identified mainly carbapenem-hydrolyzing carbapenemase OXA-58 (9).

Recently, we showed that the \( \text{bla}_{\text{NDM-1}} \) gene was carried by a composite transposon bracketed by 2 copies of \( \text{IS}A_{\text{ba125}} \) in \( A. \text{baumannii} \) (10). Cloning and sequencing of the genetic context of the \( \text{bla}_{\text{NDM-1}} \) in the first isolate showed that transposon \( \text{Tn125} \) was truncated at its 3'-end extremity by insertion sequence \( \text{IS}A_{\text{ba14}} \), giving rise to a truncated \( \text{Tn125} \) (\( \Delta \text{Tn125} \)). PCR mapping of all isolates showed that they possessed this truncated isoform of \( \text{Tn125} \), which was therefore probably no longer functional.

The identification of several clinical \( A. \text{baumannii} \) isolates that possessed the \( \text{bla}_{\text{NDM-1}} \) gene and originated from North Africa, with no obvious link to the Indian subcontinent, strongly suggests that 1 NDM-producing \( A. \text{baumannii} \) clone is probably widespread in North Africa and that it might now act as a reservoir for NDM-1. This finding might indicate that control of spread of multidrug-resistant \( A. \text{baumannii} \) would have a primary role in controlling spread of NDM-1.

This work was funded by a grant from the Institut National de la Santé et de la Recherche Médicale.

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DOI: http://dx.doi.org/10.3201/eid1905.121618

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Genomic Analysis of Salmonella enterica Serovar Typhimurium Definitive Phage Type 104

To the Editor: \( \text{Salmonella enterica} \) is among the leading causes of foodborne diseases worldwide. Multidrug-resistant \( S. \text{enterica} \) serovar Typhimurium definitive phage type (DT) 104 emerged during the early 1990s in the United Kingdom and spread worldwide thereafter (1). This phage-type strain harbors a chromosomally encoded genomic island, Salmonella Genomic Island 1, which is typically responsible for resistance to ampicillin, chloramphenicol,
streptomycin, sulfonamide, and tetracycline (2). Multilocus variable-number tandem-repeat analysis (MLVA) is an established molecular epidemiologic tool; its high-resolution power has been applied to the subtyping of a variety of bacterial species (3). An MLVA system has been developed for analyzing *S. enterica* serovar Typhimurium (4,5).

The design of an MLVA system relies on the analyzed genome sequences. In this study, we found and evaluated a variable-number tandem-repeat region, or locus, designated DT104o. The locus is specific to *S. enterica* ser. Typhimurium DT104, according to the sequence of NCTC 13348 (available from the Sanger Institute, http://www.sanger.ac.uk/resources/downloads/bacteria/salmonella.html). The repeat unit sequence was CTCAAGAA/TTCTGAG, spanning 1952121–1952274 on the reference genome or 22 repeats of 7 nt.

We used 266 apparently independent isolates of *S. enterica* serovar Typhimurium collected during 1981–2012; 103 were from human samples and 163 from non-human sources. Bacteriophage typing was performed according to Anderson’s method and scheme (6). Types of 100 isolates were in the DT104 group, comprising DT104, DT104B, and U302, the latter being related to DT104 (2); MLVA was performed by using the 5 loci (STTR3, STTR5, STTR6, STTR9, and STTR10) with slight modifications (4,5). The DT104o locus was tested by using primers o-for (5′-GTCAACATGACTGCCCCTCA-3′), labeled with NED, and o-rev (5′-TTTGCTCTTGCTCTTAGCAATC-3′); this spanned 1952367–1952043 on the reference sequence, resulting in a 325-bp product with 171-bp offset.

For all 266 isolates tested, the number of alleles and the Simpson’s index of diversity score (D) identified in each locus are summarized in the Table. The 5 common and DT104o loci displayed high discriminatory power: DT104o was specific for the DT104 group, and all 100 DT104 group isolates displayed amplified products with 13–40 repeat copy numbers; the others showed the null allele. Focusing only on the 100 DT104 group isolates, the discriminatory power of STTR9 and STTR3 were poor, whereas STTR5, STTR6, STTR10, and DT104o displayed high discriminatory powers (Table). In addition, using the 5 common loci (MLVA5) in analysis, we identified 66 types with a D value of 0.974; use of MLVA5 plus the DT104o locus (MLVA6) identified 83 types with a D value of 0.984. These results indicate that the DT104o locus is highly specific and therefore useful as an additional molecular epidemiologic marker for analyzing *S. enterica* ser. Typhimurium DT104.

