Lineage-specific evolution and gene flow in *Listeria monocytogenes* are independent of bacteriophages

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Summary

*Listeria monocytogenes* is a foodborne pathogen causing systemic infection with high mortality. To allow efficient tracking of outbreaks a clear definition of the genomic signature of a cluster of related isolates is required, but lineage-specific characteristics call for a more detailed understanding of evolution. In our work, we used core genome MLST (cgMLST) to identify new outbreaks combined to core genome SNP analysis to characterize the population structure and gene flow between lineages. Whilst analysing differences between the four lineages of *L. monocytogenes* we have detected differences in the recombination rate, and interestingly also divergence in the SNP differences between sub-lineages. In addition, the exchange of core genome variation between the lineages exhibited a distinct pattern, with lineage III being the best donor for horizontal gene transfer. Whilst attempting to link bacteriophage-mediated transduction to observed gene transfer, we found an inverse correlation between phage presence in a lineage and the extent of recombination. Irrespective of the profound differences in recombination rates observed between sub-lineages and lineages, we found that the previously proposed cut-off of 10 allelic differences in cgMLST can be still considered valid for the definition of a foodborne outbreak cluster of *L. monocytogenes*.

Introduction

*Listeria monocytogenes*, one of the most important foodborne pathogens, is the causative agent of listeriosis, an infection that particularly affects the elderly, immuno-compromised individuals, and pregnant women (Radoshevich and Cossart, 2017; Lamond and Freitag, 2018). *Listeria monocytogenes* infections can progress to life threatening conditions including septicaemia, meningitis, encephalitis and spontaneous abortion, resulting in a high mortality rate (EFSA and ECDC, 2017). In recent years, a number of large listeriosis outbreaks have been seen worldwide, including the latest one reported in 2017–2018 in South Africa, which was caused by the consumption of contaminated ready-to-eat processed meat products from one company (WHO, 2018; Smith et al., 2019). This means that surveillance of outbreaks is a major concern for national and international authorities. With the advent of low-cost whole-genome sequencing in the past few years, outbreak analysis has shifted from the previous gold standard of PFGE analysis to core genome multilocus sequencing (cgMLST) (Lüth et al., 2018). cgMLST is based upon just the genes present in the core genome of a species, or at least those present in the panel of genomes available for analysis (Ruppitsch et al., 2015). The 1701-gene cgMLST scheme (Ruppitsch et al., 2015) has a much higher capacity for resolution of isolates differences than the classical 7-locus MLST (Salcedo et al., 2003; Ragon et al., 2008). As such cgMLST is not only able to provide insight into overall population structure but is also able to contribute to short-term nomenclature and the analysis of outbreaks. In the specific case of *L. monocytogenes*, the cut-off value for outbreak detection was fixed to ≤10 cgMLST alleles difference (Ruppitsch et al., 2015). More recently after analysis of isolates with a known epidemiological link, it was proposed that...
this cut-off value should be reduced to ≤7 cgMLST allele differences (Moura et al., 2016).

Reference laboratories favour cgMLST analysis over other genomic analyses due to the easier standardization of the cgMLST scheme; however, analysis of population structure both at the outbreak and at the lineage scales also benefits from the analysis of how SNPs and recombination contribute to bacterial evolution. This is of particular interest as L. monocytogenes in not a homogeneous species but contain four separate deep-branching lineages as is evident from both MLST work and more recent global genomic analyses (Haase et al., 2014; Moura et al., 2016; Lees et al., 2019). When testing for point mutations and recombination using the 7-locus MLST data, it was found that the occurrence of recombination versus point mutations was about six times more frequent in lineage II than in lineage I (den Bakker et al., 2008). However, the knowledge is limited on population structure within sub-lineages (SL) and the gene flow between lineages at the finer scale.

Horizontal gene transfer mechanisms alter the core and accessory genome and are an important driving force in shaping bacterial population structures. With transformation not being documented for L. monocytogenes (Boreeze et al., 2000), transduction, which is well documented in many bacterial species (Popa et al., 2017), automatically gains relevance as a candidate mechanism for gene transfer. In this species interest in phages also has practical aspects due to the differences in lineage-specific (teichoic acids) and genus-specific (cell wall) phage adsorption targets (Dunne et al., 2018). Furthermore, there is significant research in the utilization of phages as a biocontrol in the food industry (Carlon et al., 2005; Hagens and Loessner, 2014). With respect to the actual contribution of bacteriophages to gene flow within Listeria, there is conflicting evidence. In addition to the relevance of phages in the shaping of the accessory genome of L. monocytogenes (Klumpp and Loessner, 2013; Kuenne et al., 2013), there is clear evidence that phages are responsible for the transfer virulence-associated genes from other species to Listeria (Chen and Novick, 2009). With this, phages have been implicated with facilitating core-genome gene transfer by generalized transduction possibly with distinct impact in distinct lineages (Hodgson, 2000; Orsi et al., 2011). Contrary to this, the short size of homologous recombination events detected does not support transduction as the main transfer mechanism (den Bakker et al., 2008).

