

## Genome-Based Characterization of *Listeria monocytogenes*, Costa Rica

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Genomic data on the foodborne pathogen *Listeria monocytogenes* from Central America are scarce. We analyzed 92 isolates collected during 2009–2019 from different regions in Costa Rica, compared those to publicly available genomes, and identified unrecognized outbreaks. Our findings suggest mandatory reporting of listeriosis in Costa Rica would improve pathogen surveillance.

*L*isteria monocytogenes is a gram-positive pathogen responsible for listeriosis, a severe foodborne infection that causes high hospitalization and mortality rates in at-risk populations, including older adults, immunocompromised persons, pregnant women, and newborns (1). *L. monocytogenes* diversity can be classified into lineages, genoseroroups, clonal complexes (CCs), and sequence types (STs), defined by multilocus sequence typing (MLST) (2).

Core-genome MLST (cgMLST) further identifies sublineages (SLs) and cgMLST types (CTs) (2). Major CCs and SLs are distributed globally and can be heterogeneous in terms of virulence; isolates from serogroup IVb (lineage I) often cause the most severe infections (2–4).

Pathogen surveillance using whole-genome sequencing (WGS) provides unprecedented resolution for identifying case clusters and contamination sources and for predicting strain virulence and antimicrobial resistance, which can aid in risk assessment (2,5). Previous studies confirmed *L. monocytogenes* in various foods in Costa Rica; reported contamination levels were 5%–20% in processed meat products and fresh cheeses (6,7). Because listeriosis is not a notifiable disease in Costa Rica, its prevalence is unknown, and diversity of *L. monocytogenes* circulating in the country is unclear.

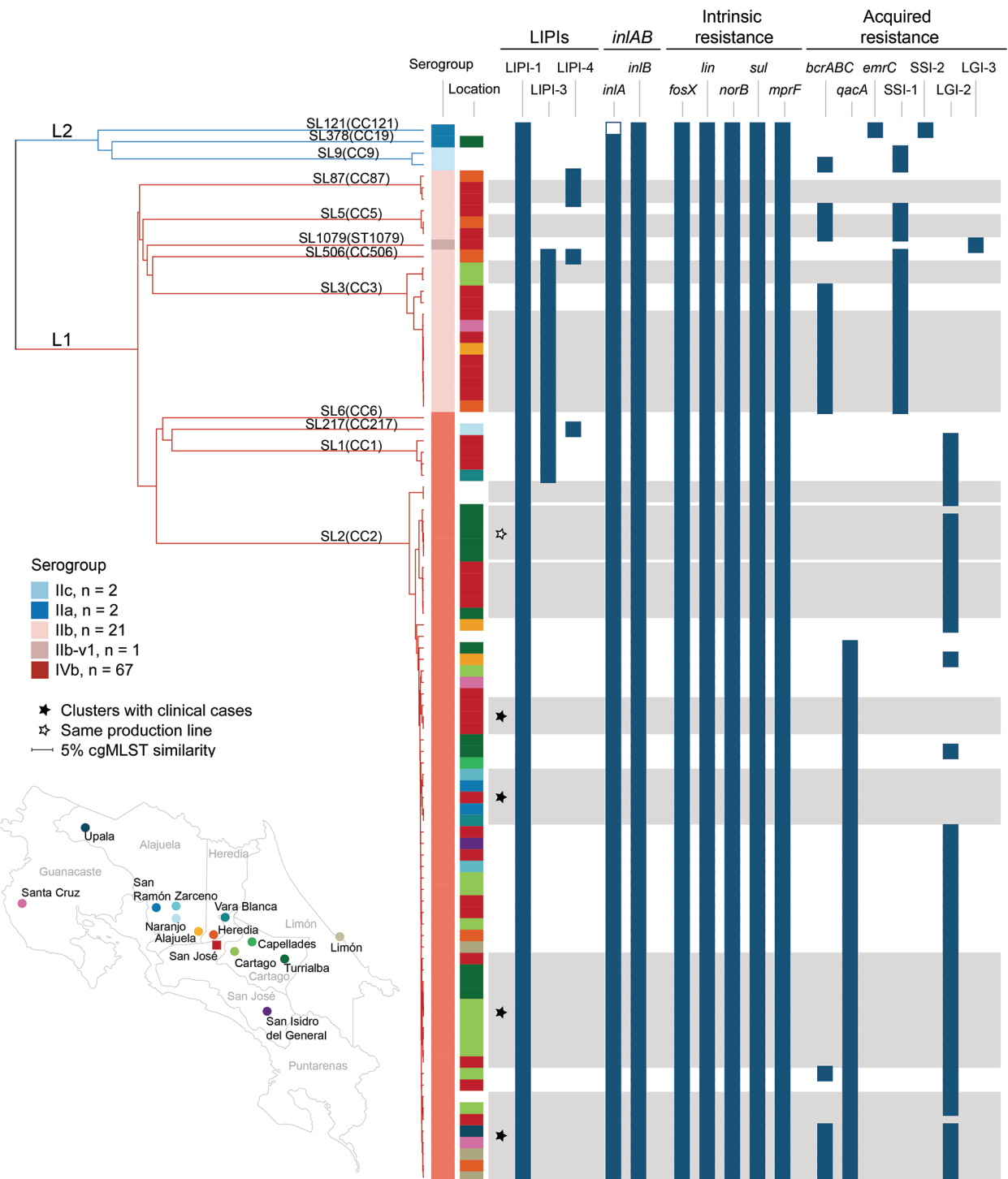
To clarify the diversity of and potential public health risk from circulating strains, we used WGS to characterize 92 isolates recovered during 2009–2019 from 16 clinical, 67 food, and 9 production environment samples in Costa Rica (Appendix, <https://wwwnc.cdc.gov/EID/article/29/12/23-0774-App1.pdf>). When location data were available, isolates were from urban areas, including the capital city San José, and from rural areas where fresh cheese production is prevalent, including Alajuela, Naranjo, San Ramón, Vara Blanca, Upala, and Turrialba. Turrialba region accounts for 70% of fresh cheese produced in Costa Rica (Figure; Appendix).

We found that isolates from lineage I (95%, n = 88) and lineage II (5%, n = 4) were unevenly distributed into 12 different SLs and CCs (Figure; Appendix Figure 1). Those isolates included a new lineage I sublineage, designated SL1079 (new MLST singleton ST1079), which was identified in an isolate from shrimp (cgMLST type L1-SL1079-ST1079-CT1669). That isolate had an atypical genoseroroup IIb profile, designated IIb-v1, that differed from the classic IIb profile by the presence of *lmo0737*. WGS confirmed the presence of *lmo0737* and flanking genes *lmo0733–39*, typically found in lineage II isolates from serogroups IIa and IIc but only occasionally found in lineage I serogroup IVb-v1 (8). Of note, 80% of isolates investigated from both clinical and food-associated sources were from sublineages SL2/CC2 (66%, n = 61) and SL3/CC3 (14%, n = 13). SL2/CC2 (serogroup IVb) and SL3/CC3 (serogroup IIb) isolates are found worldwide and are associated with invasive infections (2–4). However, they are rarely the most prevalent genotypes (2,3). Available data from other countries in Central America confirmed

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**Figure.** Single linkage dendrogram of 92 isolates generated for genome-based characterization of *Listeria monocytogenes*, Costa Rica. Dendrogram is based on core-genome multilocus sequence typing; (cgMLST) allelic profiles (1,748-locus scheme). Branches are colored according to lineages: L1, red; L2, blue. Branches are labeled according to lineages, sublineages, and clonal complexes. Information on isolates' serogroup, and resistance profiles are provided in the columns. Colors in location column correspond to dots on map. Gray bars indicate clusters of isolates with <7 allelic differences out of 1,748 cgMLST loci. Presence of selected virulence and resistance genetic traits in each isolate is represented by dark blue boxes and empty boxes denote genes with premature stop codons. More details are provided in Appendix Figure 1 (<https://wwwnc.cdc.gov/EID/article/29/12/23-0774-App1.pdf>). CC, clonal complex; L, lineage; LIPI, listeria pathogenicity island; SL, sublineage.

