Cholera is a life-threatening diarrheal disease caused by the bacterium *Vibrio cholerae*, which produces cholera toxin. The seventh cholera pandemic, caused by *V. cholerae* O1 biotype El Tor, began in Indonesia in 1961 and reached Africa in 1970 (1). Fifty years later, >100,000 cases of cholera are reported annually in sub-Saharan Africa (2).

The Central African Republic (CAR) is a land-locked and resource-limited country in central Africa; it was ranked 188/189 on the United Nations Human Development Index in 2018 (http://hdr.undp.org). CAR is relatively large but has a low population density; 2019 data estimate ≈4.75 million inhabitants, or 7.75 persons/km² (Macrotrends LLC, https://www.macrotrends.net). CAR largely has been spared by the cholera epidemic; only 4 outbreaks had been reported by 2020 (Table 1; Figure 1). The first 2 cholera outbreaks occurred during the same month in 1997 (4). In the first, 443 cases and 88 deaths were reported in southern CAR, along the Oubangui River, close to the border with the Democratic Republic of the Congo (DRC). In the second, 113 cases and 19 deaths were reported in northern CAR, close to the borders with Cameroon and Chad, after which cholera cases continued to be detected in southern CAR, along the Oubangui River. According to reports from the International Federation of Red Cross and Red Crescent Societies, 172 cholera cases and 16 deaths were reported in the region in 2011 (https://reliefweb.int/sites/reliefweb.int/files/resources/MDRCF009finalrep.pdf) and 265 cases and 20 deaths were reported in 2016 (https://reliefweb.int/report/central-african-republic/central-africa-republic-cholera-epidemic-outbreak-dref-operation).

The Institut Pasteur de Bangui in CAR performed microbial analyses to confirm the causal agent of these outbreaks and identified 30 *V. cholerae* O1 serotype Inaba isolates collected during 1997–2016 (Appendix 1 Table 1, https://wwwnc.cdc.gov/EID/article/27/1/20-0375-App1.xlsx).

We used whole-genome sequencing to fully characterize all 30 *V. cholerae* O1 isolates in terms of virulence and antimicrobial resistance determinants. We also placed these genomes within a broader phylogenetic context to elucidate their origins and evolutionary history.

**The Study**

The 30 *V. cholerae* O1 isolates were received at the Institut Pasteur, Paris, France. We performed antimicrobial susceptibility testing, whole-genome sequencing, comparative genomics, and phylogenetic analyses by using methods previously described (3,5–11) (Appendix 2, https://wwwnc.cdc.gov/EID/...
We then contextualized these 30 *V. cholerae* O1 isolates within a global collection of 1,185 seventh pandemic *V. cholerae* El Tor (7PET) genomic sequences and constructed a maximum-likelihood phylogeny of 1,215 genomes by using 9,964 single-nucleotide variants (SNVs) evenly distributed over the nonrepetitive, nonrecombinant core genome (Figure 2, panel A). Our phylogenomic analysis (Appendix 1 Tables 2–4) showed that all CAR isolates belonged to the 7PET lineage (12,13).

### Table 1. Characteristics of the *Vibrio cholerae* O1 isolates associated with outbreaks of cholera, Central African Republic*

<table>
<thead>
<tr>
<th></th>
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<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>No. deaths/no. cases†</td>
<td>88/443</td>
<td>19/113</td>
<td>16/172</td>
<td>20/265</td>
</tr>
<tr>
<td>No. isolates</td>
<td>9</td>
<td>6</td>
<td>7</td>
<td>8</td>
</tr>
<tr>
<td>7PET sublineage</td>
<td>T7</td>
<td>T9</td>
<td>T10</td>
<td>T10</td>
</tr>
<tr>
<td>Sequence type</td>
<td>69</td>
<td>69</td>
<td>515</td>
<td>515</td>
</tr>
<tr>
<td>ctxB</td>
<td>B3</td>
<td>B1</td>
<td>B1</td>
<td>B1</td>
</tr>
<tr>
<td>wbeT‡</td>
<td>A03</td>
<td>C19</td>
<td>B01</td>
<td>B01</td>
</tr>
<tr>
<td>AMR phenotypes (no. isolates)§</td>
<td>R1 (8), R2 (1)</td>
<td>R3 (6)</td>
<td>R3 (7)</td>
<td>R3 (8)</td>
</tr>
</tbody>
</table>