In the present work, we identify new outbreak isolates from a panel of genome sequences from Swiss L. monocytogenes clinical and food isolates by using cgMLST scheme, and then we used core SNPs to characterize differences in the population structure and the gene flow at a high resolution of the lineages I, II and III.

Methods

Bacterial isolates

The draft genome sequences of 158 L. monocytogenes isolates identified during surveillance by the Institute for Food Safety and Hygiene, the Swiss National Reference Laboratory, at the University of Zurich in the Swiss region of Europe mainly from 2011 to 2014 are presented in this work (Table S1). The addition of two lineage III isolates and six further outbreak isolates (Weinmaier et al., 2013; Tasara et al., 2015, 2016) creates a total database of 166 Swiss isolates. Ninety-six of the isolates were from human clinical cases (Althaus et al., 2014) and 66 isolates were isolated from a variety of food products (Ebner et al., 2015), one was from the environment and three were of unknown origin. Metadata for the isolates, including geographic location, collection data and source of the isolation, are listed in Table S1. For the comparative genomic analysis, we included 414 isolates from Germany (Halbedel et al., 2018) and 128 Dutch isolates (Kremer et al., 2017; Lees et al., 2019) (Table S2) making a total of 708 genomes. We used the German isolates because it was our reference study for the outbreaks analysis and we included the Dutch isolates because it was our reference study for the PopPUNK analysis. In addition, for the population structure and gene flow analysis, we included 36 lineage III and four lineage IV isolates from public databases (Table S3).

Serootyping of the Swiss isolates was performed using the commercial set of Listeria O-factor and H-factor antisera from Denka Seiken (Pharma Consulting, Burgdorf, Switzerland) according to the manufacturer’s instructions.

Genome sequencing

DNA extraction for the 160 Swiss isolates was performed from pure cultures of each isolate grown in Brain Heart Infusion. In brief, 1.5 ml of overnight culture was pelleted, washed with 100% acetone and re-suspended in 400 μl (10 mM Tris–HCl, 10 mM EDTA, pH 7), 10 μl lysozyme (10 mg/ml, Sigma UK) and 10 μl RNase A (10 mg/ml, Sigma UK) and incubated at 37°C for 30 min. Then, 75 μl of 10% SDS (Sigma UK) and 10 μl proteinase K (10 mg/ml, Sigma UK) were added and incubated at 65°C for 30 min. DNA was then purified using the Genomic Clean and Concentrator kit (Zymo Research), using the manufacturer’s instructions. 150 bp paired-end DNA libraries, with an average insert size of 451 bp were produced using the NEB Ultra II custom kit and sequenced in pools on the HiSeq X10 platform; this resulted in an average of 2 666 434 reads per sample, corresponding to an average of 100x coverage per genome. Adapter and poor quality sequences were removed using Trimmomatic (Ver. 0.36) (Bolger et al., 2014) and the genomes were assembled using SPAdes (Ver. 3.9.0, default parameters) (Bankevich et al., 2012) and the assembly

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statistics for each isolate are found in Table S1. For the
downstream analysis assemblies were only included if they
had a minimum depth coverage of 10-fold. Raw sequence
files and draft genomes in contig level for all 160 Swiss
isolates were submitted to the European Nucleotide Archive
(https://www.ebi.ac.uk/ena) under study accession number
PRJNA486730 (Table S1). In silico, MLST was performed
using a BLAST-based tool (https://github.com/tseemann/
mst) on de novo genome assemblies (Jolley and Maiden,
2010; Seemann, 2018).

cgMLST and SNP analysis

A set of 1701 genes well defined for cgMLST analysis
(Ruppitsch et al., 2015; Halbedel et al., 2018) were used for
the identification of cluster type, SL and lineages in a total of
708 isolates (Tables S1 and S2). The alleles for the 1701-loci
cgMLST were downloaded from the database https://www.
cgmlst.org/ncs/schema/690488/ and used as a query in
BLASTn in order to identify and extract the target gene
sequence. The parameters used in BLASTn were 100% of
query sequence coverage required, word size 11, mismatch penalty −3,
match reward 1, gap open costs 5, and gap extension costs 2 and the hit with the highest % identity were kept. The
sequences for each target gene per genome were extracted
using bedtools v2.25.0 (https://bedtools.readthedocs.io),
which creates a new FASTA file for each extracted
sequence. Using a custom-made bash script, a multi-FASTA
file was obtained for each target gene, and then the genes
were aligned using Muscle v3.8.31 (Edgar, 2004).

