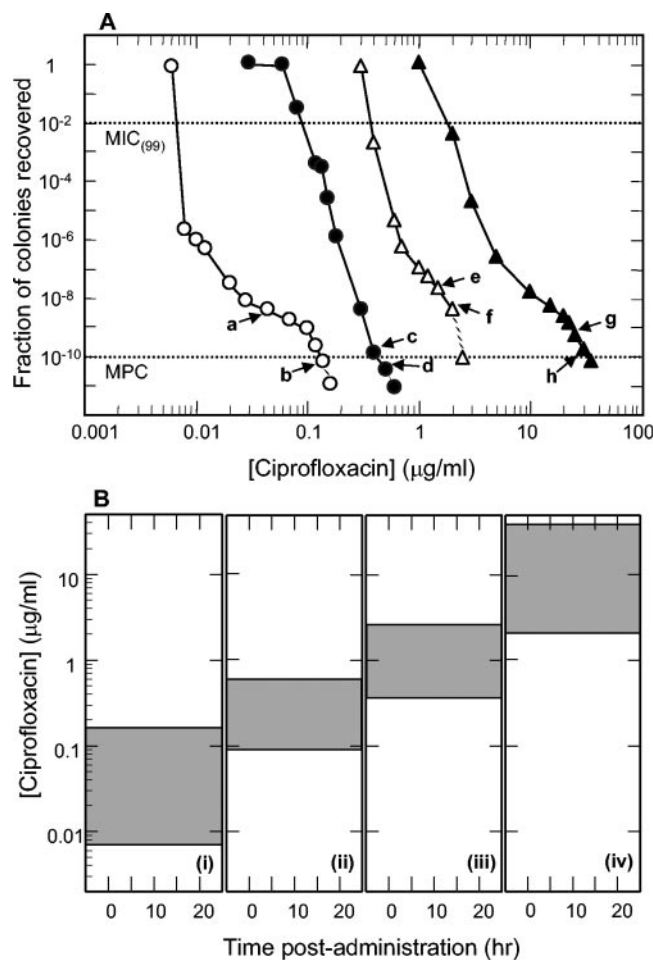


FIG. 1. Stepwise enrichment of ciprofloxacin-resistant *H. influenzae* mutants. (A) Effect of ciprofloxacin concentration on recovery of colonies. Wild-type *H. influenzae* ATCC 49247 (open circles) was applied to ciprofloxacin-containing agar at the indicated concentrations, and colonies were recovered. Arrows indicate ciprofloxacin concentrations used to obtain colonies for which nucleotide sequence information was obtained. Arrows a and b indicate recovery of Ser-84-to-Tyr GyrA variants. Strain KD2308 (arrow b) was used for second-round selection (solid circles). Cells from positions c and d contained an additional *parC* mutation that changed Ser-84 to Arg. Strain KD2322 (arrow d) was used for a third round (open triangles). Cells from positions e and f contained an additional *gyrA* mutation that changed Asp-88 to Asn. Strain KD2364 (arrow f) was used for a fourth round (solid triangles). Cells from positions g and h contained an additional *parC* mutation that changed Glu-88 to Lys. Dashed lines, drug concentrations at which no colony was recovered; dotted lines, guides for determining MIC₍₉₉₎ and MPC, as indicated. (B) Relationship of mutant selection window to serum drug concentration. Shaded areas represent mutant selection window defined by MIC₍₉₉₎ and MPC, which were obtained from data shown in panel A. (i) Wild-type cells; (ii) first-step mutant; (iii) second-step mutant; (iv) third-step mutant.

are probably the first to be enriched, as judged from the present *in vitro* studies (Table 1) and from the recovery of clinical isolates that contain only GyrA variants (2, 8, 9, 14). Since the patterns of topoisomerase changes reported for clinical isolates of *H. influenzae* (2, 5, 12) often differ from those found with the laboratory strain reported here, clinical isolates can follow a path to resistance different from that followed by the laboratory isolate.