**AMR determinants**

<table>
<thead>
<tr>
<th>Plasmid</th>
<th>IncA/C¶</th>
<th>VC_0715</th>
<th>WT</th>
<th>R169C</th>
<th>R169C</th>
<th>R169C</th>
</tr>
</thead>
<tbody>
<tr>
<td>VC_A0637</td>
<td>WT</td>
<td>Q5Stop</td>
<td>Q5Stop</td>
<td>Q5Stop</td>
<td>Q5Stop</td>
<td></td>
</tr>
<tr>
<td>SXT/R391 element</td>
<td>NT</td>
<td>ICEVchInd5</td>
<td>ICEVchInd5</td>
<td>ICEVchInd5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>gyrA</td>
<td>WT</td>
<td>WT</td>
<td>S83I</td>
<td>S83I</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*7PET, seventh pandemic *Vibrio cholerae* O1 El Tor; AMR, antimicrobial resistance; NT, not detected; WT, wild type.
‡Nomenclature according to Weill et al. (3) (Table 2).
§R1, resistance to polymyxin B and colistin; R2, resistance to polymyxin B, colistin, ampicillin, streptomyacin, sulfonamides, vibriostatic agent, trimethoprim/sulfamethoxazole, chloramphenicol; R3, resistance to polymyxin B, colistin, streptomyacin, sulfonamides, vibriostatic agent, trimethoprim/sulfamethoxazole, furazolidone, and resistance or intermediate resistance to chloramphenicol.
¶Isolate (CNRVC970079) with the R2 AMR type.

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Previous genomic studies described 12 introductions of the 7PET lineage from Southern Asia into Africa during 1970–2016 (3,6). The introduced sublineages were called T1 and T3–T13. The 2 cholera outbreaks in CAR in 1997 were caused by sublineage T7, which had been introduced into West Africa during the early 1980s, and T9, which was introduced in the late 1980s, according to Weill et al. (3). T9 isolates were identified in neighboring countries such as Chad and Cameroon (particularly northern Cameroon) before they were detected in northwest CAR, but T7 isolates were identified in Gabon and the western part of DRC, along the Congo River in Kinshasa, before being identified in CAR along the Oubangui River, a tributary of the Congo River (Figure 2, panel B; Appendix 2 Figures 1, 2).

The 2011 and 2016 outbreaks were caused by closely related bacterial populations from the same sublineage, T10, introduced into East Africa during the 1990s and later detected in the African Great Lakes Region (AGLR) (3) (Figure 2, panel B; Appendix 2 Figures 1, 2). The prevalent T10 sublineage has several clades, and the 2011 and 2016 CAR isolates are characterized by multilocus sequence type (ST) 515, a single-locus variant of ST69; ST69 is the predominant ST among 7PET isolates. In addition, these isolates have an alteration to the \textit{wbeT} gene, a 4-nucleotide deletion called B01 (Tables 1, 2), that underly the Inaba serotype. Phylogenetic data showed that the 7PET strains causing the 2011 and 2016 cholera outbreaks in CAR spread from AGLR to the western part of the DRC and CAR (Figure 2, panel B; Appendix 2 Figure 3). These genomic data are consistent with results from a recent epidemiologic study showing the spatiotemporal distribution of cholera cases during the 2011–2012 and 2015–2017 outbreaks in DRC, suggesting a spread from cholera hotspots in AGLR to major cities in the upstream section of the Congo River, followed by a downstream spread toward the densely populated CAR capital, Kinshasa, and then to the mouth of the Congo River, which opens into the Gulf of Guinea (14).