The evaluation of the cgMLST scheme was done using
standard pairwise allelic mismatches and SNPs pairwise.
The matrix of pairwise allelic mismatches per target gene
was obtained from the previously aligned sequences using
snp-dists v0.6 (https://github.com/tseemann/snp-dists),
where any differences (SNPs and/or indels) was scored
as 1 and no differences was scored as 0. Using a python
custom script, the matrices of allele pairwise differences were combined obtaining the overall allele differences in
1701-loci for each pairwise comparison. Gene absence
was scored as a missing value. A total of 1596 (93.8%) target genes were present in all 708 isolates and this set of
genes was used for the SNP pairwise analysis. A total of
1596 core genes from the gene-by-gene alignment were
concatenated according to the order of the reference
genome EGD-e (NC_003210.1) and then the matrix of
SNPs pairwise differences was obtained using the snp-
dists v0.6. The maximum-likelihood core genome phyloge-
etic tree was inferred from the concatenated alignment
core genes using the GTR model in RAxML (Randomized
Accelerated Maximum Likelihood) (version 8.2.12)
(Stamatakis, 2014). The visualization, manipulation and
annotation of the phylogenetic tree were done using the
ggtree R package v1.15.6 (Yu et al., 2017).

PopPUNK analysis

Clustering analysis of 708 isolates was performed using
the PopPUNK tool, which uses a K-mer approach to
find genetic distances between isolates and then identify
clusters of closely related isolates (Lees et al., 2019).
PopPUNK was run on the genome assemblies using the
default settings (lower k-mer length was 13). Through
PopPUNK we identified clusters of isolates, which were
then compared with the clustering obtained from the phy-
logenetic analysis that was based on SNPs.

Chromosome painting and fineSTRUCTURE analysis

In order to detect genome-wide transfer of DNA
sequence chunks through homologous recombination
events between lineages, we expanded our initial dataset
(708 isolates) with additional genomes from lineages III
(36 isolates) and IV (four isolates) taken from public data-
bases (Table S2) making a total of 748 isolates. We inferred
population structure at a finer scale among the isolates
from the core-genome SNPs data by using ChromoPainter
(version 0.04) and fineSTRUCTURE (version 2.1.1).
(Lawson et al., 2012). Before executing the analysis, we
removed isolates belonging to the same CC to generate
a dataset with a single randomly selected genome for each
CC. The genome collection for this analysis now contained
81 isolates with 15 isolates belonging to lineage I, 33 to line-
age II, 29 to lineage III and four to lineage IV. In this genome
collection, the genetic distance was more than 3000 SNPs
between the single core-genomes (1569 core genes). The
genome sequences of the isolates were mapped against the
complete reference genome N2306 (CP011004)
through the SKA tool (Harris, 2018), thus the alignment of
the whole genome was obtained. ChromoPainter infers
where DNA ‘chunks’ have been donated from a donor to
a recipient and reconstructs the chromosome haplotype of
‘recipient’ individuals as a series of chunks from the other
‘donor’ individuals in the sample. A ‘chunk’ refers to a
set of neighbouring/linked SNPs copied from a donor to a
recipient. The output from ChromoPainter is a ‘co-ancestry
matrix’, which summarizes the expected number of chunks
of DNA imported from a donor to a recipient genome.
fineSTRUCTURE uses the output of ChromoPainter
for the clustering of individuals based on the co-ancestry
matrix (Lawson et al., 2012). ChromoPainter and
fineSTRUCTURE were used according to the procedure
described in http://www.paintmychromosomes.com.

Recombination rate

The recombination rate in the core genome alignment
was assessed with Gubbins Version: 2.3.1 (Croucher
et al., 2015). Gubbins was run independently using
separate datasets for lineage I (371 isolates), lineage II (335 isolates) and lineage III (38 isolates). The whole-genome alignment was done by using the SKA tool (Harris, 2018) and complete reference genomes were used for the alignment; LL195 (HF558398) from Lineage I, Lm3136 (CP013723) from lineage II and M7 (CP002816) from lineage III. We did not estimate the recombination for lineage IV due to the small sample size of genomes deposited (four isolates). Reference-based alignments of the core genome were analysed using the default parameter of Gubbins, which identifies regions in the alignment with an atypically high density of SNPs, and that infers hypothetical recombination events. Two measures of the recombination rate were estimated: \( \rho/\theta \) estimate the relative frequency of occurrence of recombination and mutation in the history of the lineage (Milkman and Bridges, 1990), and \( r/m \) assess the relative impact of recombination and mutation in the genetic diversification of the lineage (Guttman and Dykhuizen, 1994).

Transduction in silico analysis

Listeria monocytogenes prophages were identified in the set of 81 isolates (see above) using the PHASTER tool, which provides the phage sequence annotation as either intact, questionable or incomplete (Zhou et al., 2011; Amrdt et al., 2016). The incomplete prophages were removed for further analysis. The intact and questionable prophages were validated against a well-characterized set of known prophages evaluated using MASH v2.1 (Ondov et al., 2016). This set drew on an additional 45 L. monocytogenes prophages available from a public database (http://millardlab.org/bioinformatics/bacteriophage-genomes). Through the Mash analysis, we obtained a distance matrix to produce a heatmap of similarity between the prophages, and to determine whether questionable prophage was prophage or host chromosomal sequence. The same analysis allowed us to identify those prophages from a public database with no similarity with the prophage in our dataset. The mapping of the prophages to the set of 81 isolates was done by BLASTn. The identity threshold was 72% and the coverage value was recorded and plotted as a heatmap to quantify the length of the prophage region. To test in silico the effect of phages on competence, we mapped the known insertion of phages into the core genome using the default parameter of Gubbins, which identifies regions in the alignment with an atypically high density of SNPs, and that infers hypothetical recombination events. Two measures of the recombination rate were estimated: \( \rho/\theta \) estimate the relative frequency of occurrence of recombination and mutation in the history of the lineage (Milkman and Bridges, 1990), and \( r/m \) assess the relative impact of recombination and mutation in the genetic diversification of the lineage (Guttman and Dykhuizen, 1994).