**Table.** Sequence types identified in a genome-based characterization of *Listeria monocytogenes*, Costa Rica\*

cgMLST type	CC	Serogroup	No. isolates (%)			Food type	Isolation years	Genetic resistance traits
			Total, n = 92	Clinical, n = 16	Nonclinical, n = 76			
L1-SL2-ST2-CT2715	CC2	IVb	8 (9)	4 (25)	4 (5)	Dairy, meat	2009, 2011, 2013, 2016–2017	bcrABC, qacA, LGI-2
L1-SL2-ST2-CT6120	CC2	IVb	10 (11)	2 (13)	8 (9)	Dairy	2010, 2013, 2016, 2018–2019	qacA, LGI-2
L1-SL2-ST2-CT2718	CC2	IVb	5 (5)	1 (6)	4 (5)	Dairy	2016, 2019	qacA
L1-SL2-ST1251-CT2780	CC2	IVb	3 (3)	1 (6)	2 (3)	Meat	2015–2016, 2018	qacA
L1-SL3-ST3-CT2730	CC3	IIb	9 (10)	0	9 (10)	Fish, meat	2016	bcrABC, SSI-1
L1-SL2-ST2-CT6072	CC2	IVb	5 (5)	0	5 (7)	Dairy	2019	LGI-2
L1-SL2-ST1627-CT6041	CC2	IVb	5 (5)	0	5 (7)	Dairy	2018–2019	LGI-2
L1-SL87-ST847-CT65	CC87	IIb	2 (2)	0	2 (3)	Meat	2016, 2019	NA
L1-SL3-ST1262/ST2762-CT2781	CC3	IIb	2 (2)	0	2 (3)	Dairy	2013	SSI-1
L1-SL5-ST5-CT2793	CC5	IIb	2 (2)	0	2 (3)	Fish, meat	2016	bcrABC, SSI-1
L1-SL2-ST2-CT2762	CC2	IVb	2 (2)	0	2 (3)	Mushrooms	2011	LGI-2

\*Table includes types detected in this study comprising ≥2 isolates with a cutoff of 7 allelic differences, n = 11/48. CC, clonal complex; cgMLST, core-genome multilocus sequence typing; LGI, *Listeria* genomic island; NA, not applicable; SSI, stress survival islet.

overrepresentation of SL2/CC2 and SL3/CC3 in Costa Rica (Appendix), which could be related to country's geographic location, climatic peculiarities, commercial trends, or natural reservoirs.

At the strain level, we identified 48 CTs, of which 44 (92%) were not previously reported. Eleven (23%) CTs included multiple isolates at a cutoff of 7 allelic differences of 1,748 cgMLST loci (2) (Table; Figure; Appendix Figures 1–3). Eight isolates were cgMLST type L1-SL2-ST2-CT2715, which accounted for 25% of clinical cases and spanned 9 years (Table).

Most human cases were associated with dairy products (Table). However, tracing to confirm the source of infection was not possible because most production is conducted by local farmers, often without traceability or attribution to the site of production.

Fresh cheese production is an economic staple in Costa Rica, and previous studies have reported *L. monocytogenes* detection in those products (7). Results from this study also show detection of identical strains of cgMLST type L1-SL2-ST2-CT6072 along the same production line, from raw materials to the final product, suggesting inadequate sanitation contributes to contamination (9).

*L. monocytogenes* is problematic for the food industry because it can survive and multiply under adverse environmental conditions (10). In this study, 90% of isolates carried ≥1 genetic element encoding for tolerance to disinfectants or stress. Markers of tolerance to disinfectants included qacA (51%, n = 47), bcrABC (23%, n = 21), and emrC (1%, n = 1). In addition, isolates had stress survival islet (SSI) genes, including SSI-1, conveying tolerance to low pH and high salt concentrations (21%, n =

19), and SSI-2 conveying tolerance to high pH and oxidative stress (1%, n = 1), as well as *Listeria* genomic island (LGI) genes, including LGI-2 (50%, n = 48) and LGI-3 (1%, n = 1) conveying tolerance to metals. Those tolerances can make *L. monocytogenes* elimination from production sites more difficult.

This study provides insight into the diversity of *L. monocytogenes* strains circulating in Central America and can aid national reference institutions in promoting regulatory changes to guarantee mandatory listeriosis reporting. In addition, institutions should establish mechanisms to provide low-cost microbiologic analysis. We also recommend regular sampling of risk products and training of artisanal processors.

In conclusion, strengthened WGS surveillance in Costa Rica could assist in controlling *L. monocytogenes* and provide food producers with information on strain diversity and effective means of eradication. WGS surveillance also would enable authorities to detect outbreaks and trace sources of contamination.

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This work includes multilocus sequence typing profiles publicly available on BIGSdb-*Listeria* (<https://bigsdb.pasteur.fr/listeria>).

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## References

- Swaminathan B, Gerner-Smidt P. The epidemiology of human listeriosis. *Microbes Infect.* 2007;9:1236–43. <https://doi.org/10.1016/j.micinf.2007.05.011>
- Moura A, Criscuolo A, Pouzeele H, Maury MM, Leclercq A, Tarr C, et al. Whole genome-based population biology and epidemiological surveillance of *Listeria monocytogenes*. *Nat Microbiol.* 2016;2:16185. <https://doi.org/10.1038/nmicrobiol.2016.185>
- Chenal-Francisque V, Lopez J, Cantinelli T, Caro V, Tran C, Leclercq A, et al. Worldwide distribution of major clones of *Listeria monocytogenes*. *Emerg Infect Dis.* 2011;17:1110–2. <https://doi.org/10.3201/eid1706.101778>
- Maury MM, Tsai YH, Charlier C, Touchon M, Chenal-Francisque V, Leclercq A, et al. Uncovering *Listeria monocytogenes* hypervirulence by harnessing its biodiversity. *Nat Genet.* 2016;48:308–13. <https://doi.org/10.1038/ng.3501>
- Schmid D, Allerberger F, Huhulescu S, Pietzka A, Amar C, Kleta S, et al. Whole genome sequencing as a tool to investigate a cluster of seven cases of listeriosis in Austria and Germany, 2011–2013. *Clin Microbiol Infect.* 2014;20: 431–6. <https://doi.org/10.1111/1469-0691.12638>
- Calvo-Arrieta K, Matamoros-Montoya K, Arias-Echandi ML, Huete-Soto A, Redondo-Solano M. Presence of *Listeria monocytogenes* in ready-to-eat meat products sold at retail stores in Costa Rica and analysis of contributing factors. *J Food Prot.* 2021;84:1729–40. <https://doi.org/10.4315/JFP-21-020>
- Posada-Izquierdo GD, Mazón-Villegas B, Redondo-Solano M, Huete-Soto A, Víquez-Barrantes D, Valero A, et al. Modelling the effect of salt concentration on the fate of *Listeria monocytogenes* isolated from Costa Rican fresh cheeses. *Foods.* 2021;10:1722. <https://doi.org/10.3390/foods10081722>
- Leclercq A, Chenal-Francisque V, Dieye H, Cantinelli T, Drali R, Brisson S, et al. Characterization of the novel *Listeria monocytogenes* PCR serogrouping profile IVb-v1. *Int J Food Microbiol.* 2011;147:74–7. <https://doi.org/10.1016/j.ijfoodmicro.2011.03.010>
- Castro H, Jaakkonen A, Hakkinen M, Korkeala H, Lindström M. Occurrence, persistence, and contamination routes of *Listeria monocytogenes* genotypes on three Finnish dairy cattle farms: a longitudinal study. *Appl Environ Microbiol.* 2018;84:e02000-17. <https://doi.org/10.1128/AEM.02000-17>
- Buchanan RL, Gorris LGM, Hayman MM, Jackson TC, Whiting RC. A review of *Listeria monocytogenes*: an update on outbreaks, virulence, dose-response, ecology, and risk assessments. *Food Control.* 2017;75:1–13. <https://doi.org/10.1016/j.foodcont.2016.12.016>

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## Human *Taenia martis* Neurocysticercosis, Switzerland

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Neurocysticercosis is almost exclusively caused by *Taenia solium* tapeworms. We describe a case of neurocysticercosis in Switzerland caused by infection with *Taenia martis*, the marten tapeworm, and review all 5 published cases of human infection with the marten tapeworm. In epidemiologically nonplausible cases of neurocysticercosis, zoonotic spillover infections should be suspected.

