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DOI: 10.3201/eid1410.080545

References

- Price WH. The isolation of a new virus associated with respiratory clinical disease in humans. Proc Natl Acad Sci U S A. 1956;42:892–6. DOI: 10.1073/ pnas.42.12.892
- McErlean P, Shackelton LA, Lambert SB, Nissen MD, Sloots TP, Mackay IM. Characterisation of a newly identified human rhinovirus, HRV-QPM, discovered in infants with bronchiolitis. J Clin Virol. 2007;39:67–75. DOI: 10.1016/j. jcv.2007.03.012
- Kistler A, Avila PC, Rouskin S, Wang D, Ward T, Yagi S, et al. Pan-viral screening of respiratory tract infections in adults with and without asthma reveals unexpected human coronavirus and human rhinovirus diversity. J Infect Dis. 2007;196:817–25. DOI: 10.1086/520816
- Lau SK, Yip CC, Tsoi HW, Lee RA, So LY, Lau YL, et al. Clinical features and complete genome characterization of a distinct human rhinovirus (HRV) genetic cluster, probably representing a previously undetected HRV species, HRV-C, associated with acute respiratory illness in children. J Clin Microbiol. 2007;45:3655–64. DOI: 10.1128/JCM.01254-07
- Renwick N, Schweiger B, Kapoor V, Liu Z, Villari J, Bullmann R, et al. A recently identified rhinovirus genotype is associated with severe respiratory tract infection in children in Germany. J Infect Dis. 2007;196:1754–60. DOI: 10.1086/524312
- McErlean P, Shackelton LA, Andrews E, Webster DR, Lambert SB, Nissen MD, et al. Distinguishing molecular features and clinical characteristics of a putative new rhinovirus species, human rhinovirus C (HRV C). PLoS ONE. 2008;3: e1847. DOI: 10.1371/journal.pone.0001847
- Arden KE, McErlean P, Nissen MD, Sloots TP, Mackay IM. Frequent detection of human rhinoviruses, paramyxoviruses, coronaviruses, and bocavirus during acute respiratory tract infections. J Med Virol. 2006;78:1232–40. DOI: 10.1002/ jmv.20689

- Briese T, Renwick N, Venter M, Jarman RG, Ghosh D, Köndgen S, et al. Global distribution of novel rhinovirus genotype. Emerg Infect Dis. 2008;14:944–7.
- Savolainen C, Mulders MN, Hovi T. Phylogenetic analysis of rhinovirus isolates collected during successive epidemic seasons. Virus Res. 2002;85:41–6. DOI: 10.1016/S0168-1702(02)00016-3
- Turner RB, Couch RB. Rhinoviruses. In: Knipe DM, Howley PM, editors. Fields virology. 5th ed. Philadelphia: Lippincott, Williams & Wilkins; 2007. p. 903.

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Serogroup A Neisseria meningitidis with Reduced Susceptibility to Ciprofloxacin

To the Editor: Reduced susceptibility to ciprofloxacin of Neisseria meningitidis has been reported with increasing frequency since 1992, mainly because of mutations in the quinolone resistance determining regions (QRDRs) of the gyrase and topoisomerase IV genes (1,2). Reduced fluoroquinolone susceptibility due to gyrase A mutations in serogroup A strains has previously been reported from a 2005 outbreak in Delhi, India (1). We describe 2 clinical isolates of serogroup A N. meningitidis with reduced ciprofloxacin susceptibility that were recognized in March 2003 and April 2006 in Israel, a country with low incidence of invasive meningococcal disease (<2/100,000/laboratory-confirmed cases/year) in which this serogroup accounts for <2% of cases (data from the National Center for Meningococci, Tel Hashomer, Israel).

The 2 isolates in question (M12/03 and M24/06; suffixes denote year of isolation) were compared with 2 fully susceptible strains, M44/01 and M23/00 (online Appendix Table, available from www.cdc.gov/EID/ content/14/10/1667-appT.htm). MICs were measured by Etest (AB Biodisk, Solna, Sweden) on Mueller-Hinton agar (Difco Laboratories, Detroit, MI, USA) supplemented with 5% sheep blood. Demographic information was obtained from the Israel Ministry of Health Department of Epidemiology.