Results

The genome sequences of the 166 Swiss L. monocytogenes isolates

The species Listeria monocytogenes is subdivided into four phylogenetic lineages (I–IV) (Ward et al., 2008; Orsi et al., 2011). Here, we present the draft genome sequences of L. monocytogenes isolates identified during surveillance by the Institute for Food Safety and Hygiene, the Swiss National Reference Laboratory, at the University of Zurich in the Swiss region of Europe mainly from 2011 to 2014 (Table S1). The mean length of the draft genomes was 2 948 113 bp with a mean GC% content of 37.8%. The summary data of the assembly for each genome are reported in Table S1. Core genome phylogenetic analysis of the 166 Swiss L. monocytogenes isolates found 51 isolates belonged to lineage I and 113 to lineage II (Fig. 1; Table S1). No significant association of the two lineages was found with respect to the origin (clinical or food) of the isolates (clinical isolates: lineage I 36 isolates, lineage II 60 isolates; food isolates: lineage I 14 isolates and lineage II 52 isolates) (Fig. 1; Table S1). Phenotypic testing was used to confirm the serotypes (1/2a, 1/2b, 1/2c, 3a, 3b, 3c, 4a, 4b, 4c and 4d) across the isolates. The two most prevalent serotypes were the lineage II serotype 1/2a (103 isolates) and the lineage I serotype 4b (38 isolates) (Fig. 1). MLST sequence type (ST) information showed ST1 (eight isolates) to be the most prevalent ST in lineage I and ST8 and ST121 (with seven and eight isolates respectively) in lineage II (Fig. 1; Table S1).

Genome evolution in lineages I and II

In recent years, outbreak analysis has shifted from PFGE to cgMLST and a difference of ≤10 cgMLST alleles was identified as the most appropriate cut off (Ruppitsch et al., 2015). To verify if the proposed cut off for outbreaks of ≤10 cgMLST differences was appropriate in our dataset, we included in the analysis recently published isolates from Germany (414 isolates) (Halbedel et al., 2018) and the Netherlands (128 isolates) (Kremer et al., 2017; Lees et al., 2019). To visualize the distribution and frequency of differences, we constructed a pairwise matrix of either the core genome SNPs or the cgMLST allele differences and plotted the data (Fig. 2). In the overall pairwise plots (Fig. 2E), we clearly identified differences that mapped to outbreaks, SL and lineages. When limiting the analysis to groups of isolates distinguished by fewer than 150 differences in cgMLST or SNPs (Fig. 2A and B) outbreaks were clearly identifiable on the plots both by analysing cgMLST allele difference and core genome SNPs. When choosing SNPs or cgMLST differences with the lowest frequency both approaches yielded the same cut-off of ≤12 differences separating outbreak isolates but if being more stringent cut-offs of ≤8 alleles and ≤7 SNP differences could also be considered (Fig. 2A and B). The pairwise analysis is plotted separately for diversity within sub-lineage (Fig. 2A and B), between sub-lineage within a lineage (Fig. 2C and D) and between lineages (Fig. 2E).
When we plotted the data separately for lineage I and lineage II isolates (red and blue respectively in Fig. 2) we noted differences in their genome diversity. Lineage I isolates were more diverse within sub-lineages than lineage II (Kruskal–Wallis test p-value <0.01), with a broader distribution of pairwise distances, and a less clear peak corresponding to putative outbreaks. In both lineages the outbreak cut off was comparable, but the sub-lineages had different population structures with lineage I having peaks at 36 and at 80 cgMLST differences while lineage II showed tighter distribution with peaks at 31 and 65 cgMLST differences (Fig. 2A). A comparable distribution was given by plotting SNP differences where lineage I had more SNPs within sub-lineages than lineage II (Fig. 2B).

When looking instead at the differences between SL within a lineage (cut off 1500 cgMLST alleles or 22 000 core genome SNPs) we observed the opposite arrangement with
lineage I showing less genome diversity than lineage II (Kruskal–Wallis test $p$-value <0.01) (Fig. 2C and D). cgMLST did not resolve this difference effectively as almost all the alleles were different; however, SNP differences showed dramatic diversity with lineage I between-sub-lineages differences of about 5000 SNPs and lineage II of about 15 000 SNPs differences. These differences became more evident when plotting SNPs versus a cgMLST differences (Fig. 2E). In conclusion, this suggests lineage II is composed of sub-lineages that are more highly divergent from one another than those within lineage I, but each lineage I sub-lineage contains slightly more diversity than its lineage II equivalent, which might complicate the inference of outbreaks in lineage I.