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# Genome-Based Characterization of *Listeria monocytogenes*, Costa Rica

## Appendix

### Materials and Methods

#### Bacterial Isolation

The study included 92 *L. monocytogenes* isolates previously collected from different regions throughout Costa Rica (Figure in main manuscript; Appendix Table 1) and spanning eleven years (2009–2019). Clinical isolates ( $n = 16$ ) were obtained by the Institute for Research and Teaching in Nutrition and Health (INCIENSA) in the context of the activities of the Costa Rican National Clinical Laboratory Network, comprised by laboratories of major hospitals, clinical settings, and the University of Costa Rica (UCR). Even though *L. monocytogenes* is not a notifiable disease in the country, there is an established criterium where bacterial pathogens that cause meningitis are sent to INCIENSA for confirmation of identification as well as antibiotic susceptibility tests. Isolates from food and food-production environments ( $n = 76$ ) were obtained by the National Laboratory of Veterinary Services (LANASEVE) of National Animal Health Service (SENASA), which performs microbiology analysis for the surveillance of food safety in products of animal origin for human consumption, by the INCIENSA that also monitors the microbiological quality of food for human consumption, and by the UCR and the Instituto Tecnológico de Costa Rica (ITCR) in the scope of research projects and/or routine analyses for customers, as well as in the framework of a specific study to investigate *Listeria* spp. strains previously isolated by private entities regulated and accredited by the international standard INTE/ISO/IEC 17025:2017 for microbiological analysis that guarantee the results of their sampling process.

Bacterial isolation was performed as described previously, following either the Bacteriological Analytical Manual method for *Listeria* isolation (1), for isolates obtained by the UCR, ITCR and SENASA, or the ISO 11290–1:2017 method (2), for isolates obtained by the INCIENSA.

Isolate identification was performed by proteomic analysis by matrix-assisted laser desorption/ionization time-of-flight (MALDI-TOF) mass spectrometry, using the MicroFlex LT system with MBT library DB-5989 (Bruker Daltonics, Bremen, Germany), as previously described (3).

#### **DNA Extraction and Genome Sequencing**

Isolates were cultured in Brain Heart Infusion Broth (BHI, Oxoid, Basingstoke UK) at 35°C overnight before use. DNA extraction was performed with the DNeasy Blood & Tissue Kit (QIAGEN, København Ø, Denmark-confirmed), according to the instructions provided by the manufacturer. Qubit fluorometer (Thermo FisherScientific, Waltham, MA, USA) was used to evaluate DNA quantity and purity. Library preparation was performed with the Nextera XT DNA Sample Kit (Illumina, San Diego, CA, USA), and DNA sequencing was carried out on a NextSeq 500 platform (Illumina) using 2× 150-bp paired-end runs. Reads were trimmed using fqCleanER v.21.10 (<https://gitlab.pasteur.fr/GIPhy/fqCleanER>), and assemblies were obtained using SPAdes v.3.14.0 (4) and polished with Pilon v.1.23 (5).

#### **In Silico Molecular Typing**

PCR-serogroups (6), multilocus sequence types (MLST) (7), core genome MLST (cgMLST) (8), and virulence and resistance profiles (8–14) were extracted from draft assemblies using BIGSdb-*Lm* (8,15) and BLASTN algorithm, as described before (8).

Minimum spanning trees were obtained from MLST and cgMLST profiles using BioNumerics v.7.6 (Applied-Maths, Sint-Martens-Latem, Belgium). MLST analyses also included 351 publicly available *L. monocytogenes* profiles from neighboring countries, obtained from BIGSdb-*Lm* (<http://bigsdb.pasteur.fr/listeria>; accessed on 16 February 2023). cgMLST-based dendograms were built in BioNumerics v.7.6.3 (Applied Maths, Sint-Martens-Latem, Belgium) using categorical differences and the single-linkage clustering method, and visualized with iTOL v.5 (16).

## Data Availability

Sequence data was made publicly available in NCBI/EBI/DDJJ databases (BioProject no. PRJEB20026).

## Supplementary References

1. US Food and Drug Administration. Testing methodology for *Listeria* species or *L. monocytogenes* in environmental samples. College Park (MD): The Administratio; 2015.
2. International Organization for Standardization. ISO 11290–1:2017 Microbiology of the food chain—Horizontal method for the detection and enumeration of *Listeria monocytogenes* and of *Listeria* spp., 2nd edition. Geneva: The Organization; 2017.
3. Thouvenot P, Vales G, Bracq-Dieye H, Tessaud-Rita N, Maury MM, Moura A, et al. MALDI-TOF mass spectrometry-based identification of *Listeria* species in surveillance: A prospective study. J Microbiol Methods. 2018;144:29–32. [PubMed](#) <https://doi.org/10.1016/j.mimet.2017.10.009>
4. Bankevich A, Nurk S, Antipov D, Gurevich AA, Dvorkin M, Kulikov AS, et al. SPAdes: a new genome assembly algorithm and its applications to single-cell sequencing. J Comput Biol. 2012;19:455–77. [PubMed](#) <https://doi.org/10.1089/cmb.2012.0021>
5. Walker BJ, Abeel T, Shea T, Priest M, Abouelliel A, Sakthikumar S, et al. Pilon: an integrated tool for comprehensive microbial variant detection and genome assembly improvement. PLoS One. 2014;9:e112963. [PubMed](#) <https://doi.org/10.1371/journal.pone.0112963>
6. Doumith M, Buchrieser C, Glaser P, Jacquet C, Martin P. Differentiation of the major *Listeria monocytogenes* serovars by multiplex PCR. J Clin Microbiol. 2004;42:3819–22. [PubMed](#) <https://doi.org/10.1128/JCM.42.8.3819-3822.2004>
7. Ragon M, Wirth T, Hollandt F, Lavenir R, Lecuit M, Le Monnier A, et al. A new perspective on *Listeria monocytogenes* evolution. PLoS Pathog. 2008;4:e1000146. [PubMed](#) <https://doi.org/10.1371/journal.ppat.1000146>
8. Moura A, Criscuolo A, Pouseele H, Maury MM, Leclercq A, Tarr C, et al. Whole genome-based population biology and epidemiological surveillance of *Listeria monocytogenes*. Nat Microbiol. 2016;2:16185. [PubMed](#) <https://doi.org/10.1038/nmicrobiol.2016.185>
9. Ryan S, Begley M, Hill C, Gahan CG. A five-gene stress survival islet (SSI-1) that contributes to the growth of *Listeria monocytogenes* in suboptimal conditions. J Appl Microbiol. 2010;109:984–95. [PubMed](#) <https://doi.org/10.1111/j.1365-2672.2010.04726.x>

