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## Whole-Genome Multilocus Sequence Typing of Extended-Spectrum-Beta-Lactamase-Producing *Enterobacteriaceae*

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Molecular typing has become indispensable in the detection of nosocomial transmission of bacterial pathogens and the identification of sources and routes of transmission in outbreak settings, but current methods are labor-intensive, are difficult to standardize, or have limited resolution. Whole-genome multilocus sequence typing (wgMLST) has emerged as a whole-genome sequencing (WGS)-based gene-by-gene typing method that may overcome these limitations and has been applied successfully for several species in outbreak settings. In this study, genus-, genetic-complex-, and species-specific wgMLST schemes were developed for *Citrobacter* spp., the *Enterobacter cloacae* complex, *Escherichia coli*, *Klebsiella oxytoca*, and *Klebsiella pneumoniae* and used to type a national collection of 1,798 extended-spectrum-beta-lactamase-producing *Enterobacteriaceae* (ESBL-E) isolates obtained from patients in Dutch hospitals. Genus-, genetic-complex-, and species-specific thresholds for genetic distance that accurately distinguish between epidemiologically related and unrelated isolates were defined for *Citrobacter* spp., the *E. cloacae* complex, *E. coli*, and *K. pneumoniae*. wgMLST was shown to have higher discriminatory power and typeability than *in silico* MLST. In conclusion, the wgMLST schemes developed in this study facilitate high-resolution WGS-based typing of the most prevalent ESBL-producing species in clinical practice and may contribute to further elucidation of the complex epidemiology of antimicrobial-resistant *Enterobacteriaceae*. wgMLST opens up possibilities for the creation of a Web-accessible database for the global surveillance of ESBL-producing bacterial clones.

he continuing global spread of extended-spectrum-beta-lactamase-producing Enterobacteriaceae (ESBL-E) constitutes a major public health threat in both health care and community settings (1-4). Extended-spectrum beta-lactamases (ESBL) confer resistance to the majority of beta-lactam antibiotics, including third-generation cephalosporins, which limits the options for antimicrobial therapy and results in increased morbidity and mortality and health care costs (5, 6). Infection control guidelines recommend prevention of the spread of ESBL-E in health care settings (7, 8). Molecular typing has become an important tool in infection control, as it enables the detection of nosocomial transmission of bacterial pathogens and the identification of sources and routes of transmission in outbreak settings. Different molecular typing methods have been used, ranging from fingerprintbased methods, like pulsed-field gel electrophoresis (PFGE) and amplified fragment length polymorphism (AFLP), to sequencebased methods, like multilocus sequence typing (MLST) (9–11). Although widely adopted, PFGE is labor-intensive and difficult to standardize and has limited interlaboratory reproducibility (12-14). AFLP is less time-consuming than PFGE but may have lower discriminatory power for typing of Enterobacteriaceae (13, 15). MLST targets 7 or 8 housekeeping genes, depending on the Enterobacteriaceae species involved. It is a relatively fast, accurate, and reproducible strain-typing method, which has enabled the creation of an unambiguous nomenclature for bacterial clones. However, sequence conservation in housekeeping genes limits its discriminatory power and thus its use in outbreak investigations (16–18). Whole-genome sequencing (WGS) has emerged as an ultimate typing tool that fits any bacterial species, study type, and

laboratory (19–27). Knowledge of the sequence of the entire genome, rather than of only a few loci, has greatly increased the precision of molecular epidemiologic data and allows its use in infection control, where questions of similarity of isolates focus on recently diverged isolates. Although promising, the analysis, interpretation, and interlaboratory comparability of WGS-based typing results are still challenging. Whole-genome multilocus sequence typing (wgMLST) is a WGS-based typing method that extends MLST to the genome level (20). Today, such gene-bygene comparison of WGS data has been applied successfully for several species in outbreak settings, but data for *Enterobacteriaceae* and nonoutbreak settings are still limited (22–27). The aim of this study was to develop representative wgMLST schemes for the

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most prevalent ESBL-E species and to define thresholds for wgMLST-based genetic distance that reliably distinguish between epidemiologically related and unrelated ESBL-E isolates.

#### MATERIALS AND METHODS

**ESBL-E** isolates. A national collection of ESBL-E isolates was prospectively gathered in the SoM study, a multicenter cluster-randomized study comparing different isolation strategies for known ESBL-E carriers that was performed in 14 Dutch hospitals between 2011 and 2014 (28). ESBL-E isolates were obtained from routine clinical cultures and from perianal screening cultures that were taken during 690 repeated ward-based prevalence surveys performed in the participating hospitals. Descriptive characteristics, including species, date of culture, specimen type, patient, prevalence survey, ward, and hospital, were recorded for each isolate.