**Genome diversity within sub-lineages**

To better understand this potentially confounding within-lineage population structure, the genetic diversity of the collection was analysed using MLST and PopPUNK. This revealed a close correspondence between the cgMLST-identified sub-lineages and both MLST-defined clonal complexes (CCs) and PopPUNK clusters. The differences within sub-lineages were clearly defined by both cgMLST difference and SNPs but also mapped perfectly to clusters defined by PopPUNK (Lees et al., 2019) (Fig. S1). To explore the rationale for the multiple peaks in the cgMLST and SNP analysis within SL, we plotted the values separately for the different CC/PopPUNK.
clusters/sub-lineages and PopPUNK clusters match to CC. In lineages I, CC1 included 100 isolates (89 of which were ST1), CC2 included 60 isolates (58 of which were ST2), CC5 included 56 isolates (55 of which were ST5 and one ST1063) and CC6 included 111 isolates (110 of which were ST6) (Fig. 3). Irrespective of the number of outbreak isolates (those with less than 12 cgMLST allele differences), there were different levels of SNP diversity within these CC/PopPUNK clusters/sub-lineages (Fig. 3A and B). For example, CC6 had one peak with cgMLST difference of 35 alleles and a comparable number of SNPs (Fig. 3A and B). In contrast, both CC1 and CC2 showed two clearly distinct peaks, and CC5 had three peaks (Fig. 3A and B) indicating population structure at a finer level than sub-lineages. In CC1 there were several groups of isolates that differed by about 40–50 cgMLST alleles within the groups and which differed by 80–100 cgMLST alleles between the groups. Similarly, in CC2 there were a number of groups of isolates that differed by about 40 cgMLST alleles within the groups and differed by about 60 cgMLST alleles between the groups. The few isolates in clusters not belonging to either ST1 or ST2 could not account for this peculiar within the ST population structure. In lineage II the two main two clusters were CC403, which included 54 isolates (50 of which ST403) and a large CC8 cluster with 118 isolates (102 of which ST8) (Fig. 3E). The two lineage II CC analysed (Fig. 3C and D) showed a similar population structure with a median cgMLST difference of 30 alleles for CC8 and 31 alleles for CC403. When analysing the k-mer distance using PopPUNK, data on pairwise distance could be generated in addition to the core genome, also for the accessory genome (Fig. S2A,B), with both sets of data showing quite a good overlap.

**Swiss outbreaks**

In the 166 Swiss isolates, from clinical and food sources, we could identify 15 clusters of isolates with ≤ 12 cgMLST allele differences (Table 1). Nine cases were known outbreaks and six were clusters from food (Table 1). SNP differences correlated well to cgMLST data but often yielded fewer differences as our SNP data did not include indels and were calculated on just 1594 core genes, while cgMLST was performed on all 1701 genes. Two small clusters of two and three isolates respectively were from lineage I while all others mapped to lineage II with three clusters in CC8. In three instances isolates from human and food sources are placed in the same cluster.

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**Fig. 3.** Genome diversity within clonal complexes within lineages I and II. Pairwise cgMLST and SNP differences were analysed for the major clonal complexes for each lineage. For lineage I the distribution of the pairwise cgMLST (A) and SNP (B) differences are shown for clonal complexes CC1 (red), CC2 (cyan), CC5 (purple) and CC6 (green). For lineage II the distribution of the pairwise cgMLST (C) and SNP (D) differences are shown for CC8 (yellow) and CC403 (blue). The CCs shown in panels A–D are indicated on the core genome phylogenetic tree and the arrows indicate the respective PopPUNK clusters (orange and brown) plotted on the first ring (E). The complete legend and high-resolution image of this circular tree are in Fig. S1. [Color figure can be viewed at wileyonlinelibrary.com]
When we included the German and Dutch isolates, we observed that three ‘Swiss’ outbreak clusters actually also contained isolates from Switzerland and/or Germany and/or the Netherlands.

**Recombination**

The recombination rate of *L. monocytogenes* was analysed separately for lineages I, II and III using Gubbins on 371, 336 and 38 isolates respectively. The mean of the number of polymorphisms introduced via recombination with respect to mutation (r/m) and the number of recombination events with respect to polymorphisms introduced by mutation (ρ/θ) are reported for internal and terminal branches of the lineages (Table 2). There are differences in the mean recombination ratio between the internal and external branch for lineage I and III, but there are considerably fewer for lineage II, with recombination hotspots...
(Kuenne et al., 2013) readily detectable in the core genome (Figs S3–S5). In agreement with our data above, recombination has occurred more frequently in lineages II and III where the \( r/m \) rate was 40–33 times higher when compared with the lineage I (Table 2). Graphical representations of the blocks of recombination clearly underline the higher frequency of recombination and horizontal gene transfer in lineages II and III (Figs S3–S5).