Chromosomal DNA was isolated by using the NucleoBond kit (Macherey-Nagel, Düren, Germany). The location of the QRDR in gyrase and topoisomerase IV genes was based upon prior studies in meningococci (online Appendix Table) and on the complete sequence of strain N. meningitidis Z2491 (serogroup A; Gen-Bank accession no. NC 003116). We amplified and sequenced extended regions encompassing the QRDRs by using the upstream and downstream primer pairs in gyrA (522 bases) 5'-GTTCCGCGTCAAAATATGCT-3', 5'-CCGAAATTGACGGTTTCTTC-3'; gyrB (649 bases) 5'-GGTTTGACC TGCGTGTTGTC-3', 5'-CGGCTGG GCGATATAGATG-3'; *parC* (635 5'-CACTATGGTTTGCCGT bases) TTTG-3', 5'-ATTTCGGACAACAG CAATTC-3'; and parE (610 bases) 5'-GGACAGGATGGCGATTTTG-3',5'-CGTCAGCAACTTCATCAACC-3'. PCR was performed by using Taq DNA polymerase (New England BioLabs, Beverly, MA, USA). DNA sequencing was performed using the ABI PRISM 3700 Genetic Analyzer (Applied Biosystems, Foster City, CA, USA). Screening for plasmidmediated quinolone resistance genes was carried out by multiplex PCR amplification of *qnrA*, *qnrB*, and *qnrS* as

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previously described (3). Multilocus sequence typing (MLST) was carried out by using the primers, protocols, and databases available from the Neisseria MLST website (http://pubmlst. org/neisseria) (4).

The online appendix Table shows our results and condenses previously published findings. The ciprofloxacin MIC for M24/06 was 42- to 125-fold higher than for susceptible strains and consistently 2-fold higher than that for M12/03. We have not referred to our isolates as resistant, because M12/03 would be categorized as "intermediate" by Clinical Laboratory Standards Institute breakpoints (5). The extended QRDRs in gyrA and parC of M44/01 (susceptible) were identical to those of N. meningitidis Z2491. M24/06 and M12/03 had a Thr91Ile mutation in gvrA. M24/06 also had Asn103Asp, Ile111Val, and Val120Ile mutations in gvrA (online Appendix Table; 1). In M12/03, an Ala78Val mutation was found in gyrA, and new mutations Ile474Leu and Thr365Ala were found in gyrB and parE, respectively. No parC mutations were found.

Previous reports identified chromosomal mutations in N. meningitidis (online Appendix Table). M24/06 and M12/03 possess the same Thr91Ile mutation in gyrA as a 2002 serogroup B isolate from Spain (online Appendix Table) that had a similar increase in ciprofloxacin MIC (0.12 mg/L). The Thr91Ile mutation is homologous with the Ser83Leu mutation in gyrA of Escherichia coli that is responsible for a 60-fold increase in ciprofloxacin MICs (6). Further mutations in a primary target enzyme (gyrase) have been associated with additional 2-fold increases in the MIC of ciprofloxacin (7). The level of resistance observed in M24/06 might suggest additional mechanisms. An efflux pump mechanism is unlikely; we showed no reduction in MICs in the presence of reserpine (online Appendix Table) and this organism was fully susceptible to penicillin, tetracycline, erythromycin, and Triton X-100 (data not shown). This finding suggests the absence of an efflux pump encoded by a mutated *mtrRCDE* (8). Neither M24/06 nor M12/03 had plasmid-mediated genes *qnr* genes or elevated kanamycin MICs, suggesting the presence of *aac(6')-Ib-cr*. Both of these genes can confer low-level quinolone-resistance and facilitate the emergence of higher level resistance (9) (data not shown).

