Outbreak of Listeriosis in South Africa Associated with Processed Meat


ABSTRACT

BACKGROUND
An outbreak of listeriosis was identified in South Africa in 2017. The source was unknown.

METHODS
We conducted epidemiologic, trace-back, and environmental investigations and used whole-genome sequencing to type Listeria monocytogenes isolates. A case was defined as laboratory-confirmed L. monocytogenes infection during the period from June 11, 2017, to April 7, 2018.

RESULTS
A total of 937 cases were identified, of which 465 (50%) were associated with pregnancy; 406 of the pregnancy-associated cases (87%) occurred in neonates. Of the 937 cases, 229 (24%) occurred in patients 15 to 49 years of age (excluding those who were pregnant). Among the patients in whom human immunodeficiency virus (HIV) status was known, 38% of those with pregnancy-associated cases (77 of 204) and 46% of the remaining patients (97 of 211) were infected with HIV. Among 728 patients with a known outcome, 193 (27%) died. Clinical isolates from 609 patients were sequenced, and 567 (93%) were identified as sequence type 6 (ST6). In a case–control analysis, patients with ST6 infections were more likely to have eaten polony (a ready-to-eat processed meat) than those with non-ST6 infections (odds ratio, 8.55; 95% confidence interval, 1.66 to 43.35). Polony and environmental samples also yielded ST6 isolates, which, together with the isolates from the patients, belonged to the same core-genome multilocus sequence typing cluster with no more than 4 allelic differences; these findings showed that polony produced at a single facility was the outbreak source. A recall of ready-to-eat processed meat products from this facility was associated with a rapid decline in the incidence of L. monocytogenes ST6 infections.

CONCLUSIONS
This investigation showed that in a middle-income country with a high prevalence of HIV infection, L. monocytogenes caused disproportionate illness among pregnant girls and women and HIV-infected persons. Whole-genome sequencing facilitated the detection of the outbreak and guided the trace-back investigations that led to the identification of the source.
LISTERIOSIS, a severe foodborne disease that has substantial mortality (20 to 30%), primarily affects persons with impaired cell-mediated immunity associated with pregnancy, extremes of age, underlying malignant conditions, human immunodeficiency virus (HIV) infection, chronic disease, or immunosuppressive therapy.1-5 Outbreaks are increasingly recognized,6,7 predominantly in upper-income countries where infection is more readily diagnosed,8 where existing surveillance programs facilitate early recognition,9 and where strain typing by whole-genome sequencing, which allows for identification of outbreak-linked cases and definitive attribution of the source, is accessible.10-14

An increase in the number of cases of listeriosis at two public hospitals in Gauteng Province, South Africa, during July and August 2017 prompted an investigation. Case numbers rapidly increased nationwide, and whole-genome multilocus sequence typing9 of Listeria monocytogenes isolates from patients identified a single sequence type (sequence type 6 [ST6]) in 93% of the cases. We used whole-genome sequencing and intensive epidemiologic and trace-back investigations to pursue the source of the outbreak. This report describes the key findings from the investigation.

METHODS

CASE DEFINITION

We defined an outbreak-associated case as laboratory-confirmed infection with L. monocytogenes, as determined by means of bacterial culture or polymerase-chain-reaction (PCR) assay of any clinical sample, during the outbreak period (June 11, 2017 [epidemiologic week 24 of that year], to April 7, 2018 [epidemiologic week 14]). This period was defined as the interval during which the case numbers at the national level exceeded the threshold of five cases per week. The threshold of five cases per week was determined with the use of baseline laboratory data from January 1, 2013, to December 31, 2016. All L. monocytogenes infections were initially included in the case definition, because it was not possible to definitively exclude non-ST6 cases from the outbreak event; however, for the case–control analysis, the case definition was later refined to include ST6 cases only. Pregnancy-associated cases included illness with an onset during pregnancy or within the first 2 weeks of the postpartum period and illness in the neonate. The mothers of infected neonates were not counted among those who had cases if they did not have symptomatic laboratory-confirmed listeriosis; infection in a maternal–neonatal pair was defined as laboratory-confirmed infection in both the mother and neonate and was counted as a single case. Neonatal cases were classified as early onset (diagnosed between birth and day 6) or late onset (diagnosed between days 7 and 28). A map showing the incidence of infections according to district was generated (Fig. 1).