Microbiological methods. Detection of ESBL-E in clinical cultures was performed according to national and international guidelines (29, 30). Screening cultures consisted of perianal swabs that were preenriched in a selective tryptic soy broth and subsequently cultured on a selective ESBL screening agar (EbSA) (Cepheid Benelux, Apeldoorn, the Netherlands) as described previously (31). For all oxidase-negative Gram-negative bacteria that grew on the EbSA plate, species identification was performed with Vitek MS (bioMérieux, Marcy l'Etoile, France), using the Vitek MS Knowledge Base v1.1 (until January 2013) or v2.0 (from January 2013). The production of ESBL was phenotypically confirmed by the combination disk diffusion method with cefotaxime (30  $\mu$ g), ceftazidime (30  $\mu$ g), and cefepime (30  $\mu$ g), both alone and combined with clavulanic acid (10  $\mu$ g; Rosco, Taastrup, Denmark). Test results were considered positive if the inhibition zone around the disk was ≥5 mm larger for the combination with clavulanic acid (29, 30).

WGS. EbSA plates were inoculated with phenotypically confirmed ESBL-E isolates and incubated for 18 to 24 h at 35 to 37°C. For each isolate, several colonies (about 5 µl) of the culture were suspended in 300 µl microbead solution, which was subjected to DNA extraction with the Ultraclean Microbial DNA isolation kit (Mo Bio Laboratories, Carlsbad, CA, USA). The DNA concentration and purity were measured using a NanoDrop 2000c spectrophotometer (Thermo Fisher Scientific, Waltham, MA, USA) and the Qubit double-stranded DNA (dsDNA) HS and BR assay kits (Life Technologies, Carlsbad, CA, USA). One nanogram of bacterial DNA was used for library preparation. The DNA library was prepared using the Nextera XT library preparation kit with the Nextera XT v2 index kit (Illumina, San Diego, CA, USA). The library fragment length was aimed at fragments with a median size of 575 bases and was assessed with the Genomic DNA ScreenTape assay with the 2200 Tape-Station system (Agilent Technologies, Waldbronn, Germany). Subsequently, the library was sequenced on either a MiSeq sequencer, using the MiSeq reagent kit v2 generating 250-bp paired-end reads, or on a HiSeq 2500 sequencer, using the HiSeq Rapid SBS kit v2 generating 100-bp paired-end reads (Illumina, San Diego, CA, USA). Sequencing was aimed at a coverage of at least 60-fold. MiSeq data were processed with MiSeq control software v2.4.0.4 and MiSeq Reporter v2.4 and HiSeq data with bcl2fastq2 conversion software v1.8.4 (Illumina, San Diego, CA, USA). Quality trimming of reads was performed with CLC Genomics Workbench 7.0.4 (Qiagen, Hilden, Germany) using a minimum Phred (Q) score of 28. A random and blinded sample of eight isolates was sequenced

De novo assembly. De novo assembly was performed using CLC Genomics Workbench 7.0.4 (Qiagen, Hilden, Germany) with optimal word sizes based on the maximum N50 (the largest scaffold length, N, such that 50% of the assembled genome size is contained in scaffolds with a length of at least N) value. For each assembled genome, the coverage (mean depth), the number of scaffolds, the N50, the maximum scaffold length, and the percentage of the expected genome size were determined. Assembled genomes were excluded from further analysis if they did not fulfill the following quality criteria: (i) coverage, at least 20; (ii) number of scaffolds, at most 1,000; (iii) N50, at least 15,000 bases; (iv) maximum

scaffold length, at least 50,000 bases; and (v) assembled genome size, between 90% and 115% of the expected genome size.