Because DT104o was highly variable, 5 DT104 strains were tested for the frequency of variants at each locus after 5 serial passages by using liquid culture: cultures were diluted 1:1,000 at each passage. Sixteen colonies of each strain were tested by using MLVA6 (online Technical Appendix Table 1, wwwnc.cdc.gov/EID/article/19/5/12-1395-techapp1.pdf). No variants were observed in STTR3, STTR9, or STTR10. STTR5, STTR6, and DT104o each showed 1 variant of 80 colonies. The results suggest that DT104o would not be less stable than other loci.

We also found that DT104o could provide more discriminatory power to MLVA5 in some settings (online Technical Appendix Table 2). We compared 2 settings using isolates from non-human samples. Setting 1 comprised isolates 1a and 1b from an outbreak during 1996 and isolate 1c in 2007. Isolates 1a and 1b were identical by MLVA6. Isolate 1c was identical by MLVA5 but not by MLVA6. In Setting 2, three isolates obtained in different years also were identical by MLVA5, but differed from each other by MLVA6. This suggests that MLVA6 could be useful in some epidemiologic settings such as in an outbreak investigation, though more extensive study would be required to confirm this suggestion.

The DT104o locus is located at the proximal region of fragment 180 comprised of a prophage structure, which was proven to be DT104-specific in a previous study (7). This finding is consistent with the results of our study.

In conclusion, development of an MLVA system is dependent upon the genome sequences available, and the system is usually used for molecular subtyping of a certain serotype in a particular organism. However, a specific group of strains could cause a pandemic and become a target of public health concern, as was *S. enterica* ser. Typhimurium DT104. The MLVA system could be improved by adding loci based on the genome sequence of such pandemic strains. In this study, we showed that the newly identified DT104o locus could be useful in identification and subtyping of *S. enterica* ser. Typhimurium DT104.

**Acknowledgment**

We thank the staff at all the municipal and prefectural public health institutes of Japan for providing the isolates. We also thank Nobuko Takai and Jiyoung Lee for their technical assistance.
This study was partly supported by grants-in-aid from the Ministry of Health, Labour and Welfare of Japan (H22-Shokuhin-Ippan-012, H23-Shinko-Shitei-020, H24-Shokuhin-Ippan-007, and H24-Shokuhin-Ippan-008).

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DOI: http://dx.doi.org/10.3201/eid1905.121395

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Single Genotype of Anaplasma phagocytophilum identified from Ticks, Camargue, France

To the Editor: Granulocytic anaplasmosis is a tickborne zoonosis caused by Anaplasma phagocytophilum bacteria, which are emerging in Europe. Besides infecting humans, A. phagocytophilum infect a wide range of wild and domestic mammals (1). In Europe, the Ixodes ricinus tick is the main vector for the bacteria, but A. phagocytophilum has also been detected in association with Rhipicephalus and Dermacentor spp. ticks (2). The climate and biotopes of the Mediterranean region are particularly favorable for several species of ticks and, therefore, for tickborne diseases.

Although I. ricinus ticks are rare or absent in the Mediterranean Basin, serosurveys performed on equine populations in Camargue, southern France, indicated an A. phagocytophilum seroprevalence of ≈10% (3). To investigate the prevalence and diversity of A. phagocytophilum bacteria in ticks in Camargue, we collected questing ticks from horse pastures and feeding ticks from horses.

Ticks feeding on horses were collected in randomly selected stables during 2007 (84 stables), 2008 (72 stables), and 2010 (19 stables). The stables were chosen among those where evidence of A. phagocytophilum seroconversion in horses had previously been found (3). In 2008 and 2010, questing ticks were collected by the dragging method in 19 pastures, around bushes, and in areas where horses spent the most time. Surveys were conducted in the spring, which represents the peak activity time of Ixodes ticks.

A total of 406 adult ticks were collected, representing 6 species: Rhipicephalus bursa, R. sanguineus, R. turanicus, R. pusillus, Dermacentor marginatus, and Hyalomma marginatum. Tick species were identified by morphologic criteria and molecular analyses based on mitochondrial 12S rDNA sequences (4). Total DNA was extracted from the ticks by using the NucleoSpin Tissue Kit (Macherey-Nagel, Düren, Germany) (5).

A. phagocytophilum was detected by nested PCR targeting the 16S rDNA (online Technical Appendix 1, wwwnc.cdc.gov/EID/article/19/5/12-1003-Techapp2.pdf). The prevalence of A. phagocytophilum infection did not differ significantly between species (logistic regression model, p = 0.76) but was higher among questing ticks than feeding ticks (p<0.001; odds ratio 1.15).