10. Palma F, Brauge T, Radomski N, Mallet L, Felten A, Mistou MY, et al. Dynamics of mobile genetic elements of *Listeria monocytogenes* persisting in ready-to-eat seafood processing plants in France. *BMC Genomics*. 2020;21:130. [PubMed](https://doi.org/10.1186/s12864-020-6544-x) <https://doi.org/10.1186/s12864-020-6544-x>
11. Elhanafi D, Dutta V, Kathariou S. Genetic characterization of plasmid-associated benzalkonium chloride resistance determinants in a *Listeria monocytogenes* strain from the 1998–1999 outbreak. *Appl Environ Microbiol*. 2010;76:8231–8. [PubMed](https://doi.org/10.1128/AEM.02056-10) <https://doi.org/10.1128/AEM.02056-10>
12. Mereghetti L, Quentin R, Marquet-Van Der Mee N, Audurier A. Low sensitivity of *Listeria monocytogenes* to quaternary ammonium compounds. *Appl Environ Microbiol*. 2000;66:5083–6. [PubMed](https://doi.org/10.1128/AEM.66.11.5083-5086.2000) <https://doi.org/10.1128/AEM.66.11.5083-5086.2000>
13. Lee S, Ward TJ, Jima DD, Parsons C, Kathariou S. The arsenic resistance-associated *Listeria* genomic island LGI2 exhibits sequence and integration site diversity and a propensity for three *Listeria monocytogenes* clones with enhanced virulence. *Appl Environ Microbiol*. 2017;83:e01189-17. [PubMed](https://doi.org/10.1128/AEM.01189-17) <https://doi.org/10.1128/AEM.01189-17>
14. Harter E, Wagner EM, Zaiser A, Halecker S, Wagner M, Rychli K. Stress survival islet 2, predominantly present in *Listeria monocytogenes* strains of sequence type 121, is involved in the alkaline and oxidative stress responses. *Appl Environ Microbiol*. 2017;83:e00827-17. [PubMed](https://doi.org/10.1128/AEM.00827-17) <https://doi.org/10.1128/AEM.00827-17>
15. Jolley KA, Maiden MCJ. BIGSdb: scalable analysis of bacterial genome variation at the population level. *BMC Bioinformatics*. 2010;11:595. [PubMed](https://doi.org/10.1186/1471-2105-11-595) <https://doi.org/10.1186/1471-2105-11-595>
16. Letunic I, Bork P. Interactive Tree Of Life (iTOL) v5: an online tool for phylogenetic tree display and annotation. *Nucleic Acids Res*. 2021;49(W1):W293–6. [PubMed](https://doi.org/10.1093/nar/gkab301) <https://doi.org/10.1093/nar/gkab301>
17. Moura A, Lefrancq N, Wirth T, Leclercq A, Borges V, Gilpin B, et al.; Listeria CC1 Study Group. Emergence and global spread of *Listeria monocytogenes* main clinical clonal complex. *Sci Adv*. 2021;7:eabj9805. [PubMed](https://doi.org/10.1126/sciadv.abj9805) <https://doi.org/10.1126/sciadv.abj9805>
18. Moura A, Tourdjman M, Leclercq A, Hamelin E, Laurent E, Fredriksen N, et al. Real-time whole-genome sequencing for surveillance of *Listeria monocytogenes*, France. *Emerg Infect Dis*. 2017;23:1462–70. [PubMed](https://doi.org/10.3201/eid2309.170336) <https://doi.org/10.3201/eid2309.170336>
19. Halbedel S, Prager R, Fuchs S, Trost E, Werner G, Flieger A. Whole-genome sequencing of recent *Listeria monocytogenes* isolates from Germany reveals population structure and disease clusters. *J Clin Microbiol*. 2018;56:e00119-18. [PubMed](https://doi.org/10.1128/JCM.00119-18) <https://doi.org/10.1128/JCM.00119-18>

**Appendix Table 1.** Isolate metadata and genome metrics of *Listeria monocytogenes* sequenced in this study (n = 92).

Isolate no.	No. bases after filtering	Coverage	No. contigs	Total length, bp	N50, bp	% GC	% cgMLST loci tagged	Source type	Sample type	Geographic location	Isolation year	Serogroup	Lineage	Clonal complex, MLST	Sublineage, cgMLST	cgMLST type	BIGSdb ID
CLIP 2017/00419	2.40E+08	83	38	3.01E+06	1.94E+05	37.8	99.7	F	Sausage	San José	2016	IVb	I	CC1	SL1	CT2786	ID102753
CLIP 2019/02752	3.82E+08	132	30	2.99E+06	2.62E+05	37.82	99.8	F	Cheese	Vara Blanca	2019	IVb	I	CC1	SL1	CT333	ID102785
CLIP 2019/02623	4.20E+08	145	43	3.04E+06	1.92E+05	37.8	99.9	H	CSF	San José	2018	IVb	I	CC1	SL1	CT333	ID102774
CLIP 2019/02615	5.11E+08	176	38	3.02E+06	2.55E+05	37.81	99.9	F	Cheese	San José	2019	IVb	I	CC1	SL1	CT6042	ID102766
CLIP 2017/00404	3.05E+08	105	58	3.06E+06	1.43E+05	37.82	99.9	F	Cheese	ND	2011	IVb	I	CC2	SL2	CT2715	ID102739
CLIP 2017/00409	2.52E+08	87	110	3.05E+06	7.45E+04	37.85	99.3	F	Cheese	Cartago	2013	IVb	I	CC2	SL2	CT2715	ID102744
CLIP 2017/00417	5.93E+08	205	49	3.01E+06	1.39E+05	37.88	99.8	F	Sausage	San José	2016	IVb	I	CC2	SL2	CT2715	ID102751
CLIP 2019/02742	5.20E+08	179	47	3.17E+06	1.98E+05	37.7	99.8	F	Cheese	Upala	2017	IVb	I	CC2	SL2	CT2715	ID102775
CLIP 2017/00398	4.34E+08	150	58	3.17E+06	1.50E+05	37.69	99.8	H	Blood	Guanacaste†	2013	IVb	I	CC2	SL2	CT2715	ID102733
CLIP 2017/00399	4.18E+08	144	51	3.16E+06	1.32E+05	37.69	99.8	H	Blood	Limón†	2013	IVb	I	CC2	SL2	CT2715	ID102734
CLIP 2017/00400	3.59E+08	124	40	3.10E+06	2.23E+05	37.75	99.8	H	Blood	Limón†	2013	IVb	I	CC2	SL2	CT2715	ID102735
CLIP 2017/00403	5.40E+08	186	52	3.17E+06	1.39E+05	37.69	99.8	H	CSF	Heredia†	2009	IVb	I	CC2	SL2	CT2715	ID102738
CLIP 2017/00391	4.72E+08	163	51	3.04E+06	1.61E+05	37.84	99.9	H	Blood	Cartago†	2016	IVb	I	CC2	SL2	CT2716	ID102726
CLIP 2017/00405	4.46E+08	154	38	3.05E+06	1.39E+05	37.83	99.9	H	Blood	Limón†	2010	IVb	I	CC2	SL2	CT2717	ID102740
CLIP 2019/02751	4.38E+08	151	34	3.06E+06	2.73E+05	37.84	99.9	F	Cheese	Vara Blanca	2019	IVb	I	CC2	SL2	CT2718	ID102784
CLIP 2019/02753	3.36E+08	116	29	3.06E+06	2.72E+05	37.84	99.9	F	Cheese	San Ramón	2019	IVb	I	CC2	SL2	CT2718	ID102786
CLIP 2019/02754	3.03E+08	105	26	2.97E+06	3.33E+05	37.9	99.9	F	Cheese	San Ramón	2019	IVb	I	CC2	SL2	CT2718	ID102787
CLIP 2019/02756	2.54E+08	87	58	3.01E+06	1.44E+05	37.88	99.8	F	Cheese	Zarcero	2019	IVb	I	CC2	SL2	CT2718	ID102789
CLIP 2017/00396	5.98E+08	206	51	3.01E+06	1.41E+05	37.88	99.9	H	CSF	San José	2016	IVb	I	CC2	SL2	CT2718	ID102731
CLIP 2017/00407	4.45E+08	153	58	3.06E+06	1.52E+05	37.83	99.9	H	CSF	Cartago†	2010	IVb	I	CC2	SL2	CT2719	ID102742
CLIP 2017/00390	4.06E+08	140	42	3.04E+06	2.02E+05	37.81	99.9	H	CSF	Alajuela†	2016	IVb	I	CC2	SL2	CT2720	ID102725
CLIP 2017/00392	5.23E+08	180	59	3.03E+06	1.52E+05	37.81	99.8	H	Blood	Alajuela†	2013	IVb	I	CC2	SL2	CT2721	ID102727
CLIP 2017/00401	3.46E+08	119	38	2.96E+06	1.50E+05	37.85	99.8	F	Mushroom	ND	2011	IVb	I	CC2	SL2	CT2762	ID102736
CLIP 2017/00402	2.98E+08	103	42	2.99E+06	1.38E+05	37.81	99.8	F	Mushroom	ND	2011	IVb	I	CC2	SL2	CT2762	ID102737
CLIP 2017/00426	2.63E+08	91	55	3.04E+06	1.87E+05	37.84	99.9	F	Cheese	San José†	2013	IVb	I	CC2	SL2	CT2779	ID102760
CLIP 2017/00420	1.59E+08	55	87	3.05E+06	9.85E+04	37.83	99.9	F	Sausage	San José	2016	IVb	I	CC2	SL2	CT2780	ID102754
CLIP 2017/00421	2.61E+08	90	59	3.05E+06	1.53E+05	37.83	100	F	Sausage	San José	2015	IVb	I	CC2	SL2	CT2780	ID102755
CLIP 2019/02622	3.75E+08	129	48	3.01E+06	2.38E+05	37.86	99.9	H	Blood	San José	2018	IVb	I	CC2	SL2	CT2780	ID102773
CLIP 2017/00413	5.63E+08	194	61	3.08E+06	1.29E+05	37.79	99.9	F	Cheese	San José†	2013	IVb	I	CC2	SL2	CT2787	ID102747
CLIP 2019/02613	4.08E+08	141	41	3.00E+06	1.73E+05	37.86	99.9	F	Cheese	San José	2019	IVb	I	CC2	SL2	CT6040	ID102764
CLIP 2019/02610	4.27E+08	147	67	3.05E+06	1.45E+05	37.79	99.9	F	Cheese	San José	2019	IVb	I	CC2	SL2	CT6041	ID102761
CLIP 2019/02612	3.79E+08	131	62	3.11E+06	1.61E+05	37.86	99.9	F	Cheese	San José	2019	IVb	I	CC2	SL2	CT6041	ID102763
CLIP 2019/02616	3.38E+08	117	65	3.05E+06	1.11E+05	37.79	99.8	F	Cheese	San José	2019	IVb	I	CC2	SL2	CT6041	ID102767
CLIP 2019/02618	3.65E+08	126	82	3.05E+06	1.10E+05	37.79	99.9	F	Cheese	San José	2019	IVb	I	CC2	SL2	CT6041	ID102769
CLIP 2019/02750	4.94E+08	170	50	3.05E+06	1.98E+05	37.79	99.9	F	Cheese	Turrialba	2018	IVb	I	CC2	SL2	CT6041	ID102783
CLIP 2019/02621	5.80E+08	200	51	3.08E+06	1.61E+05	37.84	99.9	H	Blood	San José	2018	IVb	I	CC2	SL2	CT6045	ID102772
CLIP 2019/02611	3.92E+08	135	76	3.09E+06	1.98E+05	37.8	99.9	F	Cheese	San José	2019	IVb	I	CC2	SL2	CT6046	ID102762
CLIP 2019/02617	3.55E+08	122	65	3.03E+06	1.26E+05	37.84	99.9	F	Cheese	San José	2019	IVb	I	CC2	SL2	CT6047	ID102768
CLIP 2019/02745	3.90E+08	135	35	2.97E+06	2.40E+05	37.91	99.9	F	Cheese	Turrialba	2018	IVb	I	CC2	SL2	CT6063	ID102778
CLIP 2019/02748	4.82E+08	166	43	3.04E+06	2.71E+05	37.85	99.9	F	Cheese	Turrialba	2018	IVb	I	CC2	SL2	CT6065	ID102781
CLIP 2019/02757	3.04E+08	105	68	3.10E+06	2.39E+05	37.79	99.7	F	Cheese	San Isidro	2019	IVb	I	CC2	SL2	CT6066	ID102790
CLIP 2019/02744	4.94E+08	170	51	3.04E+06	2.38E+05	37.85	99.8	F	Cheese	Capellades	2018	IVb	I	CC2	SL2	CT6067	ID102777
CLIP 2019/02749	4.85E+08	167	40	3.04E+06	1.59E+05	37.85	99.9	F	Cheese	Santa Cruz	2018	IVb	I	CC2	SL2	CT6069	ID102782
CLIP 2019/02747	4.08E+08	141	74	3.07E+06	1.98E+05	37.83	99.9	F	Cheese	Turrialba	2018	IVb	I	CC2	SL2	CT6070	ID102780
CLIP 2019/02764	7.56E+08	261	84	3.16E+06	3.21E+05	37.76	99.8	F	Raw milk	Turrialba	2019	IVb	I	CC2	SL2	CT6072	ID102797
CLIP 2019/02765	7.81E+08	269	59	3.12E+06	2.06E+05	37.79	99.9	F	Cheese	Turrialba	2019	IVb	I	CC2	SL2	CT6072	ID102798