WGS-based species identification, detection of ESBL-encoding genes, and MLST. Assembled genomes were uploaded to the online bioinformatics tools KmerFinder v2.0, ResFinder v2.1, and MLST v1.7 (Center for Genomic Epidemiology, Technical University of Denmark, Lingby, Denmark) (32–34). Species identification was based on the "winner takes it all" scoring method of KmerFinder. ESBL-encoding genes were reported when at least 60% of the length of the best matching gene in the ResFinder database was covered with a sequence identity of at least 90%. A nonstringent threshold for the percent alignment (60%) of ESBLencoding genes was used to account for the fact that short-read sequence data are not optimal for the assembly of plasmids, which may occasionally result in incomplete assembly of ESBL-encoding genes. Conventional MLST schemes in MLST v1.7 were based on the publicly available Pub-MLST.org database (http://www.pubmlst.org) and were available for Citrobacter freundii, Enterobacter cloacae, Escherichia coli, Klebsiella oxytoca, and Klebsiella pneumoniae (see Table S1 in the supplemental material). For E. coli, the Achtman MLST scheme was used (35). A sequence type (ST) was assigned when all MLST loci in the assembled genome perfectly matched all MLST alleles of a known ST in the MLST database, i.e., 100% sequence identity and 100% alignment for all alleles (32). When all MLST loci perfectly matched known MLST alleles but the combination of alleles was unknown in the database, isolates were classified as "unknown ST, perfect match." These isolates were considered typeable with MLST based on their allele profiles. Isolates with a sequence identity below 100% for one or more of the MLST loci were classified as "unknown ST, minor mismatch," and isolates with an alignment below 100% for one or more of the MLST loci were classified as "unknown ST, major mismatch." Isolates with minor or major mismatches were considered nontypeable with MLST.

Definition of WGS-based typing schemes. The MLST+ target definer function of Ridom SeqSphere+ (Ridom, Münster, Germany) was used to define target gene sets for all species with at least 10 isolates available. Depending on the number of available complete reference genomes, typing schemes were developed at the genus level (Citrobacter spp.), genetic-complex level (E. cloacae complex), or species level (E. coli, K. oxytoca, and K. pneumoniae). For each scheme, one annotated and publicly available complete genome (chromosome) was used as a reference genome (see Table S2 in the supplemental material). Open reading frames were extracted from the reference genomes and filtered by applying a minimum-length filter that discarded all genes with a length below 50 bases; a start codon filter that discarded all genes that contained no start codon at the beginning of the gene; a stop codon filter that discarded all genes that contained no stop codon, more than one stop codon, or a premature stop codon; an excluded sequences filter that discarded plasmid sequences; and a homologous gene filter that discarded all genes with fragments that occurred in multiple copies within the genome (with a sequence identity of at least 90% and more than 100-base overlap). Subsequently, the remaining set of reference genome genes was compared to a varying number of publicly available query genomes using the Basic Local Alignment Search Tool (BLAST) (36). The type and number of query genomes that were used differed by species and were dependent on the availability of annotated genomes (see Table S2 in the supplemental material). A gene overlap filter was used to identify reference genome genes that overlapped by more than 4 bases and to filter out the shorter gene. In addition, a stop codon percentage filter was applied to filter out genes that contained no stop codon, more than one stop codon, or a premature stop codon in more than 80% of the query genomes.

Reference genome genes that passed the gene overlap filter and the stop codon percentage filter were designated whole-genome targets. Of those, genes that were present in all query genomes with a sequence identity of at least 90% and an alignment of 100% were classified as core genome targets; the remaining genes were classified as accessory genome

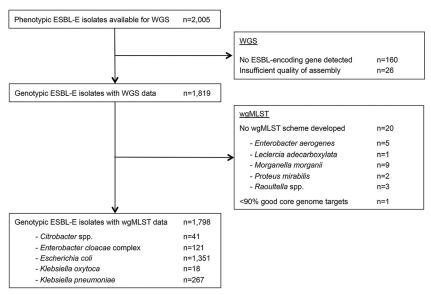


FIG 1 Availability of ESBL-E isolates for WGS and wgMLST.

targets. wgMLST schemes included all whole-genome targets; core genome MLST (cgMLST) schemes included only the core genome targets. An overview of the typing schemes, including the number of core, accessory, and whole-genome targets per typing scheme, is provided in Table S3 in the supplemental material.

Genetic distance. For each assembled genome, all whole-genome target genes that were present with a sequence identity of at least 90% and an alignment of 100% were extracted using Ridom SeqSphere+ software v 3.0 (Ridom, Münster, Germany). Genes that could not be identified in the assembled genome were classified as "missing targets." Genes with an alignment below 100% or a sequence identity below 90% were classified as "failed targets." The remaining genes were classified as "good targets." To evaluate the applicability and representativeness of the core genome target set, the percentage of good core genome targets was assessed for all assembled genomes. Assembled genomes for which the percentages of good core genome targets were below 90% were excluded from further analyses. All-to-all distance matrices, describing pairwise genetic distances, were constructed separately for core, accessory, and whole-genome target gene sets. The pairwise genetic distance was defined as the proportion of allele differences and was calculated by dividing the number of allele differences by the total number of good targets shared by both sequences, i.e., pairwise ignoring missing values.