MLST showed that M24/06 and M12/03 did not derive from a single clone after selection of the T91I mutation. M12/03 was sequence type (ST) 2 and was isolated from a recent immigrant from Russia, which is the origin of most ST 2 strains deposited in the Neisseria MLST database (29/34 records; 85%). M24/06 was ST4789 in the ST5 clonal complex, isolated from a person who had immigrated many years previously from Romania. ST4789 has been encountered only once previously, in Dhaka, Bangladesh.

Disease associated with serogroup A *N. meningitidis* has been extremely unusual in Israel (*10*) and has remained rare. This serogroup comprised only 9 (1.9%) of all 463 isolates submitted during 1997–2006 (data from the National Center for Meningococci).

The isolates described in our study confirm that serogroup A should be added to the list of meningococci with the potential for reduced fluoroquinolone susceptibility and raise the question why they have appeared in a region with particularly low serogroup A meningococcal disease incidence while frequently encountered serogroups have remained fully susceptible. The importance of continuous monitoring for reduced ciprofloxacin susceptibility in these more prevalent serogroups has been emphasized by the recent replacement of rifampin by ciprofloxacin as the preferred agent for chemoprophylaxis of meningococcal disease in adults in Israel.

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DOI: 10.3201/eid1410.080252

References

- Singhal S, Purnapatre KP, Kalia V, Dube S, Nair D, Deb M, et al. Ciprofloxacinresistant *Neisseria meningitidis*, Delhi, India. Emerg Infect Dis. 2007;13:1614–6.
- Centers for Diseases Control and Prevention. Emergence of fluoroquinolone-resistant *Neisseria meningitidis*—Minnesota and North Dakota, 2007–2008. MMWR Morb Mortal Wkly Rep. 2008;57:173–5.
- Robicsek A, Strahilevitz J, Sahm DF, Jacoby GA, Hooper DC. *qn*r Prevalence in ceftazidime-resistant *Enterobacteriaceae* isolates from the United States. Antimicrob Agents Chemother. 2006;50:2872–4. DOI: 10.1128/AAC.01647-05
- Jolley KA, Chan MS, Maiden MC. mlstdbNet–distributed multi-locus sequence typing (MLST) databases. BMC Bioinformatics. 2004;5:86. DOI: 10.1186/1471-2105-5-86
- Clinical Laboratory Standards Institute. Performance standards for antimicrobial susceptibility testing, 18th informational supplement. CLSI document M100-S18. Wayne (PA): The Institute; 2008.
- Hooper DC, Rubinstein E. Mechanisms of quinolone resistance. Quinolone antimicrobial agents. 3rd ed. Washington: American Society for Microbiology Press; 2003. p. 41.
- Bagel S, Hullen V, Wiedemann B, Heisig P. Impact of *gyrA* and *parC* mutations on quinolone resistance, doubling time, and supercoiling degree of *Escherichia coli*. Antimicrob Agents Chemother. 1999;43:868.
- Shafer WM, Veal WL, Lee EH, Zarantonelli L, Balthazar JT, Rouquette C. Genetic organization and regulation of antimicrobial efflux systems possessed by *Neisseria gonorrhoeae* and *Neisseria meningitidis*. J Mol Microbiol Biotechnol. 2001;3:219–24.
- Robicsek A, Strahilevitz J, Jacoby GA, Macielag M, Abbanat D, Hye PC, et al. Fluoroquinolone-modifying enzyme: a new adaptation of a common aminoglycoside acetyltransferase. Nat Med. 2006;12:83. DOI: 10.1038/nm1347

 Block C, Roitman M, Bogokowsky B, Meizlin S, Slater PE. Forty years of meningococcal disease in Israel: 1951– 1990. Clin Infect Dis. 1993;17:126–32.