EPIDEMIOLOGIC CASE INVESTIGATION

Clinical and demographic details and underlying medical conditions were ascertained through patient interviews or abstracted from medical records or laboratory reports with the use of a standardized case-investigation form. This investigation was reviewed in accordance with local and Centers for Disease Control and Prevention procedures for protection of human research participants and was considered nonresearch disease-control activity in a public health emergency. From November 1, 2017, all patients with newly reported cases were contacted to assess food exposures during the 4 weeks preceding the onset of illness with the use of a semistructured questionnaire. In cases in which the patient was a child, had died, or was too ill to respond, the next of kin were interviewed as proxies. In neonatal cases, history of food consumption by the mother during pregnancy was obtained.

Among the subgroup of patients with a detailed food history and available whole-genome sequencing results, a case–control analysis was performed to estimate the odds ratios for the association between specific food exposures and outbreak-associated illness. In this analysis, a case patient was defined as a person with L. monocytogenes ST6 infection and a control patient as a person with non-ST6 listeriosis during the outbreak period.

ENVIRONMENTAL AND TRACE-BACK INVESTIGATIONS

Health authorities initiated the collection of food samples from the homes of patients in mid-November 2017. When L. monocytogenes was isolated from a food sample, a trace-back investigation was conducted.
Characterization of the Outbreak Strain

*L. monocytogenes* isolates were sent to a national reference laboratory, where genomic bacterial DNA was isolated and whole-genome sequencing analysis performed as described previously.\(^{17}\) Genome assemblies were analyzed with the use of the multilocus sequence typing analysis pipeline at the Center for Genomic Epidemiology (www.genomicepidemiology.org). Data from multilocus sequence typing were used to determine clonal complexes and sequence types.\(^{15}\) Raw sequencing data were analyzed with the use of the Bacterial Isolate Genome Sequence Database for *L. monocytogenes* (BIGSdb-Lm, http://bigsdb.pasteur.fr/listeria/listeria.html) to determine sublineages and core-genome multilocus sequence types.\(^{18}\) The data exported from the BIGSdb-Lm were analyzed with BioNumerics Software, version 7.6.2 (bioMérieux) in order to perform a core-genome multilocus sequence typing–based phylogenetic analysis with the use of a single-linkage clustering algorithm.

The virulence of the ST6 strain was assessed in 7-to-10-week-old E16P KI C57BL/6 female mice, as previously described;\(^{19}\) approval was obtained from the Institut Pasteur ethics committee. We assessed the virulence of the ST6/CT4148 YA00061615 CLIP2018/00699 human isolate (L1-SL6-ST6-CT4148, in which L denotes phylogenetic lineage, SL sublineage, ST sequence type, and CT core-genome multilocus sequence type) as the South African strain, and compared it with that of the EGDe ST9 reference strain (L2-SL9-ST35-CT637; National Center for Biotechnology Information (NCBI) GenBank accession number, NC_003210)\(^{20}\) and the CLIP2009/01092 ST6 iso-
late (L1-SL6-CT451; NCBI accession number, PRJEB10792).21 Overnight culture of *L. monocytogenes* was diluted in brain–heart infusion medium to reach mid-log growth phase. The mice were inoculated intragastrically through a feeding needle with $2 \times 10^8$ colony-forming units. The infected animals were killed 4 days after inoculation, and the organs were dissected and homogenized. Serial dilutions of ground-tissue suspensions in phosphate-buffered saline were inoculated on brain–heart infusion agar plates. After 24 hours of incubation at 37°C, the colony-forming units were counted.

**Results**

**Outbreak Cases**

A total of 937 cases were reported during the outbreak period, with case numbers peaking at 41 per week in mid-November 2017 (epidemiologic week 46) (Fig. 2). ST6 was identified in 567 of 609 sequenced clinical isolates (93%). Although ST6 cases predominated during the outbreak period, smaller peaks of non-ST6 cases were noted. The number of cases decreased dramatically after recall of the implicated products on March 4, 2018. By mid-April 2018 (6 weeks after recall), fewer than 5 cases were reported weekly. Although cases were reported in all provinces, 543 of the 937 cases (58%) occurred in Gauteng Province, where the incidence reached 5 cases per 100,000 population in several districts (Fig. 1).