Classification of pairwise comparisons. Pairwise comparisons of assembled genomes were classified according to the known epidemiological link between the isolates: (i) same patient; (ii) same survey but different patients; (iii) same ward but different surveys and patients; (iv) same hospital but different wards, surveys, and patients; and (v) no known epidemiological link, i.e., different hospitals, wards, surveys, and patients.

Definition of epidemiologically related and unrelated isolates. In the absence of a gold standard molecular typing method with perfect discriminatory power, epidemiological-link data were used to define related and unrelated isolates, where maximal contrast between related and unrelated isolates was sought. Epidemiologically related isolates were defined as being obtained from the same patient, belonging to the same ST, and cultured within a time window of 30 days. The last two restrictions were added to exclude potential bias from simultaneous carriage of multiple ESBL-E isolates from different clonal lineages and within-patient microevolution of isolates over time. Epidemiologically unrelated isolates were defined as being obtained from patients without any known epidemiological link.

Statistical analysis. The Kruskal-Wallis H test and sign test were used to test for differences in median genetic distance between species and between core and accessory genomes, respectively. All analyses were performed with IBM SPSS Statistics v22 (IBM Corp., Armonk, New York,

Ethical considerations. This study was reviewed by the Medical Research and Ethics Committee (MREC) of the Elisabeth/TweeSteden Hospital (Tilburg, the Netherlands) in December 2010. The study was judged to be beyond the scope of the Medical Research Involving Human Subjects Act (in Dutch, Wet Medisch-wetenschappelijk Onderzoek met Mensen [WMO]), and a waiver of written informed consent was obtained (METC/jv/2010.034). Patients who participated in the study provided verbal consent for use of demographic, clinical, and culture data.

**Accession number(s).** All generated raw reads were submitted to the European Nucleotide Archive (ENA) of the European Bioinformatics Institute (EBI) under the study accession number PRJEB15226.

#### **RESULTS**

A total of 2,005 phenotypic ESBL-E isolates were collected and available for WGS (Fig. 1). No ESBL-encoding gene could be detected in the assembled genomes of 160 (8.0%) isolates. For 26 (1.3%) isolates, the assembly was of insufficient quality. A summary of assembly characteristics for 1,819 genotypic ESBL-E isolates with good quality WGS data is provided in Table S4 in the supplemental material. WGS-based species identification was in accordance with that of Vitek-MS for 1,808 isolates. For 11 isolates that were identified as Citrobacter spp. by Vitek-MS, KmerFinder did not provide a high-confidence result for any species, probably due to the limited number of Citrobacter sp. reference genomes in the database. No wgMLST scheme was developed for Enterobacter aerogenes (n = 5), Leclercia adecarboxylata (n = 1), Morganella morganii (n = 9), Proteus mirabilis (n = 2), or Raoultella spp. (n =3). One *E. coli* isolate for which only 71.4% of core genome targets could be detected in the assembled genome was excluded from wgMLST analyses. For the remaining 1,798 ESBL-E isolates, the percentage of core genome targets present in the assembled genomes was at least 90.0% (see Table S5 in the supplemental material), which is indicative of the representativeness of the wgMLST schemes for the ESBL-E population studied. As expected, the percentage of accessory genome targets present in the

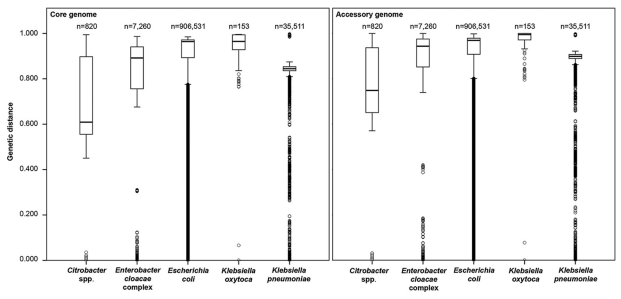


FIG 2 Distribution of genetic distances for 950,275 pairwise comparisons of extended-spectrum beta-lactamase-producing *Enterobacteriaceae*, shown separately for core genome and accessory genome. The boxes represent the interquartile (IQ) range. The whiskers extend to the highest and lowest values that are not greater than 1.5 times the IQ range. The lines across the boxes indicate the medians. The circles represent outliers, i.e., comparisons with values more than 1.5 times the IQ range. n defines the number of pairwise comparisons for each species. Kruskal-Wallis chi-square test results for difference between species were as follows: chi-square n core genome = 22,685, n < 0.0005; chi-square n = 15,980, n < 0.0005. Sign test for the difference between core and accessory genomes, n < 0.0005.

assembled genomes varied considerably within and between species.