**Table 2.** Recombination statistics among lineages.

| Lineages       | \( r/m^a \) | \( \rho/\theta^b \) | \( r/m^a \) | \( \rho/\theta^b \) |
|----------------|------------|-----------------|------------|-----------------|
| Lineage I (n = 371) | 0.0046 | 0.0002 | 0.0091 | 0.0013 | 0.0137 | 0.0015 |
| Lineage II (n = 336) | 0.3146 | 0.0149 | 0.2324 | 0.0075 | 0.5470 | 0.0224 |
| Lineage III (n = 39) | 0.1100 | 0.0038 | 0.3460 | 0.0114 | 0.4559 | 0.0151 |

\( ^a r/m \) is the ratio of base substitutions predicted to have been imported through recombination compared with those occurring through point mutations.

\( ^b \rho/\theta \) (rho/theta) is the ratio of the number of recombination events to point mutations, a measure of the relative rates of recombination and point mutation.

**Sequence exchange between lineages**

We characterized patterns of gene flow and population structure by using ChromoPainter and fineSTRUCTURE. To perform this analysis, we selected a panel of 81 isolates, with one isolate representing each CC, from all four lineages of *L. monocytogenes*. The co-ancestry matrix was visualized as a heatmap with each cell indicating the DNA ‘chunk’ counts that a recipient received from each donor (Fig. 5A). We used all isolates as potential donors in order to characterize the gene flow among the isolates. The boxes in Fig. 5A indicate evident signs of ‘gene flow’; we use the term ‘gene flow’ when there is a horizontal transfer of DNA chunks between lineages. The co-ancestry matrix revealed asymmetries in the gene flow between lineages; i.e., the four lineages imported higher numbers of chunks from other members of their own lineage. Both lineage I (\( p \)-value <0.05) and lineage II (\( p \)-value = 0.018) imported more chunk counts from lineage III than from lineages II and I respectively (Fig. S6A). There is a high genetic exchange from both lineages I and II to lineage III, with more chunk counts imported from lineage I (\( p \)-value <0.05) (Fig. S6A). Interestingly, despite the high frequency of exchanges observed overall, the analysis found little evidence for sequence exchange between lineages I and II (Fig. 5A, Fig. S6A). Lineages I and II donated more chunk counts to lineage III and lineage III donated more chunk counts to lineage II (\( p \)-value <0.05) (Fig. S6B). When we considered the total chunk counts of transferred DNA per isolate, we observed again that lineage I had fewer chunk counts from external lineage population, and that lineages

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II and III had more chunk numbers, which matched well with the lower and higher respective rates of recombination observed with Gubbins (Fig. 6A). Importantly the chunk counts also showed that each lineage predominantly imported DNA from the same lineage and that lineage III was the second ‘best’ donor in both lineages I and II (Fig. 6A). Here, we did not analyse lineage IV due to the small sample size.

**Horizontal gene transfers by transduction and transformation**

To test if bacteriophage were responsible for the horizontal gene transfer in our dataset, we mapped the prophages in our set of genomes, inferring that detection of similar phages or phage-remnants through the genomes of different CCs could be taken as a proxy for the transfer of DNA by transduction. We identified 41 intact and 47 questionable prophages across the 81 *L. monocytogenes* isolates (one representative for each CC) using PHASTER. After Mash analysis, through the heatmap of similarity, we identified 28 of the questionable prophages as chromosomal sequence and thus they were removed, also 31 lytic phages from the public database (Dorscht et al., 2009; Denes et al., 2014) were removed as none had similarity with our genomes (Table S4). We then used the remaining 60 prophages plus 13 temperate phages from the public database as a query in a BLASTn analysis with all

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81 isolates as the subject in order to predict sequence exchange between lineages mediated by prophage transduction. From BLASTn we obtained a phage coverage matrix, which was plotted as a heatmap and linked to the phylogenetic tree (Fig. 5B). Data showed that there is a high occurrence of prophages in both lineages I and II and a lower occurrence of prophages in lineage III. Some genomes, particularly in lineage II, showed extensive homology to many phages or phage-remnants present in lineages I and II. The prophages identified in lineage III had greater similarity to prophage of lineages I and II than with others in lineage III. A similar pattern was observed with the prophages from the public databases (Fig. 5B); however, for these, we had the limitation that we did not have access to the lineage data from the isolates from which they were originally sourced. When we quantified the total coverage of an individual phage throughout all isolates of a given lineage, we observed that lineage II phages scored much higher than lineage I (p-value <0.05), followed by lineage III (Fig. 6D, Fig. S7) indicating that phages from lineage II show much higher levels of sequence identity to the phages present in the other lineages. To check the impact of phage on competence for genetic transformation, we mapped the incidence of insertion of prophage into the competence regulator gene comK in the 81 isolates (Table S5). We detected a phage-interrupted comK gene in 2/15 isolates in lineage I, 10/33 isolates in lineage II, 1/29 in lineage III and 1/4 in lineage IV, however, none of these genomes showed any evidence of a reduction in DNA exchange (Fig. 5A).