Isolate no.	No. bases after filtering		Coverage	No. contigs	Total length, bp	N50, bp	% GC	% cgMLST loci tagged	Source type	Sample type	Geographic location	Isolation year	Serogroup	Lineage	Clonal complex, MLST	Sublineage, cgMLST	cgMLST type	BIGSdb ID
	No. bases after filtering	Coverage																
CLIP 2019/02759	1.24E+09	426	120	3.11E+06	3.60E+05	38	99.9	PE	Drain	Turrialba	2019	IVb	I	CC2	SL2	CT6072	ID102792	
CLIP 2019/02760	7.47E+08	258	47	3.12E+06	3.60E+05	37.77	99.9	PE	Tank	Turrialba	2019	IVb	I	CC2	SL2	CT6072	ID102793	
CLIP 2019/02761	7.06E+08	243	62	3.15E+06	3.63E+05	37.81	99.9	PE	Cooling chamber	Turrialba	2019	IVb	I	CC2	SL2	CT6072	ID102794	
CLIP 2017/00372	1.76E+08	61	90	3.12E+06	9.53E+04	37.69	99.9	PE	Soil	Cartago†	2016	IVb	I	CC2	SL2	CT6116	ID102721	
CLIP 2017/00424	5.53E+08	191	32	3.04E+06	1.98E+05	37.81	99.9	F	Cheese	Heredia	2013	IVb	I	CC2	SL2	CT6117	ID102758	
CLIP 2019/02755	3.66E+08	126	37	3.04E+06	3.55E+05	37.84	99.9	F	Cheese	Zarcero	2019	IVb	I	CC2	SL2	CT6118	ID102788	
CLIP 2017/00412	2.08E+08	72	60	3.06E+06	1.35E+05	37.8	99.8	F	Cheese	Cartago	2013	IVb	I	CC2	SL2	CT6119	ID102746	
CLIP 2017/00411	2.77E+08	96	57	3.08E+06	2.81E+05	37.8	99.8	F	Cheese	Cartago	2013	IVb	I	CC2	SL2	CT6120	ID102745	
CLIP 2017/00415	2.78E+08	96	51	3.04E+06	1.55E+05	37.84	99.9	F	Cheese	Cartago†	2013	IVb	I	CC2	SL2	CT6120	ID102749	
CLIP 2017/00418	2.74E+08	94	48	3.04E+06	1.50E+05	37.84	99.9	F	Cheese	San José	2013	IVb	I	CC2	SL2	CT6120	ID102752	
CLIP 2017/00423	2.62E+08	90	63	3.08E+06	1.39E+05	37.8	99.9	F	Cheese	Cartago	2013	IVb	I	CC2	SL2	CT6120	ID102757	
CLIP 2019/02614	2.43E+08	84	90	3.08E+06	9.05E+04	37.81	99.4	F	Cheese	San José	2019	IVb	I	CC2	SL2	CT6120	ID102765	
CLIP 2019/02746	3.45E+08	119	59	3.08E+06	1.61E+05	37.81	99.8	F	Cheese	Turrialba	2018	IVb	I	CC2	SL2	CT6120	ID102779	
CLIP 2019/02762	6.86E+08	236	475	3.27E+06	2.73E+05	39.67	99.8	F	Cheese	Turrialba	2019	IVb	I	CC2	SL2	CT6120	ID102795	
CLIP 2019/02763	9.18E+08	317	136	3.12E+06	3.58E+05	38.07	99.9	F	Raw milk	Turrialba	2019	IVb	I	CC2	SL2	CT6120	ID102796	
CLIP 2017/00393	4.76E+08	164	59	3.08E+06	1.50E+05	37.8	99.9	H	Blood	Cartago†	2016	IVb	I	CC2	SL2	CT6120	ID102728	
CLIP 2017/00408	4.09E+08	141	66	3.08E+06	1.36E+05	37.81	99.9	H	CSF	Cartago†	2010	IVb	I	CC2	SL2	CT6120	ID102743	
CLIP 2017/00406	6.63E+08	228	60	3.07E+06	1.38E+05	37.79	99.9	H	CSF	Cartago†	2015	IVb	I	CC2	SL2	CT6121	ID102741	
CLIP 2017/00389	3.43E+08	118	41	3.05E+06	1.39E+05	37.83	99.9	F	Meat	ND	2016	IVb	I	CC2	SL2	CT6122	ID102724	
CLIP 2019/02758	8.26E+08	285	31	2.91E+06	4.33E+05	37.91	99.8	F	Cheese	Naranjo	2019	IVb	I	CC217	SL217	CT222	ID102791	
CLIP 2017/00395	3.03E+08	104	41	3.00E+06	1.95E+05	37.79	99.9	F	Sausage	ND	2016	IVb	I	CC6	SL6	CT2761	ID102730	
CLIP 2017/00383	3.27E+08	113	59	3.03E+06	1.55E+05	37.82	99.9	F	Shrimp	San José†	2016	IIb-v1	I	ST1079	SL1079	CT1669	ID102722	
CLIP 2017/00357	3.37E+08	116	53	3.14E+06	1.58E+05	37.75	99.8	F	Ground beef	San José†	2016	IIb	I	CC3	SL3	CT1674	ID102707	
CLIP 2017/00366	3.20E+08	110	51	3.10E+06	1.97E+05	37.79	99.9	PE	Drain	San José	2016	IIb	I	CC3	SL3	CT1674	ID102716	
CLIP 2017/00358	2.38E+08	82	45	3.05E+06	1.34E+05	37.81	99.8	F	Chicken wings	San José	2016	IIb	I	CC3	SL3	CT2730	ID102708	
CLIP 2017/00360	3.67E+08	127	43	3.05E+06	1.86E+05	37.81	99.8	F	Pork	Heredia	2016	IIb	I	CC3	SL3	CT2730	ID102710	
CLIP 2017/00361	5.29E+08	183	44	3.05E+06	1.87E+05	37.81	99.8	F	Pork	San José†	2016	IIb	I	CC3	SL3	CT2730	ID102711	
CLIP 2017/00363	4.02E+08	139	51	3.05E+06	1.42E+05	37.81	99.7	F	Cheese	San José	2016	IIb	I	CC3	SL3	CT2730	ID102713	
CLIP 2017/00364	4.97E+08	171	44	3.05E+06	1.69E+05	37.81	99.7	F	Ground beef	San José†	2016	IIb	I	CC3	SL3	CT2730	ID102714	
CLIP 2017/00365	2.15E+08	74	43	3.05E+06	1.97E+05	37.81	99.8	F	Tilapia	Guanacaste†	2016	IIb	I	CC3	SL3	CT2730	ID102715	
CLIP 2017/00359	3.82E+08	132	40	3.05E+06	3.71E+05	37.81	99.8	PE	Drain	Alajuela	2016	IIb	I	CC3	SL3	CT2730	ID102709	
CLIP 2017/00370	3.05E+08	105	40	3.05E+06	1.97E+05	37.81	99.8	PE	Drain	San José	2016	IIb	I	CC3	SL3	CT2730	ID102719	
CLIP 2017/00371	5.02E+08	173	42	3.05E+06	2.36E+05	37.81	99.7	PE	Drain	San José	2016	IIb	I	CC3	SL3	CT2730	ID102720	
CLIP 2017/00422	2.31E+08	80	39	3.04E+06	2.53E+05	37.92	99.7	F	Cheese	Cartago	2013	IIb	I	CC3	SL3	CT2781	ID102756	
CLIP 2017/00425	2.84E+08	98	39	3.04E+06	1.97E+05	37.93	99.7	F	Cheese	Cartago	2013	IIb	I	CC3	SL3	CT2781	ID102759	
CLIP 2017/00367	2.62E+08	90	79	3.11E+06	2.36E+05	37.79	99.9	PE	Cooling chamber	San José	2016	IIb	I	CC5	SL5	CT2783	ID102717	
CLIP 2017/00362	4.62E+08	159	74	3.11E+06	1.35E+05	37.77	99.9	F	Tilapia	San José	2016	IIb	I	CC5	SL5	CT2793	ID102712	
CLIP 2017/00368	5.08E+08	175	59	3.10E+06	1.53E+05	37.79	99.9	F	Beef	Heredia†	2016	IIb	I	CC5	SL5	CT2793	ID102718	
CLIP 2017/00414	2.13E+08	73	77	2.93E+06	1.20E+05	37.86	99.4	F	Cheese	Heredia†	2013	IIb	I	CC506	SL506	CT2776	ID102748	
CLIP 2019/02619	3.78E+08	130	39	2.97E+06	3.04E+05	37.88	99.9	F	Chorizo	Heredia	2019	IIb	I	CC87	SL87	CT6044	ID102770	
CLIP 2017/00416	2.36E+08	81	38	2.92E+06	1.99E+05	37.91	99.9	F	Sausage	San José	2016	IIb	I	CC87	SL87	CT65	ID102750	
CLIP 2019/02620	3.96E+08	137	39	2.97E+06	2.09E+05	37.88	99.9	F	Chorizo	San José	2019	IIb	I	CC87	SL87	CT65	ID102771	
CLIP 2017/00397	4.04E+08	139	28	3.09E+06	3.73E+05	37.87	100	F	Sausage	ND	2016	IIc	II	CC9	SL9	CT13239	ID102732	
CLIP 2017/00388	2.98E+08	103	44	3.04E+06	2.02E+05	37.8	100	F	Pork	ND	2016	IIc	II	CC9	SL9	CT1668	ID102723	