The sequence identity of the ESBL-encoding genes that were identified was 100% for 1,751 (97.6%) ESBL-E isolates. For the remaining 47 (2.4%) isolates, the percent sequence identity with known ESBL-encoding genes ranged from 99.1% to 99.9%. The percent alignment of the ESBL-encoding genes was 100% for 1,789 isolates (99.5%) and ranged from 66.6% to 97.9% for the other 9 (0.5%) isolates.

The all-to-all genetic distance matrices comprised 950,275 unique pairwise comparisons. Figure 2 shows the species-specific distributions of genetic distance for core and accessory genome targets. Pairwise genetic distances ranged from 0.000 to (approximately) 1.000 for both core and accessory genomes and for all species. However, the distribution of genetic distances differed significantly between species (P < 0.0005) and between core and accessory genomes (P < 0.0005). For the core genome, median genetic distances ranged from 0.609 for Citrobacter spp. to 0.965 for K. oxytoca and, for the accessory genome, from 0.748 for Citrobacter spp. to 0.995 for K. oxytoca. For all species, the median genetic distance was higher for the accessory genome than for the core genome. Summary statistics of the distribution of numbers of alleles compared; numbers of different alleles; and genetic distances for core genome, accessory genome, and whole genome are presented by species in Table S6 in the supplemental material. To assess the added value of the accessory genome in terms of genetic distance, the difference between whole-genome and core genome genetic distances was calculated for each pairwise comparison (Fig. 3). For the majority of pairwise comparisons (82.0%) [779,316/950,275]), the whole-genome genetic distance was larger than the core genome distance, although the absolute difference differed between species. For the remaining comparisons, the inclusion of accessory genome targets reduced the genetic distance. Five E. coli and three *K. pneumoniae* isolates were sequenced in duplicate. The results were found to be reproducible, with genetic distances ranging from 0.000 to 0.001 for both core and whole-genome targets.

Table 1 provides the distributions of genetic distances for isolates with the same (conventional) ST and isolates with different

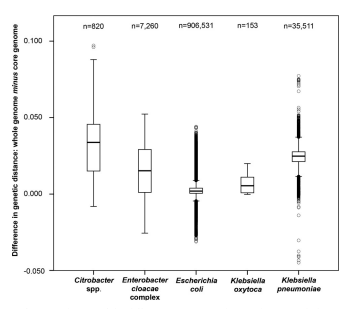


FIG 3 Distribution of the differences in genetic distance between whole genome and core genome for 950,275 pairwise comparisons of extended-spectrum beta-lactamase-producing *Enterobacteriaceae*. The boxes represent the IQ range. The whiskers extend to the highest and lowest values that are not greater than 1.5 times the IQ range. The lines across the boxes indicate the medians. The circles represent outliers, i.e., pairwise comparisons with values more than 1.5 times the IQ range. n defines the number of pairwise comparisons for each species. Kruskal-Wallis chi-square test results for difference between species were as follows: chi-square = 69,760, P < 0.0005.

TABLE 1 Genetic distances for pairwise comparisons of ESBL-E isolates within and between conventional MLST STs<sup>a</sup>

MLST and species	Within ST $(n = 68,482)$			Between ST ( $n = 834,045$ )		
	Pairwise comparison $(n)$	Genetic distance			Genetic distance	
		Median	Range	Pairwise comparison $(n)$	Median	Range
cgMLST						
Citrobacter spp.	14	0.000	[0.000-0.036]	31	0.897	[0.537-0.917]
E. cloacae complex	430	0.006	[0.000-0.104]	5,135	0.881	[0.088-0.987]
E. coli	67,108	0.022	[0.000-0.507]	796,847	0.966	[0.019-0.985]
K. oxytoca	1	0.066	[0.066-0.066]	65	0.992	[0.789-0.996]
K. pneumoniae	929	0.020	[0.000-0.542]	31,967	0.844	[0.090-0.998]
wgMLST						
Citrobacter spp.	14	0.000	[0.000-0.027]	31	0.913	[0.558-0.933]
E. cloacae complex	430	0.007	[0.000-0.129]	5,135	0.902	[0.102-0.988]
E. coli	67,108	0.027	[0.000-0.520]	796,847	0.968	[0.021-0.986]
K. oxytoca	1	0.070	[0.070-0.070]	65	0.994	[0.795-0.996]
K. pneumoniae	929	0.022	[0.000-0.566]	31,967	0.870	[0.104-0.997]

<sup>&</sup>lt;sup>a</sup> Excluding 47,748 (5.0%) pairwise comparisons for which no ST could be assigned to either of the isolates.

