ETX-M–Type β-Lactamases Affect Community Escherichia coli Treatment, Greece

To the Editor: In recent years, a new group of extended-spectrum β-lactamases (ESBLs), the CTX-M–type enzymes, has emerged among Enterobacteriaceae (1). These enzymes are much more active against cefotaxime than against ceftazidime and exhibit greater susceptibility to tazobactam than to clavulanate. Currently, the CTX-M family includes at least 30 alleles, which may be clustered on the basis of sequence similarity into four major evolutionary lineages (2). These enzymes were first detected in South America, Germany, and France, and subsequent reports found them to be in several European countries as well as in the Far East and North and South America (1–5). All of these studies have described the production of CTX-M–type β-lactamases in Enterobacteriaceae that have been isolated from patients with hospital infections, whereas surveys assessing countrywide prevalence of CTX-M clusters in community-acquired infections have not been carried out. We report the dissemination of various CTX-M–type enzymes in clinical isolates of Escherichia coli recovered from community-acquired infections in two large regions of Greece.

The microbiology databases of two healthcare systems in Greece that serve as tertiary care centers for their region (University Hospital of Larissa and Hippokration University Hospital of Thessaloniki) were prospectively searched from January to September 2003. From almost 75,000 outpatient visits, we tested E. coli isolates that were recovered from patients with community-acquired infections and classified as ESBL producers by the E-test ESBL screen method with cefotaxime and ceftazidime plus clavulanate. Community-acquired E. coli infections were defined as those contracted outside a hospital environment for persons with no history of hospitalization, surgery, or outpatient care during the previous 30 days.

ESBL-positive isolates for which the cefotaxime MICs were at least eightfold higher than those of ceftazidime by agar dilution were saved and stored at –70°C. Results of antimicrobial susceptibility tests and ESBL screening and confirmatory tests were used to characterize the phenotypes of the isolates. Molecular analysis included polymerase chain reaction (PCR) detection with primers producing an 873-bp amplicon of the blaCTX-M gene (6), sequencing on both strands of PCR products, pulsed-field gel electrophoresis of XbaI chromosomal digests, plasmid analysis, and transferability experiments of cefotaxime resistance.

Treatment outcomes were assessed by reviewing the medical records of all patients for whom a culture yielded a strain of E. coli producing a CTX-M–type ESBL. Patients’ data included demographic details, presence of existing illness, symptoms, laboratory test results, history of surgery, and exposure to extended-spectrum cephalosporins <30 days before the positive culture. The antimicrobial treatment regimen was recorded, including the agent or agents administered, the duration of treatment, and clinical response.

During the study period, 14 community-acquired E. coli isolates (10 in the region of Larissa and 4 in the region of Thessaloniki) were recovered; the E-test ESBL screen test confirmed that these isolates were ESBL producers, and cefotaxime MICs were at least eightfold higher than those of ceftazidime. A CTX-M–type determinant was detected by PCR in 10 isolates (6 from Larissa and 4 from Thessaloniki). Sequencing their amplicons revealed that three of them were CTX-M-1 producers, and seven were CTX-M-3 producers. Genotyping showed that all of these isolates were unrelated. MICs of cefotaxime were always >128 µg/mL, whereas MICs of ceftazidime ranged from 0.5 µg/mL to 16 µg/mL. All isolates were sensitive to cefoxitin and piperacillin/tazobactam. However, the two laboratories reported them as being ESBL producers and recommended that β-lactam antibiotics, with the exception of carbapenems, not be used in their treatment. Several isolates exhibited additional resistance to co-trimoxazole, ciprofloxacin, tetracycline, or gentamicin, but all were susceptible to the other aminoglycosides. The blaCTX-M determinants were transferable to E. coli by conjugation in all but one case, along with other antimicrobial resistance determinants, with transfer frequencies that ranged from 8.3 x 10⁻² to 2.2 x 10⁻². Transconjugants contained one to three plasmids of varying size, ranging from 10 to 130 kb.

CTX-M–positive isolates were recovered from five children and five adults. Patients did not have typical risk factors, except for a chronic hematologic malignancy in one patient. None of the patients had severe urinary tract infections and received courses of amikacin or ciprofloxacin. One female patient had a purulent perianal infection. Results of blood and wound cultures from samples obtained at admission yielded the CTX-M–positive E. coli strain. She was given amikacin and clindamycin, and her condition gradually improved.

β-lactam antibiotics are the most common antimicrobial agents used in the community setting. The documented CTX-M–positive isolates exhibited plasmid-mediated resistance that affected the antimicrobial activity of all penicillins and cephalosporins as well as of several alternative antimicrobial agents used to treat community-acquired E. coli
infections. The spread of CTX-M–positive bacteria considerably changes the way we think about treating community-acquired infections and limits the oral antibiotics that may be administered. This finding has major implications for treating children, who should not be given fluoroquinolones and tetracyclines.

The observation that different bla_{CTX-M} alleles, located on plasmids of different sizes, were involved in clinical infections caused by distinct E. coli clones implies that CTX-M enzymes may become widespread in the community. A possible association of bla_{CTX-M} genes with insertion sequences like ISEcp1B might have contributed to the enhanced expression and mobilization of bla_{CTX-M} genes among E. coli isolates (7). The apparent dissemination of CTX-M producers could represent a substantial barrier in the treatment of community-acquired infections. Additionally, severely ill patients treated in the outpatient setting may transmit such resistant organisms to hospitalized patients.

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LETTERS

Age and Transmissible Spongiform Encephalopathies

To the Editor: Bacchetti (1) notes “Our findings suggest that the possibility should not be discounted that biological factors peaking in the third decade of life may promote variant Creutzfeldt-Jakob disease (vCJD) prion replication and consequent development of disease.” Such age specificity of disease risk may be a general feature of transmissible spongiform encephalopathies, which suggests that a general mechanism should be sought. A likely candidate for this mechanism is senescence-related immune system defects.

In a study of scrapie outbreaks in four sheep flocks, the incidence of clinical cases peaked in sheep 2–3 years of age, despite very different forces–of– infection at work and very large differences in disease incidence (2). Similar age specificity has been observed in cattle infected with bovine spongiform encephalopathy (3), which is believed to be the causal agent of variant Creutzfeldt-Jakob disease. There is evidence that an age-specific peak in prevalence also occurs in chronic wasting disease, a laterally transmitted spongiform encephalopathy of North American cervids, specifically elk, mule deer, and white-tailed deer. For example, data on prevalence of chronic wasting disease in mule deer (Figures 4B and 4A of [4]) suggest the existence of age-specific peaks. In aggregate, these observations suggest that a general mechanism might produce the marked decline in disease risk as age increases.

In 1979, Dickinson and Outram (5) conjectured that, in some experiments, scrapie responsiveness is the opposite of what one normally expects with an infection, “raising the possibility that, far from being inimical, some part of the host’s immune system is essential and may even play the role of a Trojan Horse for these agents when infection occurs by a peripheral route.” This theory appears well founded for transmissible spongiform encephalopathies in general. Disease-associated forms of resistant prion protein (PrP^{Res}) are likely transported from the gut to lymphoid tissue by cells such as migrating intestinal dendritic cells (6). Once in the lymphoid tissue PrP^{Res} appears to be amplified by follicular dendritic cells (6) and then enters the nervous system. Defects in either the complement pathway or follicular dendritic cells result in resistance to peripheral scrapie infection (7,8), and this resistance likely occurs for peripheral transmissible spongiform encephalopathy infections in general.