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Identification of All Dengue Serotypes in Nepal

To the Editor: Nepal is situated on the southern slopes of the Himalayas, surrounded by India on 3 sides and China to the north. Nepal's altitude ranges from 8,848 m in the Himalayas to 90 m in the Terai, the southern, low, flatland bordering India. Nepal is a disease-endemic area for many vector-borne diseases, including malaria, kala-azar, Japanese encephalitis, and lymphatic filariasis. Because of the porous border between Nepal and India, social, cultural, and economic activities in cross-border areas are common.

Dengue is an emerging disease in Nepal; presumably transmission is moving north from India into the Terai (1-5). The first report of dengue virus isolation or RNA (serotype 2 with nucleotide homology closest to a dengue virus type 2 isolate from India) was in 2008 involving a Japanese patient returning from Nepal in October 2004 (5). Entomologic investigations from the 1980s showed *Aedes albopictus* in the Terai plains, but *Ae. aegypti* has not been previously reported.

After Indian outbreaks now known to include all 4 dengue serotypes (6), a team from the Epidemiology and Disease Control Division, Kathmandu, investigated suspected cases of dengue fever during September–October 2006 in Banke, the district bordering Uttar Pradesh, India. The team collected blood samples from persons in Banke and, subsequently, from persons in a number of other districts and sent them to the National Public Health Laboratory in Kathmandu or the Armed Forces Research Institute of Medical Sciences, Bangkok, Thailand for analysis with ELISA, reverse transcription–PCR, (RT-PCR), or both.

Case definitions for dengue fever were adopted based on World Health Organization guidelines (7). Blood samples were obtained from patients with an acute febrile illness of 2-7 days' duration and with >2 of the following manifestations: headache, retro-orbital pain, muscular or joint pain, and rash. If laboratory tests were positive, cases were confirmed. Results were confirmed by ELISA performed at the Armed Forces Research Institute of Medical Sciences as previously described (8). Positive results were immunoglobulin (Ig) M ≥40 units or IgG >100 units. RT-PCR was performed by extracting RNA from 140 µL of each serum sample using QIAGEN Viral RNA Extraction Kit per manufacturer's instructions (QIA-GEN, Germantown, MD, USA). RT-PCR and nested PCR were conducted according to the Lanciotti protocol (9) with the following modifications. Reverse transcriptase from avian myeloblastosis virus (Promega, Madison, WI, USA) was used in the first round RT-PCR. The concentrations of the primers used in the RT-PCR and nested reactions were reduced from 50 pmol to 12.5 pmol per reaction, and the number of nested PCR amplification cycles was increased to 25.

Serum specimens were obtained from 70 suspected case-patients from 16 districts from October 13 through December 3, 2006; 25 confirmed cases (13 by ELISA, 10 by RT-PCR, and 2 by both tests) came from 9 districts (Table). The average age was 29 years (range 5–65 years); 80% of the casepatients were men. Three patients had a history of travel to India, but clusters of dengue fever cases reported in October (Banke and Dang districts) indicated local transmission was occurring among patients with no travel history. The Terai districts accounted for 80% of cases. Entomologic collections done indoors and outside at 5 different sites reporting suspected cases identified *Ae. albopictus* and *Ae. aegypti* in all 5 districts.

These clinical and laboratory test results confirmed the presence of all 4 dengue serotypes. Notably, patients from the Dang district had no travel history outside the Dang valley. Because Aedes spp. have been identified in Dang, the data strongly suggest the existence of an endemic cycle of dengue. Underreporting is expected in the absence of diagnostic facilities at the field level. It is unclear whether the predominance of male patients is indicative of greater outdoor as opposed to indoor transmission. Of note, Ae. albopictus has been found in the country since the 1980s; in this study, we found Ae. aegypti in Nepal. Men typically wear short-sleeved clothes due to hot and humid conditions and, therefore, are frequently exposed to mosquito bites. However, men may also access the healthcare system more frequently. The ages of case-patients point to a relative lack of dengue immunity among the older population, and this finding is consistent with a new introduction of dengue. Because dengue hemorrhagic fever appears when >1 serotype becomes endemic to an area (10), the presence of all 4 serotypes portends the emergence of more severe dengue disease in Nepal.