Isolate no.	No. bases after filtering	Coverage	No. contigs	Total length, bp	N50, bp	% GC	% cgMLST loci tagged	Source type	Sample type	Geographic location	Isolation year	Serogroup	Lineage	Clonal complex, MLST	Sublineage, cgMLST	cgMLST type	BIGSdb ID
CLIP 2017/00394	3.41E+08	117	38	3.09E+06	2.01E+05	37.79	100	F	Frozen vegetables	ND	2016	Ila	II	CC121	SL121	CT909	ID102729
CLIP 2019/02743	3.86E+08	133	18	2.83E+06	3.82E+05	37.93	99.9	F	Cheese	Turrialba	2018	Ila	II	CC19	SL378	CT6064	ID102776

\*BIGSdb, Bacterial Isolate Genome Sequence Database (17); cgMLST, core-genome multilocus sequence typing; F, food; H, human; ID, identification; MLST, multilocus sequence typing; ND, not done; PE, production environment.

†Geographic information available only at the level of the province.

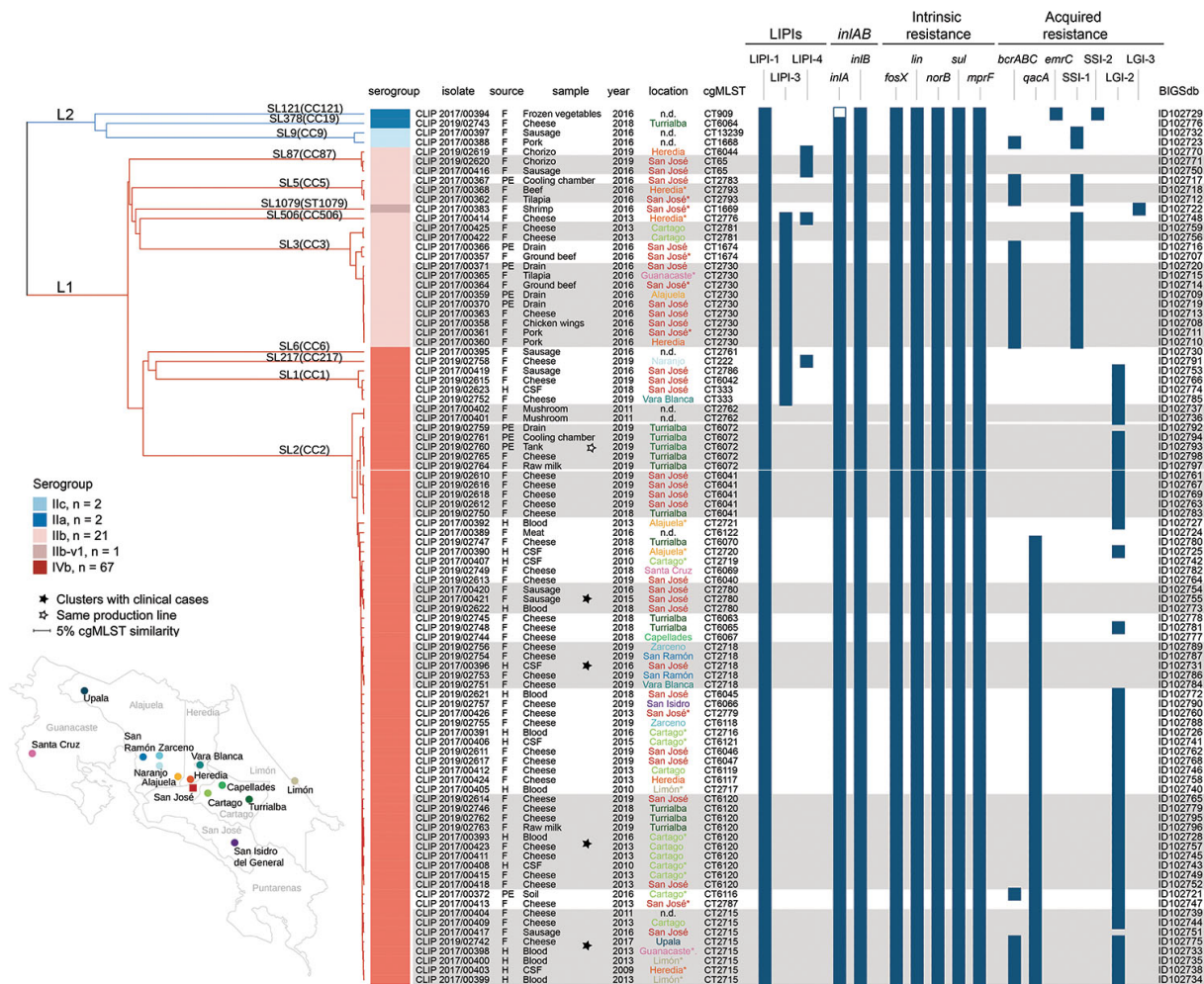
**Appendix Table 2.** Previously reported cgMLST types detected in this study (cutoff of 7 or less allelic differences out of 1748 cgMLST loci, Institut Pasteur scheme)

