Pulsed-field gel electrophoresis (PFGE) analysis of *Listeria monocytogenes* isolates from different sources and geographical origins and representative of the twelve serovars

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**Abstract**

Multiplex-PCR (MPCR) serogrouping and pulsed-field gel electrophoresis (PFGE) subtyping analysis are currently used by several public and private laboratories for the characterization of *Listeria monocytogenes*. In this study a set of 80 *L. monocytogenes* isolates belonging to the twelve serovars was used to investigate (i) the typeability of the rare serovars, (ii) the ability of PFGE analysis with *Apa*I and *Asc*I to differentiate serovars within MPCR serogroups and (iii) the association of molecular types with the specific source or geographical origin of the isolates. With the exception of three isolates (rare serovars 4a and 4c) that were not amenable to restriction with *Apa*I, all the other analyzed isolates were subtyped by both enzymes. PFGE discriminated the 80 isolates into 62 combined *Apa*I and *Asc*I PFGE patterns (pulsotypes), but could not differentiate serovars within MPCR serogroups, in which isolates from different serovars displaying the same pulsotype were found. Clustering analysis suggests that for some pulsotypes grouping according to Portuguese origin or source can be suggested. On the other hand, some *L. monocytogenes* clones are widely distributed. Two pulsotypes from Portuguese human isolates were identical to the ones displayed by human outbreak clones in the UK and in the USA and Switzerland, respectively, although they were not temporally matched.

Computer-assisted data analysis of large and diverse PFGE type databases will improve the correct interpretation of subtyping data in epidemiological studies and in tracing routes and sources of contamination in the food industry.

**Keywords:** *Listeria monocytogenes*; Sources; Geographical origin; Genetic diversity; Serovars; Genetic lineages; PFGE; Multiplex-PCR (MPCR) serogrouping

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**Introduction**

*Listeria monocytogenes* is a foodborne facultative intracellular pathogenic bacterium responsible for listeriosis in humans and animals, particularly among immunocompromised individuals. Thirteen serovars (1/2a, 1/2b, 1/2c, 3a, 3b, 3c, 4a, 4ab, 4b, 4c, 4d, 4e, 7)
have been reported for *L. monocytogenes* [5]. Some of these serovars have been recovered from patients and foods (1/2a, 1/2b, 1/2c and 4b) more frequently [14] than the others considered as rare serovars. Although serotyping [20] is less discriminatory than molecular typing methodologies, it has been used in epidemiological studies as a universal technique for the characterization of *L. monocytogenes*. Recently, different authors have proposed PCR based methods for serogrouping *L. monocytogenes* isolates [1,5,24].

The multiplex-PCR (MPCR) assay proposed by Doumith et al. [5] is based on the use of five sets of primers targeting group-specific genes that enable the clustering of *L. monocytogenes* strains into five phylogenetic groups (serogroups) correlated with the serovars (except serovar 4ab which is extremely rare) [5]. The set of primers targeting the *prs* gene (putative phosphoribosyl pyrophosphate synthetase) fragment (370 bp) is universal to all *Listeria* species. In addition to this fragment, strains from group 1 (1/2a−3a) generate one amplified fragment from gene *lmo0737* (unknown function) (691 bp), strains from group 2 (1/2c−3c) produce two fragments from gene *lmo0737* (691 bp) and gene *lmo1118* (unknown function) (906 bp), strains from group 3 (1/2b−3b−7) amplify a fragment from ORF2819 (putative transcriptional regulator) (471 bp), and strains from group 4 (4b−4d−4e) generate two fragments from ORF2819 (471 bp) and from ORF2110 (putative secreted protein) (597 bp). However, the proposed MPCR assay does not distinguish *L. monocytogenes* serovar 1/2a from 3a, 1/2c from 3c, 1/2b from 3b and 7, 4b from 4d and 4e. Isolates from serovars 4a and 4c (grouped as L in the present work) and isolates from the other *Listeria* species are also undistinguishable by this approach.

In order to better differentiate *L. monocytogenes* isolates, different subtyping methods have been used [10,22,23] although pulsed-field gel electrophoresis (PFGE) is the current gold standard method for molecular subtyping of most foodborne bacteria, including *L. monocytogenes* [8]. Results from different subtyping studies suggest that three genetic divisions or lineages correlated with the flagellar (H) antigen type exist in *L. monocytogenes* [2,15,19,23]. Lineage I consists of strains (flagellar antigen types b and d) that are more likely to cause human disease than isolates classified into lineages II (antigen type a or c) and III (serovars 4a and 4c and some 4b strains). However, clustering analysis based on these studies usually uses isolates from the more frequent serovars (1/2a, 1/2b, 1/2c and 4b).

Eighty isolates collected in Portugal and out of Portugal from different sources (including humans, animals, food and environment), and belonging to twelve serovars were analyzed in this study. We used MPCR serogrouping and PFGE subtyping to investigate (i) the typeability of the rare serovars, (ii) the ability of PFGE to differentiate serovars of *L. monocytogenes* within MPCR serogroups and (iii) the association of molecular types with specific sources or geographical origin (from Portugal versus out of Portugal). To our knowledge, this is the first report on clustering analysis of PFGE patterns of a group of isolates representing the twelve serovars of *L. monocytogenes*.