STs. As expected, wgMLST and cgMLST enabled further differentiation of isolates that belong to the same ST. The broadest within-ST ranges in genetic distance were observed for *E. coli* and *K. pneumoniae*, with ranges from 0.000 up to 0.520 and from 0.000 up to 0.566 (wgMLST), respectively. Although median genetic distances clearly differed for within-ST and between-ST comparisons, the ranges of genetic distances overlapped for *E. cloacae*, *E. coli*, and *K. pneumoniae*, indicating that some isolates with different STs can be genetically more closely related than isolates with the same ST. For 47,748 (5.0%) pairwise comparisons, no ST could be assigned to either of the isolates because of minor or major mismatches for at least one of the seven alleles involved.

The distributions of genetic distances for epidemiologically related and unrelated isolates were clearly different. Figure 4 shows the distributions of wgMLST-based genetic distances for epidemiologically related and unrelated ESBL-producing E. coli. Comparable distributions were observed for epidemiologically related and unrelated isolates of the other ESBL-E species (see Fig. S1 in the supplemental material) and for cgMLST-based genetic distances (data not shown). Based on these distributions for epidemiologically related and unrelated isolates, thresholds for genetic distance were set at the minimal value that classified 100% of epidemiologically related isolate pairs as clonally related (Table 2). The absence of a pair of epidemiologically related K. oxytoca isolates in the ESBL-E strain collection precluded the definition of thresholds for K. oxytoca. Thresholds differed between species, with two to three times higher values for E. coli than for the other three species. Thresholds for cgMLST- and wgMLST-based genetic distances were similar for Citrobacter spp. and the E. cloacae complex but differed slightly for E. coli (a lower threshold for wgMLST) and K. pneumoniae (a lower threshold for cgMLST). For all genetic distance thresholds, the corresponding percentage of presumed epidemiologically unrelated isolate pairs that were classified as clonally related was low (0.1% to 0.5%), which indicates high discriminatory power, i.e., the ability to assign different types to different isolates (37).

Thresholds for genetic distance were subsequently used to type all 1,798 ESBL-E isolates, and the results were compared with those of *in silico* MLST (Table 3). Epidemiological concordance was observed for all three typing methods, i.e., the percentage of

isolates identified as clonally related was highest for within-patient comparisons and decreased with waning degrees of epidemiological linkage. The typeability, i.e., the percentage of isolates to which a genotype could be assigned, was significantly higher for cgMLST and wgMLST (99.98%) than for *in silico* MLST (95.0%; P < 0.0005) (37). In addition, cgMLST and wgMLST had higher discriminatory power than *in silico* MLST, i.e., classification of presumed epidemiologically unrelated isolates as clonally related was significantly less frequent for cgMLST (0.5%) and wgMLST (0.3%) than for *in silico* MLST (7.1%, P < 0.0005). Extending cgMLST to wgMLST resulted in a 0.2% increase in the overall percentage of correct classification of presumed epidemiologically

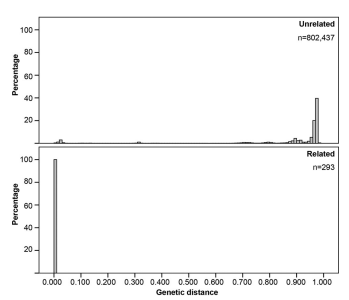


FIG 4 Distributions of wgMLST-based genetic distances for pairwise comparisons of epidemiologically related and unrelated extended-spectrum beta-lactamase-producing *Escherichia coli* isolates. Epidemiologically related isolates were defined as being obtained from the same patient, belonging to the same ST, and cultured within a time window of 30 days. Epidemiologically unrelated isolates were defined as being obtained from patients without a known epidemiological link.

**TABLE 2** Thresholds for genetic distance with corresponding classification of epidemiologically unrelated ESBL-E isolates<sup>a</sup>

		Threshold-based classification of epidemiologically unrelated isolates <sup>c</sup>			
	Threshold for genetic	Pairwise comparison	Clonally related		
MLST and species	distance <sup>b</sup>	(n)	n	%	
cgMLST					
Citrobacter spp.	0.0030	693	1	0.1	
E. cloacae complex	0.0035	6,256	31	0.5	
E. coli	0.0105	802,437	3,999	0.5	
K. pneumoniae	0.0035	30,808	75	0.2	
wgMLST					
Citrobacter spp.	0.0030	693	1	0.1	
E. cloacae complex	0.0035	6,256	17	0.3	
E. coli	0.0095	802,437	2,485	0.3	
K. pneumoniae	0.0045	30,808	96	0.3	