cgMLST Type (CC, serogroup)	Source (item)		Studies from other countries	Source lab	NCBI/EBI/DDJJ accession no.	Reference no.
	This study	Other studies				
<b>Lineage I</b>						
L1-SL217-ST217-CT222 (CC217, IVb)	F (dairy)	H, F (salad)	US	CDC	SRR1021894, SRR1027089	(8)
L1-SL1-ST1-CT333 (CC1, IVb)	H, F (dairy)	H	US	CDC	SRR1043171, SRR7057542	(8,17)
L1-SL87-ST847-CT65 (CC87, IIb)	F (meat)	F (avocado)	MX	FDA	SRR975360	(8)
<b>Lineage II</b>						
L2-SL121-ST121-CT909 (CC121, IIa)	F (frozen vegetables)	H, F, PE, FE, A	CL, DK, FR, DE, LV, NO, PL, NL, UK, US	KMAHVH, ANSES, ECDC, IP, UWLMO, UKHSA, FDA, RKI	ERR1304231, ERR1738638, ERR1738650, ERR2522041, ERR2522263, ERR2522276, ERR2522284, ERR2522292, ERR2522294, ERR2522295, ERR2522297, ERR2522312, ERR2522337, ERR2522338, ERR2522353, ERR2522363, ERR2522371, ERR2522810, ERR3040059, ERR3040061, ERR3040067, ERR3040072, ERR3040073, ERR3040075, ERR3040076, ERR3040078, ERR3040083, ERR3040104, ERR4176463, ERR4176497, ERR4176536, ERR4176606, ERR4176666, ERR4176794, ERR4648214, SRR10753542, SRR10753574, SRR10753599, SRR10753600, SRR10753602, SRR11004717, SRR13072765, SRR14783257, SRR15245555, SRR15245949, SRR20053048, SRR2040689, SRR4052010, SRR4052014, SRR4052068, SRR4052071, SRR4052072, SRR4052073, SRR4052161, SRR4124934, SRR4124940, SRR5133499, SRR5318940,	(8,10,18,19)

cgMLST Type (CC, serogroup)	Source (item)		Studies from other countries	Source lab	NCBI/EBI/DDJJ accession no.	Reference no.
	This study	Other studies				
			SRR5526029, SRR5526031, SRR5526035, SRR5526085, SRR5526092, SRR5526094, SRR5526097, SRR5526099, SRR5526101, SRR5526105, SRR5526120, SRR5526126, SRR5526128, SRR5526130, SRR5526135, SRR5526137, SRR5526138, SRR5526142, SRR5526145, SRR5526155, SRR5647016, SRR5647027, SRR5647029, SRR5647030, SRR6966182, SRR7403111, SRR7410624, SRR7429735, SRR7429781, SRR7440567, SRR7440606, SRR7440615, SRR7440618, SRR7440621, SRR7440634, SRR7440649, SRR7440688, SRR7440921, SRR7441075, SRR7441146, SRR7441209, SRR7441250, SRR7441287, SRR7547848, SRR7827106, SRR7828078, SRR7839361, SRR7841130, SRR7841155, SRR7841205, SRR7842337, SRR7850202, SRR7850362, SRR7850428, SRR7850452, SRR7866225, SRR7866376, SRR7866514, SRR7866571, SRR7866588, SRR7866636, SRR7866672, SRR7866727, SRR7866750, SRR7866891, SRR7873595, SRR7873692, SRR7873717,			

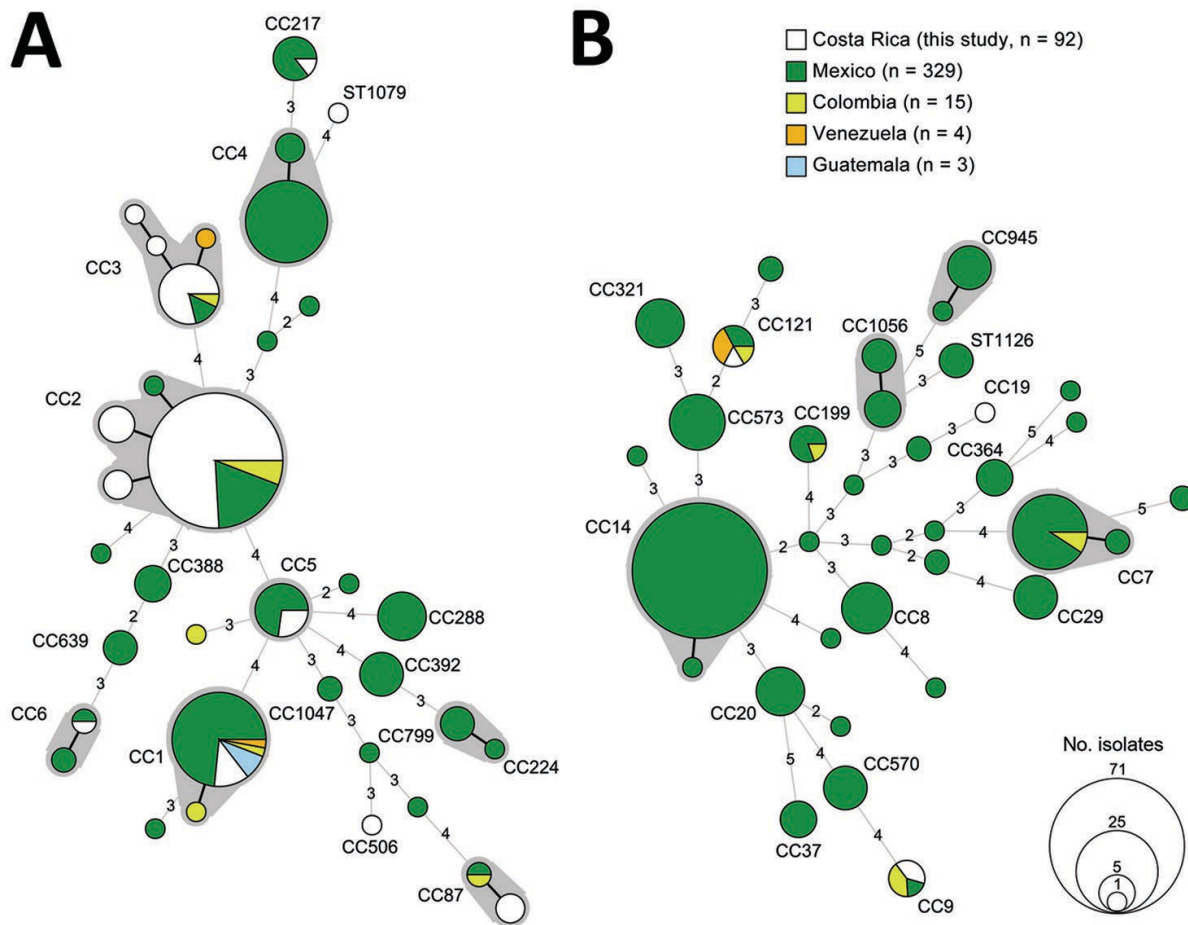
cgMLST Type (CC, serogroup)	Source (item)		Studies from other countries	Source lab	NCBI/EBI/DDJJ accession no.	Reference no.
	This study	Other studies				
SL21(CC121)					SRR7873744, SRR8235628, SRR8239064, SRR8239220, SRR8239239, SRR8655304, SRR8979699, SRR9004322, SRR9010038	
SL378(CC19)						
SL9(CC9)						
SL87(CC87)						
SL5(CC5)						
SL1079(ST1079)						
SL506(CC506)						
SL3(CC3)						
SL2(CC2)						
SL6(CC6)						
SL217(CC217)						
SL1(CC1)						