### Table 2. Alterations of the \textit{wbeT} (formerly \textit{rfbT}) gene in \textit{Vibrio cholerae} El Tor isolates from the Central African Republic*

<table>
<thead>
<tr>
<th>Alteration no.</th>
<th>Alteration type</th>
<th>Genetic alteration</th>
<th>Protein consequence†</th>
<th>7PET sublineage</th>
</tr>
</thead>
<tbody>
<tr>
<td>A03</td>
<td>Premature stop codon</td>
<td>G133T</td>
<td>G45-to-STOP</td>
<td>T7</td>
</tr>
<tr>
<td>B01</td>
<td>Premature stop codon</td>
<td>Deletion TGTAC (nt 24–28)</td>
<td>Frameshift after N7; then STOP</td>
<td>T10</td>
</tr>
<tr>
<td>C19</td>
<td>Amino acid substitution</td>
<td>G674A</td>
<td>C225Y</td>
<td>T9</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>T9</td>
</tr>
</tbody>
</table>

*7PET, seventh pandemic \textit{Vibrio cholerae} O1 El Tor.
†The codon number and single-letter amino acid abbreviation are shown (STOP corresponds to a stop codon).
Except during the outbreak in northwest CAR in 1997, all cholera cases were reported along or close to the Oubangui River, suggesting that 7PET strains probably moved from area to area along the river and with the displacement of human populations. The risk factors in these remote areas are unknown, but the prevailing conditions, such as poor hygiene and sanitary conditions, overcrowding, lack of latrines, and drinking water from the Oubangui River, likely would increase the risk for transmission via the fecal–oral route, as evidenced by the high attack rates observed at several sites when the 2011 outbreak began (15). Nevertheless, since the declaration of the first case in 1997, the small number (<1,000) of cholera cases in CAR contrasts with the much larger numbers in central Africa (2). The low population density of CAR, its poor transport infrastructure, and poor trading links are probably key factors limiting disease spread (15). Phylogenetic analyses showed no other isolates from Africa were derived from CAR isolates in the aftermath of the 4 outbreaks, which also suggests that the transmission of cholera is impeded in this country. Of note, all 4 outbreaks were caused by serotype Inaba 7PET strains. This serotype has a nonmethylated form of lipopolysaccharide caused by an alteration to the \textit{wbeT} gene (3) (Table 2). The implication of this serotype in all 4 outbreaks suggests that these 7PET sublineages circulated regionally for some time, long enough to acquire this alteration to the \textit{wbeT} gene, before reaching CAR (Appendix 2 Figures 1–3).

All CAR isolates in this study displayed resistance to polymyxin B, consistent with the susceptibility pattern reported for the El Tor biotype until recently (6). All but 1 of the isolates collected along the Oubangui River in 1997 were susceptible to all other antimicrobial drugs tested; the outlying isolate contained the extended-spectrum β-lactamase \textit{bla\textsubscript{SHV-2a}} gene on an IncA/C2 plasmid (Table 1; Appendix 1 Table 3). No susceptible isolates have been collected in CAR since. All the other isolates display mutations of the \textit{VC\textsubscript{0715}} and \textit{VC\textsubscript{A0637}} genes, conferring nitrofuran resistance, and carry an SXT/R391 genomic element called ICE\textit{V chol}Nd5, encoding resistance to streptomycin (\textit{strAB}), sulfonamides (\textit{su}l2), trimethoprim, the O/129 vibriostatic agent (\textit{dfrA}1), and trimethoprim-sulfamethoxazole (\textit{su}l2 and \textit{dfrA}1). The 2011 and 2016 CAR isolates also recently acquired \textit{gyrA} mutations (Table 1), resulting in resistance to nalidixic acid (Appendix 1 Table 3).

**Conclusions**

Strains from 3 7PET sublineages caused 4 cholera outbreaks identified in CAR during 1997–2016. The southern and southeastern parts of CAR are higher risk areas for cholera outbreaks, particularly when cases are reported in the western part of DRC. These findings highlight the need for an effective surveillance system, and for coordinated communication actions on cholera that target healthcare professionals and the populations living along the Oubangui River, to prevent and control cholera outbreaks in CAR.