<sup>&</sup>lt;sup>a</sup> The absence of a pair of epidemiologically related *K. oxytoca* isolates precluded the definition of thresholds for *K. oxytoca*.

unrelated isolates as clonally unrelated (P < 0.0005). In addition, reclassification analysis of individual comparisons showed that extension to wgMLST improved classification for 1,551 of 840,338 (0.2%) epidemiologically unrelated isolate pairs, i.e., presumed epidemiologically unrelated isolate pairs were reclassified from clonally related (cgMLST) to clonally unrelated (wgMLST) (Table 4). On the other hand, 44 of 840,338 (0.005%) presumed epidemiologically unrelated isolate pairs were classified as clonally related by wgMLST, while they were classified as clonally unrelated by cgMLST.

#### **DISCUSSION**

In this study, representative genus-, genetic-complex-, and species-specific wgMLST schemes were developed for the five most prevalent *Enterobacteriaceae* species in clinical practice and used to type a large national collection of ESBL-E isolates obtained from patients in Dutch hospitals. Genus-, genetic-complex-, and species-specific thresholds for genetic distance that accurately distinguished between epidemiologically related and unrelated isolates were defined for *Citrobacter* spp., the *E. cloacae* complex, *E. coli*, and *K. pneumoniae*. wgMLST was shown to have higher discriminatory power and typeability than *in silico* MLST.

After the introduction of WGS as a typing tool with high potential, gene-by-gene comparison of WGS data has been applied successfully for several species in outbreak settings (22–27). To our knowledge, this large multicenter study is the first to apply wgMLST for typing of *Enterobacteriaceae* in nonoutbreak settings and to include a comprehensive evaluation of its performance in a large collection of isolates without a clear epidemiological link. The availability of detailed epidemiological data enabled the definition of thresholds for wgMLST-based genetic distance that can

TABLE 3 Typing results for 950,275 pairwise comparisons of ESBL-E isolates with different degrees of epidemiological linkage by MLST, cgMLST, and wgMLST

	Pairwise comparison					
Epidemiological link between isolates	Clonally related		Clonally unrelated		Nontypeable <sup>a</sup>	
and typing method	n	%	n	%	n	%
Same patient $(n = 726)$						
MLST	550	75.8	136	18.7	40	5.5
cgMLST	564	77.7	161	22.2	1	0.1
wgMLST	565	77.8	160	22.0	1	0.1
Same survey $(n = 1,180)$						
MLST	168	14.2	930	78.8	82	6.9
cgMLST	129	10.9	1,051	89.1	0	0.0
wgMLST	131	11.1	1,049	88.9	0	0.0
Same ward ( $n = 14,291$ )						
MLST	1,165	8.2	12,465	87.2	661	4.6
cgMLST	329	2.3	13,960	97.7	2	0.0
wgMLST	301	2.1	13,988	97.9	2	0.0
Same hospital ( $n = 93,740$ )						
MLST	6,908	7.4	82,414	87.9	4,418	4.7
cgMLST	921	1.0	92,813	99.0	6	0.0
wgMLST	778	0.8	92,956	99.2	6	0.0
Other hospital ( $n = 840,338$ )						
MLST	59,691	7.1	738,100	87.8	42,547	5.1
cgMLST	4,106	0.5	836,088	99.5	144	0.0
wgMLST	2,599	0.3	837,595	99.7	144	0.0

 $<sup>^</sup>a$  MLST, unknown ST, minor or major mismatch (n=47,748); cgMLST/wgMLST, no cutoff available for K. oxytoca (n=153).

be used to distinguish between epidemiologically related and unrelated isolates in hospital settings and provided estimates of the probability of classifying presumed epidemiologically unrelated isolates as clonally related in settings with low levels of transmission.

Available studies on WGS-based typing have shown that reliable phylogenies with good resolution can be inferred from the core genome (16). It has been suggested, though, that resolution at the isolate level may be further improved by the additional targeting of more variable genes from the accessory genome (16). In the present study, wgMLST resulted in a larger genetic distance between isolates than cgMLST for the majority of isolate comparisons and in less frequent classification of presumed epidemiologically unrelated isolates as clonally related and thus had slightly higher discriminatory power. Based on these findings, the use of

TABLE 4 Reclassification of clonal (un)relatedness of epidemiologically unrelated ESBL-E isolates after extension of cgMLST to wgMLST

	Pairwise comparison by wgMLST (n)			
Pairwise comparison by cgMLST	Clonally related	Clonally unrelated	Nontypeable	
Clonally related	2,555	1,551	0	
Clonally unrelated	44	836,044	0	
Nontypeable	0	0	$144^{a}$	

<sup>&</sup>lt;sup>a</sup> All nontypeable isolates were *K. oxytoca* isolates.

