Fluoroquinolone-resistant Streptococcus pneumoniae

To the Editor: In pneumococci, resistance to fluoroquinolones is associated with chromosomal mutations in the quinolone-resistance–determining regions (QRDR) of type II topoisomerase enzymes, predominantly gyrA and parC. Several mutations have been described in these enzymes, but only a few have been shown by in vitro studies to confer resistance: S81F or Y, C, or I and E85K in gyrA; E474K in gyrB; A63T, S79F or Y or L and D83G or N in parC; and E474K and D435N or H in parE (1–5). Other frequently described mutations are K137N in parC and 1460V in parE, which appear to not contribute to fluoroquinolone resistance because they are commonly found in susceptible strains, and no evidence exists for their conferring fluoroquinolone resistance in vitro. We describe here a pneumococcal strain that was isolated from a 66-year-old white man with chronic obstructive pulmonary disease (COPD).

The patient was admitted to the hospital with a presumed exacerbation of COPD. He had been discharged from the hospital 2 days earlier, having recovered from a similar manifestation of this disease. His treatment history was 250 mg/day oral levofloxacin for 7 days while in the hospital and levofloxacin for 10 days as an outpatient for a similar lower respiratory tract infection 3 months earlier.

On this second admission he was given levofloxacin, 250 mg intravenously, once a day. He was treated with a low dosage because he was in renal failure. The patient continued to worsen and was transferred to the intensive care unit, where ceftriaxone, 1 g intravenously once a day, was given along with levofloxacin. He improved on the combination therapy and was discharged without sequelae.

Cultures of the patient’s blood and sputum grew Streptococcus pneumoniae. The isolate from blood was resistant to levofloxacin (MIC 8 mg/L) and ciprofloxacin (MIC 8 mg/L), yet susceptible to gatifloxacin (MIC 1 mg/L) and ceftriaxone (MIC 0.38 mg/L), with intermediate resistance to penicillin (MIC 1.5 mg/L). The resistant isolate was of serotype 6A and of multilocus sequence type 376, which is the North Carolina A-23 clone (http://www.sph.emory.edu/PMEN/index.html).

Efflux testing that compared the ciprofloxacin MICs in the presence and absence of reserpine (10 mg/L) showed no evidence of an overexpressed efflux pump. We sequenced the QDRDs (gyrA, gyrB, parC, parE) and the entire gyrA and parC genes of the resistant strain isolated from blood by using previously described primers (2). Sequencing showed a S79Y mutation in parC and a Q118K (CAA→AAA) mutation in gyrA. Sequencing of the entire gyrA and parC genes confirmed that no additional amino acid substitutions were outside the QRDRs. The entire gyrA gene PCR product was transformed directly into the susceptible pneumococcal reference strain R6 by a standard transformation protocol (4). Transformants were selected on plates containing increasing concentrations of ciprofloxacin and, in a second step, were transformed with the entire parC gene of the resistant strain.

The ciprofloxacin and levofloxacin MICs of R6 transformed with the gyrA gene of the resistant isolate containing the new Q118K mutation were 4 and 2 mg/L, respectively. After additional transformation of these transformants with parC of the resistant isolate containing the S79Y mutation, the selected double transformants exhibited the same MICs as the original clinical isolate (8 mg/L for ciprofloxacin and levofloxacin).

The transformation of parC alone conferred an intermediate increase in the MICs (ciprofloxacin 2 mg/L, levofloxacin 4 mg/L). All transformants were confirmed by sequencing.

To determine the biologic cost associated with the different resistance mutations in vitro, each fluoroquinolone-resistant mutant was competed against the fluoroquinolone-susceptible parent strain R6 (with an independent streptomycin resistance marker) as described by Johnson et al. (6). The outcome was evaluated as the change in the ratios of the competing strains as a function of the number of generations. Each competition was performed in triplicate by using independent starting cultures of each competing strain. Compared with the wild-type R6 strain, the relative fitness values for the gyrA, parC, and double mutants were 1.06, 1.03, and 0.93, respectively.

These data indicate that a single mutation in either parC or gyrA does not impose a substantial fitness burden. In contrast, the double-mutation parC S79Y and gyrA Q118K was associated with a slower growth rate. Similar results of relative fitness for single (parC S79Y and gyrA S81F) and double mutations were observed by Gillespie et al. (7).

Development of resistance to fluoroquinolones is a stepwise process, involving spontaneous mutations in the genes encoding the target enzymes DNA gyrase and the topoisomerase IV. Mutants with mutations in 1 of the enzymes are estimated to arise at a frequency of 1 to 10⁻⁷ (1). Therefore, fluoroquinolone resistance due to selection of spontaneous mutants during treatment may be related to the number of bacterial cells in the population under selective pressure. Patients with COPD are frequently colonized by high bacterial loads. COPD has been identified in several recent studies as an independent risk factor for fluoroquinolone resistance (8,9). Low doses of fluoroquinolones
may also lead to an increased risk for resistance selection (10). Because the Q118K mutation has not been previously described, this new mutation was probably selected by the current or antecedent treatments rather than by an infection with a resistant widely disseminated clone.

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Spring Scrub Typhus, People’s Republic of China

To the Editor: Pingtan Island, in the eastern Fujian Province, People’s Republic of China, has been a traditional focus of summer scrub typhus. In the early 1950s, the health of the residents was compromised by scrub typhus, with an incidence of 1,000 cases/100,000 population and a case-fatality rate of 13.6%. With the understanding of the pathogen and application of effective treatments (1,2), the epidemic was brought under control. Documentation showed that from 1960 through 1990, the annual incidence of scrub typhus maintained a level of 50–100 cases/year. Since 1990, cases have decreased sharply.

The usual epidemic season for scrub typhus on the island was summer. The first outbreak of spring scrub typhus occurred in 2000 in the town of Beicuo; 10 cases were reported. Beicuo, population 10,000, is located in southwestern Pingtan Island. The first patient visited the local hospital on April 6, 2000, with a high fever, cough, and headache. The initial exclusion of scrub typhus because of the spring time frame made the diagnosis difficult until a typical black eschar was found on the patient’s waist. This case called attention to spring scrub typhus and led to the timely diagnosis and treatment of the subsequent cases. By 2005, a total of 28 spring cases were on file. An indirect immunofluorescence antibody method with Gilliam strain antigen, as described (3,4), was applied to the above samples for serologic analysis. Samples with antibody titers >64 were considered diagnostic. All 28 case-patients were identified as having antibodies to Orientia tsutsugamushi 8–20 days after the onset of the disease (Table).

The number of cases of spring scrub typhus from 2000 to 2005 were