the sample was deposited on an agar plate flooded with a suspension of a antimicrobial susceptible strain of Micrococcus luteus. After 24 h of incubation at 37°C, presence of antimicrobial activity in the sample was evident by a visible area of growth inhibition of M. luteus around the sample. Procedures for DNA extraction and 16S rRNA gene amplification and sequencing have been detailed (4).

Gram staining of the sample showed numerous polymorphonuclear leukocytes and gram-negative bacteria. Culture of the sample remained sterile after 20 days of incubation, and antimicrobial susceptibility was found. The 16S rRNA gene amplification and sequencing determined a 1,493 nucleotide sequence. This sequence had 99.7% nucleotide similarity with that of L. amnionii (GenBank accession no. AY078425), which corresponded to a difference of 4 nucleotide. The 16S rRNA gene sequence of the detected bacterium was deposited under accession no. AY489565. L. amnionii was previously recovered in anaerobic culture of the amniotic fluid of a woman after intrauterine fetal demise (2). It was isolated on blood and chocolate agar under anaerobic conditions and showed very small gray colonies of <1 mm. This slow-growing bacterium was lost after two subcultures, and no isolate is available for further description (2). In that case and in the case reported here, the patients had uneventful recoveries after an amoxicillin plus clavulanic acid antimicrobial regimen was given. This bacterium and our isolate are related to, but different from, L. sanguinegens.

Leptotrichia is a small genus closely related to Fusobacterium and comprises slow-growing, gram-negative, filamentous, anaerobic bacterial flora of the oral cavity and genital tract (5). Species included in the genus are L. buccalis, L. trevisanii, L. sanguinegens, and L. amnionii (2,6).

All Leptotrichia species are extremely fastidious and cannot be grown easily on conventional microbiologic media or by conventional methods. As evidenced by sequences available in the GenBank database, most of the 16S rRNA gene sequences are from cloned DNA from complex flora but not from bacterial isolates. Leptotrichia species have been suspected to play a role in periodontal disease. However, Leptotrichia species have only been associated with serious systemic disease, usually in immunocompromised patients (7,8). Bacteremia caused by L. sanguinegens in pregnant women has also been reported (9). More widespread use of polymerase chain reaction amplification and sequencing of the 16S rRNA gene for identification or detection of fastidious pathogens in humans will likely provide verification of several new pathogens that are now part of normal human flora.

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Cholera in Mozambique, Variant of Vibrio cholerae

To the Editor: Cholera outbreaks caused by toxigenic Vibrio cholerae serogroup O1 frequently occur in many sub-Saharan African countries. The serogroup O1 is classified into two biotypes, classical and El Tor. The seventh and current pandemic of cholera is caused by the El Tor biotype; the classical biotype is believed to be extinct. The classical and El Tor biotypes of V. cholerae O1 are closely related in their O-antigen biosynthetic genes but differ in other regions of the genome. The genomic structure of the CTXΦ filamentous phage (1), in which the cholera toxin genes are contained, differs between the classical and El Tor biotypes. CTXclassΦ is found in classical strains, CTXElTorΦ is
present in El Tor and O139 strains, and CTXΦ is found in resurgent O139 strains. The diversity of CTXΦ among biotypes is mainly due to the variations in the repeat sequence elements, particularly in the rstR gene region (2).

While conducting surveillance in the cholera treatment center in Beira, the second largest city in Mozambique, we examined 175 rectal swabs or stool samples from January 7 to March 8, 2004, using standard published procedures. During this period, we isolated 58 strains of V. cholerae O1. The isolates were transported to the Enteric Microbiology Unit of the International Center for Diarrhoeal Disease Research in Dhaka, Bangladesh (ICDDR,B), for further phenotypic and genotypic characterization to determine serotype, biotype, and presence of important virulence genes. All 58 strains were identified as V. cholerae O1 of the Ogawa serotype. Forty strains selected for detailed characterization were resistant to polymyxin B, agglutinated chicken cells, yielded a positive Voges-Proskauer reaction, were positive for the El Tor hemolysin by the tube agglutination method, and were sensitive to group IV El Tor antibiotics which stimulate antitoxic antibodies and the vibriostatic compound 0/129.