**Clinical Information**

A total of 465 of the 937 cases (50%) were associated with pregnancy: 406 cases (43%) occurred in neonates and 59 (6%) in pregnant girls and women. Nine maternal–neonatal pairs were identified. Early-onset disease occurred in 95% of the neonatal cases. Of the 937 cases, 229 (24%)
occurred in patients 15 to 49 years of age (excluding those who were pregnant) (Table 1). With the exclusion of the 59 girls and women known to be pregnant, female patients were overrepresented in the age group of 15 to 49 years (140 of 229 [61%]). A total of 83 cases (9%) occurred in persons 65 years of age or older. Overall, all but 2 patients were hospitalized, and no health care–associated infections were documented.

HIV status was known in 415 of the 937 cases (44%). In 204 pregnancy-associated cases with known HIV status, 77 patients (38%) had positive HIV status, which included HIV exposure in 60 of 158 neonates (38%) and HIV infection in 17 of 46 pregnant girls and women (37%). Among the remaining 211 patients, 97 (46%) were infected with HIV. Among the 114 patients (excluding neonates) who were infected with HIV, 82 (72%) had available data on the CD4 T-lymphocyte count; the median count was 194 cells per cubic millimeter (interquartile range, 91 to 387). Maternal CD4 T-lymphocyte counts were known for 12 HIV-exposed neonates (20%); the median count was 479 cells per cubic millimeter (interquartile range, 322 to 575). After adjusting for age and sex, we found that the odds of ST6 infection were 48% lower among HIV-infected patients than among those without HIV infection, although odds lower than 78% or higher than 25% are also compatible with our data (odds ratio for ST6 infection, 0.52; 95% CI, 0.22 to 1.25). HIV-infected patients older than 1 month of age were 2.55 times as likely to have meningitis (odds ratio, 2.55; 95% CI, 1.38 to 4.72).

The outcome was known for 728 patients (78%), among whom 193 deaths were reported (case-fatality ratio, 27%). HIV infection was associated with a 53% increased odds of death among patients older than 1 month, after adjustment for age and sex (odds ratio, 1.53; 95% CI, 0.75 to 3.15). Of the 4 maternal deaths reported, the underlying risk factors were known in 1 patient (diabetes mellitus and HIV infection). Fetal loss occurred in 27 of the 59 pregnant girls and women (46%).

CASE–CONTROL ANALYSIS
A total of 109 patients were interviewed. Consumption of polony (a ready-to-eat processed meat containing chicken, pork, beef, or any combination of these, similar to bologna) was reported by 93 patients (85%), and the brands produced by Facility A were the most commonly reported. Sequence data were available for 76 of the 109 patients: 65 had ST6 infections (case patients) and 11 had non-ST6 infections (control patients). The food items most strongly associated with ST6 infection included polony (odds ratio, 8.55; 95% CI, 1.66 to 43.35) and frozen chicken (odds ratio, 4.90; 95% CI, 1.04 to 25.55) (Table 2). Of the 57 case patients who reported eating polony, 50 (88%) reported eating brands manufactured at Facility A, although several patients reported eating several brands or did not specify a particular brand.

TRACE-BACK AND ENVIRONMENTAL INVESTIGATIONS
On January 13, 2018, febrile gastroenteritis developed in 10 children from a nursery in Gauteng Province. Several stool samples were collected, and one yielded L. monocytogenes ST6. Sandwiches prepared and eaten at the nursery were the only common food exposure, and polony was the common ingredient. Polony was recovered from the nursery refrigerator, and L. monocytogenes ST6 was identified in the polony produced at Facility A.

On February 2, 2018, an environmental investigation was conducted at Facility A, located in Limpopo Province. Production of the polony entailed grinding and mixing raw ingredients, stuffing the emulsion into clipped nylon casings, cooking the polony loaves in hot water, and cooling the loaves in a brine chiller. Several areas were in poor repair, and many opportunities for cross-contamination of food products were identified, including condensation, unrestricted movement of workers, and prolonged reuse of brine for chilling.

L. monocytogenes was isolated from 47 of the 317 environmental samples (15%) collected at Facility A. A total of 34 of the 47 typed isolates (72%) were identified as ST6. These isolates originated from samples collected at several facility sections (precooking and postcooking), including from food-contact surfaces, non–food-contact surfaces, and chilling brine. L. monocytogenes ST6 was detected in 2 of 13 samples of unopened polony loaves collected at the facility.
Table 1. Characteristics of Patients with Laboratory-Confirmed Listeriosis during the Outbreak Period (June 11, 2017, to April 7, 2018).*