When testing the gene flow between lineages using Chromopainter the data clearly showed a correlation between the number of chunks of exchanged DNA and their size (Fig. 6A–C). In each of the three lineages incoming DNA originating from the same lineage is detected more frequent (higher chunk counts) and larger (chunk size) (Fig. 6A–C). Of the three lineages, lineage III shows by far the highest numbers of recombination events (Fig. 6C; note different X-axis scale). Interestingly, when we compare incoming DNA from other lineages the data show that lineage III is also the main donor to both lineages I and II (Fig. 6A and B, Fig. S6A). Finally, through Pearson correlation, we confirm an inverse correlation (p = 0.023) between the prophage coverage and the chunk counts donated from lineages II and III to lineage I, and from I and III to lineage II.

Discussion

The cgMLST scheme set up for L. monocytogenes typing is currently the most widely accepted approach for the identification of outbreaks (Ruppitsch et al., 2015; Moura et al., 2016; Halbedel et al., 2018). This is because cgMLST has a greater discriminatory power when compared with MLST or PFGE (Lüth et al., 2018). In this study, we explored the cut-off for outbreak identification by performing a pairwise comparison with a 1701-locus cgMLST scheme (Ruppitsch et al., 2015) against 708 de novo assembled genomes using BLASTN. Our work analysed 158 genomes of Swiss human and food L. monocytogenes isolates (this study) to which we added, for comparison, six known Swiss outbreak isolates (Weinmaier et al., 2013; Tasara et al., 2015, 2016), 414 German human isolates (Halbedel et al., 2018) and 128 Dutch human isolates (Kremer et al., 2017; Lees et al., 2019). Our results show a cut-off for the definition of outbreaks between 7 and 12 allelic or SNPs differences. Using the same set of cgMLST target genes the cut-off for the determination of outbreak isolates was previously identified as ≤10 alleles differences by analysing 42 and 424 genomes respectively (Ruppitsch et al., 2015; Halbedel et al., 2018). In contrast, a lower threshold of ≤7 alleles was identified by Moura et al. (2016) using 1696 genomes with a 1748 gene cgMLST scheme. The use of different sets of cgMLST has not been found by others to be critical for the general identification of clonal groups or outbreak isolates (Chen et al., 2016). While we agree with this cut-off, especially when plotting all isolates on the same graph, it should be noted that there are also more subtle differences to be seen when mapping pairwise differences of different lineages and sub-lineages/clonal-complexes. Our data clearly show that dependent upon the sub-lineage analysed, the most likely cut-off could be anywhere between 7 and 12 cgMLST differences when using the 1701-locus cgMLST scheme (Fig. 3). Plotting core genome SNP differences shows the same pattern of variation when the different lineages and sub-lineages are plotted separately. Our pairwise data analysis, in common with previous observations (Chen et al., 2016), does not allow us to propose defining a clonal-complex specific cut-off, however, we would underline the necessity to consider the possibility of occurrence of outbreaks with a slightly higher cgMLST or SNP diversity. SNP analysis in core-genomes was used in previous work for the characterization of the genome diversity within SL but this was limited to CC1 and CC9 only (Moura et al., 2016). In this study, we have greatly expanded this and characterized the genome diversity within lineages for CC1, CC2, CC5 and CC6 for lineage I and CC8 and CC403 for lineage II.

While the web-based identification of CC is straightforward, the definition of SL is not as obvious. To identify a computationally more approachable method, we tested the PopPUNK program for clustering clonal isolates (Lees et al., 2019). Our data show that PopPUNK clustering matches perfectly to isolates grouped into CC. Importantly, this overlaps perfectly with the SL identified by cgMLST that are defined by having less than <150 alleles or SNPs differences (Moura et al., 2016). One important aspect is
that PopPUNK is much faster, more flexible and computationally less heavy for the definition of SL, which confirms it as an ideal tool for the analysis of large datasets (Lees et al., 2019).

Nine cases of listeriosis were recorded in Switzerland in 2011 as due to imported cooked ham (Hächler et al., 2013), and in this study, we have sequenced the genomes for the isolates from these cases. A follow-up study reported a genome sequence for the isolate N1546, which was associated with this listeriosis outbreak in 2011 (Tasara et al., 2015). In this study, we have identified a total of 11 isolates belonging to this outbreak; these include the seven isolates previously associated with the outbreak, in addition, four isolates newly reported in this study, three of which were from a food source (meat). Another isolate N2306, which is associated with the listeriosis outbreak in 2013–2014 (Tasara et al., 2016), also fell into a cluster with one of the newly sequenced isolates. Overall, using the cgMLST scheme and SNP approach, we have identified nine Swiss outbreaks of which, three included isolates with food and human sources. The fact that we have identified three outbreaks, which include isolates from Switzerland and Germany, shows that a posteriori genome analysis can highlight cross-border outbreaks, which may fail to be noted by national surveillance programs. In this case, inclusion of the large collection of French genomes (Moura et al., 2017) may well have shown other cross-border outbreaks.