This work was supported by the Institut Pasteur, Santé Publique France and the French Government’s Investissement d’Avenir program, Laboratoire d’Excellence Integrative Biology of Emerging Infectious Diseases (grant no. ANR-10-LABX-62-IBEID).

**About the Author**

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**References**


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- US CDC Real-Time Reverse Transcription PCR Panel for Detection of Severe Acute Respiratory Syndrome Coronavirus 2
- Coronavirus Disease Outbreak in Call Center, South Korea
- Investigation and Serologic Follow-Up of Contacts of an Early Confirmed Case-Patient with COVID-19, Washington, USA
- Characteristics and Outcomes of Coronavirus Disease Patients under Nonsurge Conditions, Northern California, USA, March–April 2020
- Tuberculosis in Internationally Displaced Children Resettling in Harris County, Texas, USA, 2010–2015
- Epidemiology of Legionnaires’ Disease, Hong Kong, China, 2005–2015
- Rise in Babesiosis Cases, Pennsylvania, USA, 2005–2018
- Sporadic Creutzfeldt-Jakob Disease among Physicians, Germany, 1993–2018
- Analysis of MarketScan Data for Immunosuppressive Conditions and Hospitalizations for Acute Respiratory Illness, United States
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- SARS-CoV-2 Phylogenetic Analysis, Lazio Region, Italy, February–March 2020

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Seventh Pandemic *Vibrio cholerae* O1 Sublineages, Central African Republic

Appendix 2

**Antimicrobial Susceptibility Testing**

Antimicrobial susceptibility was determined by the disk diffusion method, on Mueller-Hinton agar (Bio-Rad, https://www.bio-rad.com), in accordance with the guidelines of the Antibiogram Committee of the French Society for Microbiology (1). The following antimicrobial drugs (Bio-Rad) were tested: ampicillin, cefalotin, cefotaxime, streptomycin, chloramphenicol, azithromycin, sulfonamides, trimethoprim-sulfamethoxazole, vibriostatic agent O/129, tetracycline, nalidixic acid, ciprofloxacin, nitrofurantoin, polymyxin B, and colistin. *Escherichia coli* CIP 76.24 (ATCC no. 25922) was used as a control. The MICs of nalidixic acid and ciprofloxacin were determined by Etests (bioMérieux, https://www.biomerieux.com).

**Whole-Genome Sequencing**

We analyzed 30 *Vibrio cholerae* O1 biotype El Tor isolates from the Central African Republic (CAR) by whole-genome sequencing. Fourteen of these isolates were collected between 1997 and 2011 and were sequenced in a previous study (2). The 24 *V. cholerae* O1 biotype El Tor isolates sequenced here consisted of the remaining 16 isolates from CAR and included all isolates from 2016 and 8 isolates from neighboring countries, Chad and the Democratic Republic of the Congo (Appendix 1 Table 1).

Total DNA was extracted with the Maxwell 16-cell DNA purification kit (Promega, https://www.promega.com) in accordance with the manufacturer's recommendations. Whole-genome sequencing was carried out at the Biomics and PIBnet sequencing platforms of the Institut Pasteur, the genotyping and sequencing core facility of the Institut du Cerveau et de la Moëlle Epinière (Paris, France), or at Macrogen Korea (Seoul, South Korea), on Illumina

**Additional Genomic Data**

Raw sequence files from 1,164 seventh generation pandemic *V. cholerae* El Tor (7PET) genomes described by Weill et al. (2,3) and Irenge et al. (4) were downloaded from the European Nucleotide Archive (ENA; https://www.ebi.ac.uk/ena) and included in this study (Appendix 1 Table 1). Twelve assembled genomes also described by Weill et al. (2,3) were downloaded from GenBank (https://www.ncbi.nlm.nih.gov/genbank; Appendix 1 Table 1) and included in this study. We generated 100 bp overlapping simulated reads from the 12 assembled genomes with fasta_to_fastq.pl (https://github.com/ekg/fasta-to-fastq/blob/master/fasta_to_fastq.pl).