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>No. of Cases (N = 937)</th>
<th>Pregnancy-Associated Cases</th>
<th>Cases According to Age Group</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Neonates (N = 406)</td>
<td>Pregnant Patients (N = 59)</td>
</tr>
<tr>
<td>Female sex — no./total no. of patients with known sex (%)</td>
<td>516/918 (56)</td>
<td>208/392 (53)</td>
<td>59/59 (100)</td>
</tr>
<tr>
<td>Risk factor — no./total no. of patients with known risk factor (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HIV§</td>
<td>174/415 (42)</td>
<td>60/158 (38)</td>
<td>17/46 (37)</td>
</tr>
<tr>
<td>Malignant condition</td>
<td>20/188 (11)</td>
<td>NA</td>
<td>0/42</td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>20/187 (11)</td>
<td>NA</td>
<td>1/42 (2)</td>
</tr>
<tr>
<td>Renal failure</td>
<td>10/187 (5)</td>
<td>NA</td>
<td>0/42</td>
</tr>
<tr>
<td>Immuno compromised condition other than HIV¶</td>
<td>10/186 (5)</td>
<td>NA</td>
<td>0/42</td>
</tr>
<tr>
<td>Alcohol abuse</td>
<td>10/187 (5)</td>
<td>NA</td>
<td>0/42</td>
</tr>
<tr>
<td>Liver failure</td>
<td>7/187 (4)</td>
<td>NA</td>
<td>1/42 (2)</td>
</tr>
<tr>
<td>Died — no./total no. of patients with known outcome (%)</td>
<td>193/728 (27)</td>
<td>86/308 (28)</td>
<td>4/51 (8)</td>
</tr>
<tr>
<td>Culture site — no./total no. of patients (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blood</td>
<td>677/937 (72)</td>
<td>376/406 (93)</td>
<td>35/59 (59)</td>
</tr>
<tr>
<td>Cerebrospinal fluid</td>
<td>197/917 (21)</td>
<td>19/406 (5)</td>
<td>1/59 (2)</td>
</tr>
<tr>
<td>Other‖</td>
<td>63/917 (7)</td>
<td>11/406 (3)</td>
<td>23/59 (39)</td>
</tr>
</tbody>
</table>

* The completeness of responses on the case-investigation forms varied among the patients; therefore, the patient denominators vary among the characteristics assessed. HIV denotes human immunodeficiency virus, and NA not applicable.
† This age group excludes patients who were known to be pregnant.
‡ A dash indicates that data were not available.
§ HIV-exposure status (i.e., maternal HIV status) was reported for neonates.
¶ Immunocompromising conditions other than HIV included the receipt of glucocorticoid therapy or chemotherapeutic agents.
‖ Other sample types included stool, urine, synovial fluid, tissue-biopsy specimen, pus, umbilical cord, placenta, amniotic fluid, cervical swab, and gastric aspirate.
and subsequently from polony loaves sold in retail stores.

**Characterization of the Outbreak Strain**

Whole-genome sequencing was performed in 710 human *L. monocytogenes* isolates (8 isolates collected in 2015; 37 in 2016; 455 in 2017; and 210 in 2018) and in 1061 food and environmental *L. monocytogenes* isolates collected between September 1, 2017, and March 31, 2018. On the basis of multilocus sequence typing, 567 of 609 isolates (93%) from the outbreak cases were identified as ST6, and the remainder represented 14 other sequence types. A total of 34 environmental isolates from Facility A and 19 isolates from food produced at Facility A were identified as ST6.

Whole-genome sequencing data for 386 ST6 isolates were analyzed with the use of core-genome multilocus sequence typing. Four human ST6 isolates that were recovered in 2017 and 2018 differed by at least 14 alleles, but the remaining 382 differed by no more than 7 alleles (out of 1748 loci included in the scheme) (Fig. 3A). This maximum 4-allelic difference is within the 7-allelic difference threshold that defines potentially epidemiologically linked isolates, as described by Moura et al. These 382 isolates, including 336 human isolates and 19 food and 27 environmental isolates from Facility A, were assigned to the same core-genome multilocus sequence type (CT4148; complete genotype, L1-SL6-ST6-CT4148). The oldest South African CT4148 isolates date from September 2015 and are related to a cluster of three cases of listeriosis in

### Table 2. Case–Control Analysis of the Association between Specific Food Exposures and *Listeria monocytogenes* ST6 Infection.