\*A, animal; ANSES, Agence Nationale de Sécurité Sanitaire de L'Alimentation, de L'Environnement et du Travail, FR; CC, clonal complex; CDC, US Centers for Disease Control and Prevention; CL, Chile; DE, Germany; DK, Denmark; ECDC, European Centre for Disease Prevention and Control; F, food; FDA, US Food and Drug Administration; FE, farm environment; FR, France; H, human; IP, Institut Paster, FR; KMAHVH, Klinisk Mikrobiologisk Afdeling, DK; LV, Latvia; MX, Mexico; NO, Norway; PE, food production environment; PL, Poland; RKI, Robert Koch Institut, DE; UK, United Kingdom; UKHSA, UK Health Security Agency, UK; US, United States; UWLMO, University of Warmia and Mazury in Olsztyn, PL.

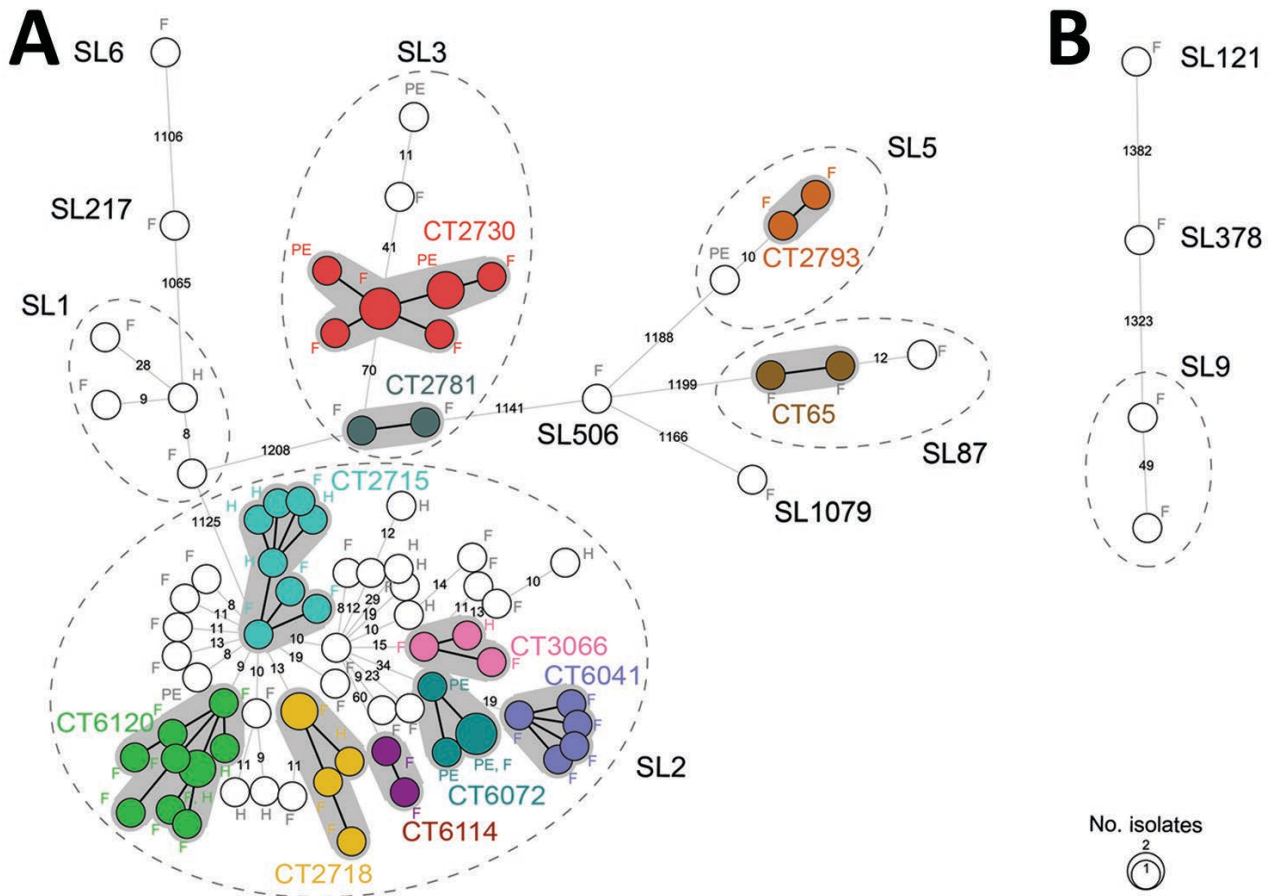


**Appendix Figure 1.** Single linkage dendrogram of 92 isolates generated for genome-based characterization of *Listeria monocytogenes*, Costa Rica. Dendrogram based on cgMLST allelic profiles (1,748-locus scheme). Branches are colored according to lineages: L1, red; L2, blue. Branches are labeled according to lineages, sublineages, and clonal complexes. Information on isolates' name, serogroup, source, sample type, cgMLST type, year of isolation, location, and BIGSdb IDs are provided in the columns. Colors in location column correspond to dots on map. Gray bars indicate clusters of isolates

with  $\leq 7$  allelic differences out of 1,748 cgMLST loci. presence of selected virulence and resistance genetic traits in each isolate is represented by squared dark blue boxes and empty boxes denote genes with premature stop codons. BIGSdb, Bacterial Isolate Genome Sequence Database (<https://bigsdb.pasteur.fr>); CC, clonal complex; cgMLST, core-genome multilocus sequence typing; F, food; H, human; ID, identification; L, lineage; LIPI, listeria pathogenicity island; SL, sublineage; ST, sequence type.



**Appendix Figure 2.** Minimum spanning tree of 92 *Listeria monocytogenes* isolates, Costa Rica, 2009–2019. Trees are based on cgMLST allelic profiles (7-locus scheme). A) Lineage I isolates; B) lineage 2 isolates. Publicly available *L. monocytogenes* isolates (n = 351) from neighboring countries in the Caribbean region were also included. Circles represent different profiles and sizes are proportional to the number of isolates within. Branch lengths are proportional to the allelic differences between the profiles which are indicated in the branches. For simplicity, allelic differences of 1 are omitted and represented by thicker branch lines. Clonal complexes with >1 profile are surrounded by gray shading and labeled if detected in this study or if they contain  $\geq 5$  isolates. CC, clonal complex; cgMLST, core-genome multilocus sequence typing; ST, sequence type.



**Appendix Figure 3.** Minimum spanning tree of 92 *Lysteria monocytogenes* isolates, Costa Rica, 2009–2019. Trees are based on cgMLST allelic profiles (1,748-locus scheme). A) Lineage I isolates; B) lineage II isolates. Circles represent different profiles and sizes are proportional to the number of isolates within. Labels next to circles indicate the source of isolates. Branch lengths are proportional in logarithmic scale to allelic differences between profiles, which are also indicated in the branches. For simplicity, allelic differences of  $\leq 7$  are omitted and represented by thicker branch lines. Clusters with  $>1$  profile are highlighted in colors; labels correspond with cgMLST type and are delimited by gray shadows. Dashed ellipses delimitate sublineages with  $>1$  isolate and labeled with corresponding sublineage. cgMLST, core-genome multilocus sequence typing; CT, cgMLST type; F, food; H, human, PE; production environment; SL, sublineage.