<sup>&</sup>lt;sup>b</sup> Thresholds were set at the minimal value for genetic distance that classified 100% of epidemiologically related ESBL-E isolates as clonally related.

<sup>&</sup>lt;sup>c</sup> Epidemiologically related isolates were those obtained from the same patient, belonging to the same ST, and cultured within a time window of 30 days. Epidemiologically unrelated isolates were those obtained from patients without a known epidemiological link. Clonally related isolates were those at a genetic distance below or equal to the threshold value.

wgMLST is preferred over cgMLST for the *Enterobacteriaceae* species studied here. However, extension with the accessory genome was shown to result in a decrease in genetic distance for a minority of isolates and in rare cases even in classification of presumed epidemiologically unrelated isolates as clonally related.

Thresholds for genetic distance that distinguished epidemiologically related from unrelated isolates differed between species and were highest for *E. coli*. These high thresholds for *E. coli* may reflect higher mutation and recombination rates or may be related to within-patient microevolution of coexisting isolates with common ancestry. Estimates of the mutation and recombination rates of *Enterobacteriaceae* are limited and range from 0 to 10 mutations per genome per year (38, 39). These rates reflect fixed substitution rates; short-term mutation (polymorphism) rates may be higher and are known to increase under antibiotic pressure or other environmental stress (40).

Due to the lack of *K. oxytoca* isolate pairs that fulfilled the definition of epidemiologically related isolates, thresholds for genetic distance for the species are still to be defined. The distribution of genetic distances for epidemiologically unrelated *K. oxytoca* isolates was comparable to that of the other *Enterobacteriaceae* species and might be used to evaluate proposed thresholds in future studies.

wgMLST was shown to have high discriminatory power. With genetic distance thresholds set at the minimal value that classified 100% of epidemiologically related isolates as clonally related, only 0.3% of presumed epidemiologically unrelated isolates were classified as clonally related. Thus, wgMLST may enable high-resolution detection of nosocomial transmission of antimicrobial-resistant *Enterobacteriaceae*. Classification of presumed epidemiologically unrelated isolates as clonally related may be due to a residual lack of resolution of wgMLST but may also reflect the presence of an unidentified epidemiological link in the hospital or community setting. Other possible explanations may be related to clone-specific characteristics, such as a low mutation rate, converging evolution, or recent introduction into the human population.

Interestingly, about 20% of within-patient comparisons were found to pertain to clonally unrelated isolates, indicating that patients are cocolonized with multiple ESBL-producing isolates. This finding may reflect multiple acquisitions of ESBL-E but may also be the result of within-patient horizontal transfer of ESBL-encoding mobile genetic elements.

An important strength of wgMLST is the use of alleles and not nucleotide polymorphisms as units of comparison. Irrespective of the number of nucleotide polymorphisms involved, each allelic change is numbered as a single event, i.e., an allelic change is related to at least one point mutation, but can also contain several nucleotide changes. Thus, allele-based methods largely reduce the conflicting signals of recombination in determining genetic relatedness of bacteria (41-45). Another major advantage of this gene-by-gene allelic profiling method is the immediate and automated assignment of novel allele variants, which enables a standardized and universal nomenclature and is a prerequisite for successful interlaboratory exchange of data (46). Most studies on wgMLST have used the absolute number of allele differences to compare isolates, which is highly dependent on the number of targets that are compared (22, 23, 26, 27). In the present study, differences in the numbers of targets compared were taken into account by defining genetic distance as the proportion of allele differences. This enabled the unbiased comparison of genetic distances for different pairwise comparisons and the definition of thresholds for genetic distance.

A limitation of allele-based typing is that the analysis is reduced to coding regions only, as second-generation sequencing platforms currently in use produce only relatively short reads that do not facilitate high-quality assembly of highly repetitive intergenic regions. This might change when third-generation sequencing instruments that produce much longer reads from a single molecule become widely available. Nevertheless, the current wgMLST approach will be sustainable, as it will maintain backward compatibility with the expansion of typing schemes, as we see today with the *in silico* extraction of conventional MLST sequence types from WGS data.

In conclusion, the wgMLST schemes developed in this study facilitate high-resolution WGS-based typing of the most prevalent ESBL-producing species in clinical practice and may contribute to further elucidation of the complex epidemiology of antimicrobial-resistant *Enterobacteriaceae*. wgMLST opens up possibilities for the creation of a Web-accessible database for the global surveil-lance of ESBL-producing bacterial clones.

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