The population structure in L. monocytogenes had been previously evaluated using the STRUCTURE program to analyse seven house-keeping genes. These studies on MLST loci found that the genetic diversity between lineages was due to mutation rather than recombination (Ragon et al., 2008; Haase et al., 2014). In contrast, previous work, which involved the examination of two virulence genes, had found that the rate of recombination was six times higher than the mutation rate (den Bakker et al., 2008). In our work, which used 1569 core genes, we have identified a higher recombination rate in lineages II and III, 40 and 33 times higher respectively, than in lineage I; these figures match well to the data of den Bakker (den Bakker et al., 2008). Our work is the first study to demonstrate that horizontal gene transfer occurs predominantly within a lineage, and that lineage III shows the highest rate of within-lineage exchange. In addition, lineage III was also observed to be the most proficient donor of DNA chunks. The surprisingly low gene flow between lineages I and II indicates a, so far, an unexplained barrier to horizontal gene transfer between these lineages. As restriction-modification systems in Listeria are specific to STs (Chen et al., 2017; Croix et al., 2017) these barriers to horizontal gene transfer cannot account for the observed lineage-specific DNA exchange. Bacteriophages have also been proposed to be responsible for DNA transfer in Listeria (Orsi et al., 2011). The preferential within-lineage transfer of DNA we have observed could have matched well to the lineage-specific serovars of wall teichoic acids to which Listeria phages adsorb and which therefore determine their serovar-dependent host range (Dunne et al., 2018). In contrast, our results from the prophages mapping showed the opposite findings, with lineage III harbouring almost no bacteriophages whilst being the most proficient acceptor and donor of core genome chunks. Given that the detection of phage sequences was inversely proportional to the amount of recombination the hypothesis that horizontal gene transfer in Listeria is mainly due to transduction had to be excluded. The observation of the preferential gene flow to and from lineage III in absence of phage detection would exclude for Listeria phage-related mechanisms as mayor contributors to core genome evolutions, as also for example in E. coli (Tree et al., 2014). This observation would also exclude other phage-related mechanisms of genome evolution in Listeria as for example the genome hypermobility through lateral gene transfer by transduction, as recently reported for the related species Staphylococcus aureus (Chen et al., 2018) for which phage-mediated gene transfer to Listeria had been documented (Chen and Novick, 2009). The potential contribution of the conjugation of plasmids and genomic islands to the horizontal gene flow was not evaluated here as their contribution to homologous recombination within the core genome is considered generally less significant.

The other hypothesis for horizontal gene transfer, i.e. that it most likely occurs through transformation, has been examined previously by analysis of MLST locus recombination based on the relatively short size of recombining DNA (den Bakker et al., 2008). The well-described insertion of phages into the competence regulator comK (Loessen et al., 2000; Kuenne et al., 2013) was detected in a third of lineage II isolates and few other isolates however, this did not correlate with an appreciable reduction in horizontal gene transfer in these isolates. It is important to underline that the phage insertion into comK in the isolates tested for gene flow was generally representative of all isolates (as they were selected randomly) of those specific CCs, thus representing a stable trait that could have a potential impact on CC evolution. This does not rule out transformation as the mechanism for gene transfer in the core genome because the comK-integrated phages have been shown to excise efficiently during stress allowing comK transcription and ComK-mediated gene regulation in conditions like intracellular infection of macrophages (Molloy, 2012; Rabinovich et al., 2012). The observation that the gene flow shows clearly that within lineage gene exchange is favoured over the transfer of DNA to or from other lineages is consistent with
homologous recombination, the main mechanism for horizontal gene transfer in the core genome, being facilitated by the lower diversity between isolates in the same lineage. While we have no direct evidence for the mechanism of core genome evolution, the fact that phage presence inversely correlates to recombination, indicates that transformation, even if it has never been experimentally tested in \textit{Listeria monocytogenes}, should be viewed as the most likely mechanism.

In conclusion, our data show important differences in the evolution of the four lineages of \textit{Listeria monocytogenes}, with differences in recombination rate and flow of core genome genes. Furthermore, the finding that phage presence inversely correlates with core genome gene flow indicates that transformation and not transduction is the most likely mechanism for the evolution of the core genome. From an epidemiological point of view, where the aim is for a clear definition of a cut-off for an outbreak, our work demonstrates that the proposed cut-offs of 10 cgMLST differences can still be considered valid irrespective of the different population structures found in the different \textit{Listeria monocytogenes} lineages.

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Data availability

The genome sequences were deposited as reads at the European Nucleotide Archive ENA with Study accession PRJEB17888 and accession 4421STDY6601709 to 4421STDY6601879 and as FASTA files at GenBank as Bioproject PRJNA486730 with the accession numbers QYDA0000000 to RDTB0000000.

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Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site:

**Table S1.** Swiss *L. monocytogenes* isolates.

**Table S2.** Germany and The Netherlands *L. monocytogenes* isolates.

**Table S3.** *L. monocytogenes* lineage III and IV isolates.

**Table S4.** *Listeria monocytogenes* bacteriophages.

**Appendix S1.** Lineage specific evolution and gene flow in *Listeria monocytogenes* are independent of bacteriophages.