**Genomic Sequence Analyses**

The paired-end reads and simulated paired-end reads were mapped onto the reference genome of *V. cholerae* O1 El Tor N16961 (GenBank accession nos. LT907989 and LT907990) with Snippy v4.1.0/BWA-MEM v0.7.13 (https://github.com/tseemann/snippy). Single-nucleotide variants (SNVs) were called with Snippy v4.1.0/Freebayes v1.1.0 (https://github.com/tseemann/snippy) under the following constraints: mapping quality of 60, a minimum base quality of 13, a minimum read coverage of 4, and a 75% read concordance at a locus for a variant to be reported. An alignment of core genome SNVs was produced in Snippy v4.1.0 for phylogeny inference. Short reads were assembled with SPAdes version 3.1.0 (5).

In silico multilocus sequence typing (MLST) for *V. cholerae* was performed with MLST version 2.0. (https://cge.cbs.dtu.dk/services/MLST), on assembled sequences for the entire dataset (6). The various genetic markers were analyzed with BLAST version 2.2.26. against reference sequences of the O1 *rfb* gene, *ctxB*, *wbeT*, and the whole locus of VSP-II, as previously described (2,3,7).

The presence and type of acquired antimicrobial resistance genes (ARGs) or ARG-containing structures were determined with ResFinder version 3.1.0. (https://cge.cbs.dtu.dk/services/ResFinder), BLAST analysis against GI-15, Tn7, and SXT/R391 integrative and conjugative elements, and PlasmidFinder version 2.0.1. (https://cge.cbs.dtu.dk/services/PlasmidFinder). The presence of mutations in the genes
encoding resistance to quinolones (gyrA, parC), resistance to nitrofurans (VC_0715 and VC_A0637), or restoring susceptibility to polymyxin B (vprA) was investigated by manual analysis of the sequences assembled de novo with BLAST (https://blast.ncbi.nlm.nih.gov), as previously described (2,3,7).

**Phylogenetic Analysis**

Repetitive (insertion sequences and the TLC-RS1-CTX region) and recombinogenic (VSP-II) regions in the alignment were masked (2). Putative recombinogenic regions were detected and masked with Gubbins version 2.3.4 (8). A maximum likelihood (ML) phylogenetic tree was built from an alignment of 9,964 chromosomal SNVs, with RAxML version 8.0.20., under the GTR model with 200 bootstraps (9). The final tree was rooted on the A6 genome and visualized with iTOL version 5 (https://itol.embl.de) or FigTree version 1.4.2. (http://tree.bio.ed.ac.uk/software/figtree).

**Data Availability**

Short-read sequence data were submitted to the ENA (http://www.ebi.ac.uk/ena), under study accession no. PRJEB36666 (see Appendix 1 Table 4 for accession nos.).

**References**


Appendix 2 Figure 1. Maximum likelihood phylogeny for the T7 seventh pandemic Vibrio cholerae El Tor sublineage. Bootstrap values $>90\%$ are shown at the branch nodes. The genomes from the Central African Republic are shown in red. Acquisition of the genetic variation of the $wbeT$ (also named $rfbT$) gene implicated in the Inaba serotype, is indicated by an arrow.
Appendix 2 Figure 2. Maximum likelihood phylogeny for the T9 seventh pandemic *Vibrio cholerae* El Tor sublineage. Bootstrap values >90% are shown at the branch nodes. The genomes from the Central African Republic are shown in red. Acquisition of the genetic variation of the *wbeT* (also named *rfbT*) gene implicated in the Inaba serotype, is indicated by an arrow. Red asterisks indicate the derived isolates are Ogawa revertants.
Appendix 2 Figure 3. Maximum likelihood phylogeny for the T10 seventh pandemic *Vibrio cholerae* El Tor sublineage containing sequence type 515 (ST515) *V. cholerae* O1 isolates from the Central African Republic. Bootstrap values greater than or equal to 90% are shown at the branch nodes. The genomes from the Central African Republic are shown in red. Acquisition of the genetic variation of the *wbeT* (also named *rfbT*) gene implicated in the Inaba serotype, is indicated by an arrow.