<table>
<thead>
<tr>
<th>Food Consumed</th>
<th>Case Patients (N = 65)</th>
<th>Control Patients (N = 11)</th>
<th>Odds Ratio (95% CI)‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polony</td>
<td>57 (88)</td>
<td>5 (45)</td>
<td>8.55 (1.66–43.35)</td>
</tr>
<tr>
<td>Kota‡</td>
<td>9 (14)</td>
<td>0</td>
<td>ND</td>
</tr>
<tr>
<td>Frozen chicken</td>
<td>42 (65)</td>
<td>4 (36)</td>
<td>4.90 (1.04–25.55)</td>
</tr>
<tr>
<td>Any delicatessen-style ready-to-eat meat</td>
<td>57 (88)</td>
<td>8 (73)</td>
<td>2.67 (0.37–14.38)</td>
</tr>
<tr>
<td>Apples</td>
<td>51 (78)</td>
<td>6 (55)</td>
<td>3.86 (0.77–18.16)</td>
</tr>
</tbody>
</table>

* A total of 109 patients were interviewed, among whom sequence data were available for 76 patients (65 [86%] had *L. monocytogenes* ST6 infections [case patients] and 11 [14%] had non-ST6 listeriosis [control patients]). ND denotes not determined (unable to calculate odds ratio because of zero value in cell).
† The odds ratio is for the association between the specific food item and *L. monocytogenes* ST6 infection.
‡ Kota is a sandwich-type fast food containing polony.
A  Population Structure of the South African L. monocytogenes ST6 Outbreak–Associated Isolates

Origin of isolates (no. of isolates)
- Human (340)
- Production environment (27)
- Food (19)

N=174

L1-SL6-ST6-CT443

14

N=29

N=1

L1-SL6-ST6-CT4148

B  L. monocytogenes SL6 Phylogeny

Region of Isolates (no. of isolates)
- Asia (1)
- Europe (754)
- North America (259)
- South Africa (1)
- South America (9)
- Unknown (93)

1998 — U.S. hot dog–associated outbreak

2017–18 — South African polony–associated outbreak

2013 — U.S. French-style cheese–associated outbreak

2018 — European frozen corn–associated outbreak

2002 — U.S. turkey deli meat–associated outbreak

Percentage of Similarity

97 98 99 100

N=174

N=1

N=29

N=1

N=1
Western Cape Province; this finding suggests a potential epidemiologic link to the 2017–2018 outbreak (Fig. 3A).

Whole-genome sequencing data were compared with information from curated databases through international networks and with isolates representative of previous major ST6 outbreaks to detect possible matches, and none were found (Fig. 3B, and Fig. S1 in the Supplementary Appendix, available with the full text of this article at NEJM.org); genome sequences for 10 L. monocytogenes ST6 isolates associated with this South African outbreak have been deposited at the NCBI GenBank repository under the accession numbers QEXB00000000 to QEXK00000000 (Bio-Project number, PRJNA451422; and BioSample numbers, SAMN08970424 to SAMN08970415). Outbreak ST6 isolates had a hypervirulent phenotype, as described previously for L. monocytogenes ST6, but were not more virulent than a strain representative of ST6 (Fig. S2).

**CONTROL MEASURES**

On March 4, 2018, the Minister of Health announced the outbreak source. Facility A products were traced and recalled from distributors and retailers, and the public was advised to return products for reimbursement. Facility A was closed immediately. The World Health Organization assisted in recalling the products that had been exported to 15 African countries (Angola, Botswana, Democratic Republic of Congo, Ghana, Lesotho, Madagascar, Malawi, Mauritius, Mozambique, Namibia, Nigeria, eSwatini, Uganda, Zambia, and Zimbabwe). A single case of listeriosis was reported in Namibia during March 2018, but the patient’s isolate was confirmed as non-ST6; no other countries reported cases during the period from January 1, 2017, to September 3, 2018, when the outbreak was declared over.

**DISCUSSION**

Contaminated polony that was produced at a single facility was the cause of a large national outbreak of listeriosis predominantly associated with an L. monocytogenes ST6 strain in South Africa. Evidence supporting the cause includes a strong association between eating polony and ST6 infection, the detection of ST6 isolates in polony recovered from the refrigerator in the nursery, the detection of ST6 isolates in unopened polony loaves collected at the production facility, the detection of ST6 isolates in environmental samples collected from the production facility, a decline in ST6 cases after a recall of the products and closure of the facility, and the fact that outbreak-associated ST6 isolates from patients as well as food and environmental isolates from the facility belonged to a single core-genome multilocus sequence type.