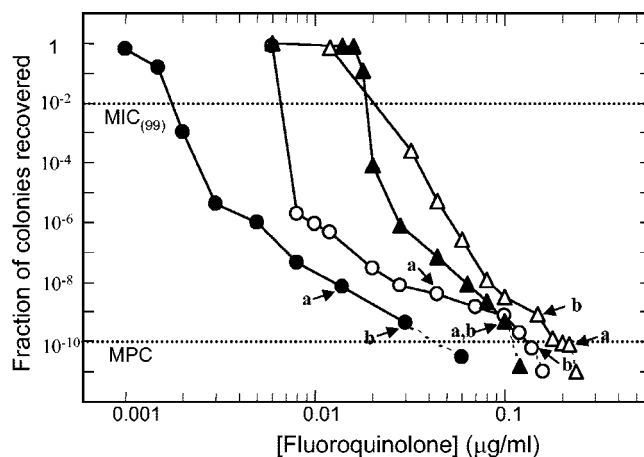


FIG. 2. Effect of fluoroquinolone concentration on the recovery of first-step, resistant mutants. Wild-type *H. influenzae* strain ATCC 49247 was applied to agar plates containing the indicated concentrations of garenoxacin (solid circles), ciprofloxacin (open circles), levofloxacin (solid triangles), and moxifloxacin (open triangles). After incubation for 5 to 6 days, colonies were counted, and the fraction recovered relative to the input CFU was determined. Dashed lines, concentrations at which no colony was recovered; dotted lines, MIC₍₉₉₎ and MPC, as indicated. Arrows a and b indicate conditions used to isolate colonies from which the DNA sequence was determined (see Table 1).

The selection window for ciprofloxacin with wild-type cells was below serum drug concentrations measured at steady state in human volunteers receiving twice-daily doses of 500 mg (10). Comparable statements can be made about the three other compounds examined; this is consistent with the prevalence of fluoroquinolone resistance being low with *H. influenzae* (3, 4). Resistance may arise largely from sporadic situations in which abnormally low doses (dosing errors) and/or patient characteristics, such as chronic lung disease (16), cause drug concentration to fall into the selection window and allow enrichment of spontaneous *gyrA* mutants. Since increasing the time that drug concentration is inside the selection window should increase mutant enrichment, the development of resistance is expected to accelerate with each successive mutation. A key to preventing fluoroquinolone resistance in *H. influenzae* may be to block enrichment of the first *gyrA* mutation by strictly avoiding low-dose fluoroquinolone use.

The progressive, stepwise acquisition of resistance alleles,

TABLE 1. Selection of resistant mutants by various quinolones

Quinolone	MIC ₍₉₉₎ ^a (mg/liter)	MPC (mg/liter)	GyrA change in QRDR ^b for mutant:	
			a	b
Garenoxacin	0.0018	0.06	S84 to Y	S84 to Y
Levofloxacin	0.019	0.12	S84 to Y	S84 to Y
Ciprofloxacin	0.007	0.16	S84 to Y	S84 to Y
Moxifloxacin	0.023	0.24	D88 to Y	S84 to Y

^a MIC using agar is 1.05 to 1.5 times higher than MIC₍₉₉₎ using agar (Fig. 2); broth MIC is higher than agar MIC by factors of 1, 2, and 3.7 for levofloxacin, ciprofloxacin, and moxifloxacin, respectively (15).

^b Quinolone concentrations used to obtain mutants are indicated in Fig. 2. Abbreviations: D, aspartic acid; S, serine; Y, tyrosine. First letter represents wild-type amino acid at the position indicated by the number. In each case the mutation changed the amino acid to Y.

which has been likened to hill climbing (1), is likely to be common to many bacterial species. For example, *gyrA* and *parC* mutations accumulate in *Neisseria gonorrhoeae* and *Escherichia coli*, albeit with the additional occurrence of drug efflux mutations (1, 11, 18). For some species, such as *Streptococcus pneumoniae*, only two steps appear to be required for resistance (13), and only a single gyrase mutation may be necessary for resistance with *Mycobacterium tuberculosis* (6). Slowing the development of fluoroquinolone resistance may require adjusting dosing strategies to take into account the number of mutational steps involved in resistance.

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