By using polymerase chain reaction (PCR), we established that all 40 strains carried the ctxA gene (constituent gene of the CTX prophage) and the tcpA gene (the El Tor type), a constituent gene of the vibrio pathogenicity island. We then focused on the rstR gene because of its diversity between the two biotypes. All of the 40 El Tor strains produced a 500-bp PCR product of the rstR gene of the classical type (rstRclass), despite belonging to the El Tor biotype. Nucleotide sequence analysis of the rstR gene of two representative Mozambique strains showed 100% homology to the classical rstR gene of classical reference strain O395. The amino acid sequence of the B-subunit of classical and El Tor biotypes have identical to those of the classical CT-B subunit of the Mozambique strains varied from the El Tor CT-B subunit at positions 39 (histidine replaces tyrosine in El Tor), and 68 (threonine replaces isoleucine in El Tor), and the amino acid residues at these positions are identical to those of the classical CT-B subunit (Figure). The nucleotide sequences obtained for ctxB of the two Mozambique strains B33 and B65 were deposited in GenBank under accession no. AY648939 and AY648940, respectively. Therefore, the Mozambique strains of V. cholerae O1 displayed typical traits of the El Tor biotype overall but carried the classical CTX prophage.

Our findings that El Tor strains of V. cholerae O1 from Mozambique are carrying the classical prophage shows the presence of genetic materials associated with the classical biotype in Mozambique. Further, these findings provide the first circumstantial evidence of transmission of the classical CTXΦ prophage. The CTX prophages in El Tor strains give rise to infectious phage particles (1), but neither of the two CTX prophages integrated at two different sites of the classical genome give rise to phage particles (4). Subsequent studies have shown that, although the genes of the classical prophages encode functional forms of all of the proteins needed for production of CTXΦ, the CTX prophage does not yield virions because of the atypical arrangement of its prophage arrays (4).

Genetic hybrids between El Tor and classical biotypes of O1 V. cholerae were reported among sporadic isolates earlier in Bangladesh (5) and were named the Matlab variants after the place where they were first isolated. The Mozambique strains of V. cholerae likely evolved from an El Tor strain, which shed its CTX phage and acquired the classical prophage. Alternatively, strains like the Matlab variant may have spread to the African continent. Whether introducing the CTX prophage in the El Tor genome background will increase pathogenicity, affect genomic stability, or enhance the epidemic-causing potential is uncertain. This subtle genetic change might also alter the effectiveness of current cholera vaccines which stimulate antitoxic as well as antibacterial immunity.
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Correction, vol. 10, no. 10

In "Scrub Typhus in the Republic of Palau, Micronesia," by A. Mark Durand et al. (p. 1840), Table 2 was incorrect. The correct version appears below and online at http://www.cdc.gov/ncidod/eid/vol10no10/04-0288.htm#table2. We regret any confusion this error may have caused.

Table 2. Orientia tsutsugamushi IgG and IgM antibody titers for six southwest islanders with prolonged fever and abdominal distressa

| Patient no. | Antibody type | Acute-phase titer | Convalescent-phase titer |
|-------------|---------------|------------------|--------------------------|
| 1           | IgG           | 1:2,048          | NA                       |
|             | IgM           | 1:16,384         | NA                       |
| 2           | IgG           | NA               | 1:32,768                 |
|             | IgM           | NA               | 1:2,048                  |
| 3           | IgG           | 1:262,144        | 1:262,144                |
|             | IgM           | 1:4,006          | 1:1,024                  |
| 4           | IgG           | 1:65,536         | 1:65,536                 |
|             | IgM           | 1:1,024          | 1:2,048                  |
| 5           | IgG           | 1:8,000          | 1:64,000                 |
|             | IgM           | NA               | NA                       |
| 6           | IgG           | 1:4,000          | 1:64,000                 |
|             | IgM           | NA               | NA                       |

Ig, immunoglobulin; NA, result not available.

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