Ready-to-eat processed meats are a well-known vehicle for listeriosis outbreaks. Polony is a low-cost, readily available food popular across all socioeconomic groups in South Africa and is used in kota, a fast food favored in urban areas. Polony has a shelf life of 5 months and is produced in large quantities by several manufacturers for local consumption and export.

Unique features of this outbreak include its recognition in a middle-income country with a high prevalence of HIV infection and a high fertility rate. HIV infection is a well-recognized risk factor for listeriosis. In 2017, the prevalence of HIV infection in South Africa was 12.6%, and an estimated 7.9 million people were living with HIV infection. The prevalence of HIV infection in the age group of 15 to 49 years was 26.3% among female persons and 14.8% among male persons. Of the nearly 1.2 million girls and women who gave birth in 2017, approximately 265,000 (22%) were infected with HIV.

In this outbreak, HIV infection was the most common predisposing condition among patients younger than 65 years of age. We also found that a large percentage of pregnant girls and women were infected with HIV (37% [17 of 46]), and a similar percentage of neonates were exposed to HIV (38% [60 of 158]). The percentage of female patients was highest in the age group of 15 to 49 years, which suggests that possible unrecognized HIV infection or pregnancy were predisposing conditions. Culture-confirmed or PCR assay–confirmed L. monocytogenes infections were reported in more cerebrospinal fluid samples from HIV-positive patients than from HIV-negative patients. It is possible that HIV infection is associated with an increased mortality, but given the missing data and study design, the analyses were not powered to detect a difference between the HIV-infected group and the HIV-uninfected group.

The size and velocity of the outbreak were
noteworthy and most likely resulted from the wide distribution of large volumes of contaminated products in a large population vulnerable to invasive listeriosis. Likely explanations for the predominance of cases in Gauteng Province include consumer behavior, food preferences, and higher socioeconomic status in this densely populated province, but increased reporting or differential health-seeking and physician-testing behavior may have been contributing factors. We found no evidence that this ST6 outbreak–associated strain was more virulent than a strain representative of ST6.21

No outbreak-associated cases were detected in the 15 low-to-middle-income countries that imported polony from Facility A. However, cases were probably missed because of nonspecific clinical presentation, physician-testing behavior, limited diagnostic capacity at the laboratory, and lack of surveillance for listeriosis. The burden of listeriosis is most likely higher than is currently recognized in low-income and middle-income countries, particularly those with large populations of people living with HIV.

Before this outbreak, listeriosis was not required to be reported and not under surveillance in South Africa. A national surveillance system has since been implemented, and all isolates from patients are analyzed by means of whole-genome sequencing. This outbreak catalyzed a revision of local food-safety regulations; certification through the Hazard Analysis and Critical Control Point system is now a legal requirement for ready-to-eat meat producers, and microbiologic criteria for L. monocytogenes in ready-to-eat foods are under review.

This outbreak investigation had several limitations. Not all clinical isolates were available for whole-genome sequence typing. Microbiologic investigations are not routinely conducted in pregnant women with mild, nonspecific febrile illness or in those who have had miscarriages or stillbirths; therefore, the number of patients with pregnancy-related listeriosis was most likely underestimated. With respect to the period under study, limited case-investigation forms were available, and data were of varying completeness. Insufficient clinical data prohibited a description of clinical syndromes. Sources of the data on HIV status included laboratory reports and reports from the patients themselves, which probably resulted in underreported positive status. On the basis of the findings from core-genome multilocus sequence typing, it is likely that the ST6 cluster in 2015 was associated with the 2017–2018 outbreak; however, incomplete histories of food consumption and trace-back data precluded a conclusive epidemiologic link.

The findings showcase the power of complementary epidemiologic data and whole-genome sequence typing for detecting and investigating foodborne disease outbreaks and show that whole-genome sequencing technology can be ably implemented and used in developing countries. As the global shift to whole-genome sequence typing for foodborne-pathogen surveillance accelerates, developing countries should build the capacity to leverage this technology in a rapidly evolving landscape of food-safety concerns. Targeted health communication for the prevention of listeriosis among pregnant girls and women and HIV-infected persons in developing countries may help mitigate the risk of disease in these vulnerable groups.

The findings and conclusions in this report are those of the authors and do not necessarily represent the official position of the Centers for Disease Control and Prevention.

Disclosure forms provided by the authors are available with the full text of this article at NEJM.org.

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APPENDIX

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