Acknowledgments

We thank laboratory personnel at the National Public Health Laboratory and the Armed Forces Research Institute of Medical Sciences for their expertise.

Publisher: CDC; Journal: Emerging Infectious Diseases Article Type: Letter; Volume: 14; Issue: 10; Year: 2008; Article ID: 08-0252 DOI: 10.3201/eid1410.080252; TOC Head: Letter

Appendix Table. Activity of ciprofloxacin against serogroup A Neisseria meningitidis with reduced and full susceptibility to fluoroquinolones and comparison of mechanisms associated with elevated MICs reported in N. meningitidis

Location (isolates)	Serogroup	Source*	Ciprofloxacin MIC. mg/L	gyrA mutation	<i>parC</i> mutation	gyrB mutation	parE mutation
France 1996-98†	В	STD clinic	0.125	Asp-95→Glv	None		
Australia 1998‡	С	Blood, CSF	0.25	Asp-95→Asn	None		
Spain 2002§	В	CSF	0.12	Thr-91→lle	None		
Argentina 2002¶	Y	CSF	0.12	None	None	None	None
Argentina 2002¶	В	Blood, CSF	0.06	Asp-95→Asn	None		
Spain 2007#**	Nongroupable	Sputum	0.25	Asp-95→Asn	None		His-495→Asn
				Asn-103→Asp lle-			
				111→Val			
				Val-120→Ile			
India 2005 (1)	A (12 outbreak			Thr-91→lle			
	isolates)			Asn-103 \rightarrow Asp			
				lle-111→Val			
				Val-120 \rightarrow lle			
USA 2007–08 (<i>2</i>)	B (3 cases)	CSF	0.19	Thr-91→lle			
Israel 2001	Α	Blood	0.001	None	None	None	None
(M44/01)††							
Israel 2000	A	Blood	0.003	None	None		
(M23/00)							
Israel 2003	A	Blood, CSF	0.064	Thr-91→lle	None	lle-474→Leu	Thr-365→Ala
(M12/03)				Ala-78→Val			
Israel 2006	A	Blood	0.125	Thr-91→lle	None	None	None
(M24/06)				Asn-103→Asp lle-			
				111→Val			
				Val-120→lle			

*STD, sexually transmitted disease; CSF, cerebrospinal fluid.

+Casin I, Gandry B, Lassau F, Janier M, Lagrange P, Collatz E. Decreased susceptibility to penicillin G (Pen), tetracyclines (Tet), and fluoroquinolones (Fq), and characterization of DNA gyrase mutations, in oropharyngeal and anogenital isolates of Neisseria meningitidis (Nme) in patients presenting at an STD clinic. In: Program and Abstracts of the 39th Interscience Conference on Antimicrobial Agents and Chemotherapy; San Francisco, CA; 1999 September 26-29; Abstract 2101. Washington: American Society for Microbiology; 1999. ‡Shultz TR, Tapsall JW, White PA, Newton PJ. An invasive isolate of *Neisseria meningitidis* showing decreased susceptibility to quinolones.

Antimicrob Agents Chemother. 2000;44:1116.

§Alcala B, Salcedo C, de la Fuente L, Arreaza L, Uria MJ, Abad R, et al. Neisseria meningitidis showing decreased susceptibility to ciprofloxacin: first report in Spain. J Antimicrob Chemother. 2004;53:409.

Corso A, Faccone D, Miranda M, Rodriguez M, Regueira M, Carranza C, et al. Emergence of Neisseria meningitidis with decreased susceptibility to ciprofloxacin in Argentina. J Antimicrob Chemother. 2005;55:596-7.

#Enriquez R, Abad R, Salcedo C, Perez S, Vazquez JA. Fluoroquinolone resistance in Neisseria meningitidis in Spain. J Antimicrob Chemother. 2008;61:286-90.

**Nine additional strains reported; shown is a strain with mutations in gyrA similar to those found in isolate in present study.

††Identifiers of the National Center for Meningococci.