**Materials and methods**

**Bacterial strains and growth media**

Eighty *L. monocytogenes* isolates (Fig. 1) including 13 culture collection strains, previously classified by slide agglutination serotyping, and belonging to the twelve serovars were characterized in this study. The strains were from food (*n* = 29), from human cases of listeriosis (*n* = 25), from animals (*n* = 4), from environment (*n* = 5) and from uncertain sources (*n* = 17) (Fig. 1). The geographical origin of the isolates was ascribed as from Portugal (P) or from out of Portugal (O), as it was in most of the cases the only information we had. Some of the *L. monocytogenes* strains were kindly provided by the Faculté de Médecine de Tours, Université François-Rabelais (L’U.F.R), France, Laboratoire des Listeria, Institut Pasteur (I.P.) Paris, France, Universidade do Algarve (UA), Portugal and Università degli studi di Udine (USU), Italy. The 13 reference strains were from Collection de l’Institut Pasteur (CIP), Collección Española de Cultivos Tipo (CECT) and National Collection of Type Cultures (NCTC): CIP104794 (= NCTC7973) and NCTC7973 (serovar 1/2a); CECT936 and NCTC10887 (serovar 1/2b); CECT911 (= ATCC19113) (ATCC—American Type Culture Collection) and NCTC9862 (serovar 1/2c); CECT913 (serovar 3a); CECT937 and CIP78.35 (serovar 3b); CECT911 (= ATCC19113) (ATCC—American Type Culture Collection) and NCTC9862 (serovar 1/2c); NCTC5105 (serovar 4a); CECT937 and CIP78.35 (serovar 3b); CECT934 (= ATCC19114) (serovar 4a); CECT 4032 (NCTC11994) and NCTC10527 (serovar 4b); CIP78.39 (= ATCC19116) (serovar 4c).

All strains were kept on Tryptic Soy Broth (TSB, Difco, Detroit, USA) containing 15% glycerol and stored at −80 °C. Before the assays, each *L. monocytogenes* selected isolate was streaked on TSA-YE (Oxoid, Hampshire, UK) plates, and incubated for 18 h at 37 °C. Queries of the PathogenTracker database (http://www.pathogentracker.net) which contains subtype information for 648 *L. monocytogenes* (as of 19/06/2008) from various sources were conducted to probe the distribution of 22 out of 45 Portuguese isolates.

**Multiplex-PCR (MPCR) serogrouping**

MPCR serogrouping was performed using as PCR templates lysates obtained from bacterial colonies,
The five primer sets for target fragments from genes \textit{lmo0737}, \textit{lmo1118}, ORF2819, ORF2110 and \textit{prs} were synthesized by MWG-Biotech AG Switzerland, according to the sequence data of Doumith et al. [4]. The resulting PCR products were resolved on 1.4\% (w/v) agarose gels in 1 \times \text{TAE} buffer, at 2.3 V/cm for 60 min. The gels were stained with ethidium bromide (Sigma) and analysed by using Bio-Rad Gel Doc 2000\textsuperscript{TM} imaging system (Bio-Rad Laboratories, Segrate, Milan Italy).

### PFGE

The 80 \textit{L. monocytogenes} isolates were characterized by DNA macrorestriction analysis using PFGE in accordance with the Pulse-Net standardized protocol [9]. Bacterial cultures were embedded in chromosomal grade agarose (SeaKem Gold Agarose, Cambrex, New Jersey, USA), lysed, washed and the DNA was \textit{in situ} digested in separate reactions with 10 U of \textit{AciI} (New England Biolabs, Massachusetts, USA) at 37°C and with 20 U of \textit{ApaI} (Roche Diagnostics, Mannheim, Germany) at 30°C overnight, respectively. The resolution of the generated DNA fragments was obtained with 1\% SeaKem Gold Agarose gels in 0.5 \times \text{Tris-Borate EDTA} Buffer (Invitrogen, Paisley, UK) at 14°C and 6 V/cm with time ramped for 4-40 s over 22 h using a CHEF DR II System (BioRad, Hercules, USA). Gels were stained with ethidium bromide at a final concentration of 10 \mu g/ml (Sigma, St. Louis, USA) and pattern images were acquired in Bio-Rad Gel Doc 2000\textsuperscript{TM} imaging system.
imaging system (Bio-Rad Laboratories, Milan, Italy). The TIF images were normalized by aligning the peaks of the size standard strain Salmonella enterica serovar Braenderup H9812 which was loaded on three lanes in each gel. This strain was kindly provided by Centre National de Référence des Listeria, Institut Pasteur, France.

Data analysis

PFGE patterns were analyzed and compared using the GelCompar II version 4.5 (Applied Maths, Kortrijk, Belgium). The levels of similarity were based on the Dice correlation coefficient. For cluster analysis of the patterns the unweighted-pair group matching algorithm (UPGMA) was used with a tolerance of 1.50% and optimization of 1.50% as recommended by Martín et al. [13]. The quality of the cluster analysis was verified by calculating the cophenetic correlation value (r) for the dendrogram. Eight control strains of L. monocytogenes were used to determine the experimental variation between duplicate experiments. For both enzymes, the minimum level of repeatability of the macrorestriction conditions was calculated by running DNA samples from duplicated restrictions of the DNA of each strain. On the basis of the obtained results, a cut-off value of similarity was established for typing identical strains with identical outputs.

The Simpson’s index of discrimination (SID) of PFGE, expressed as a percentage, was determined as described by Hunter and Gaston [12].

Results and discussion

MPCR and PFGE analysis are currently used by several public and private laboratories for serogrouping and subtyping L. monocytogenes. The first method constitutes a practical alternative to slide agglutination serotyping and separates L. monocytogenes isolates belonging to the twelve serovars into five distinct serogroups. PFGE is considered the gold standard method for subtyping foodborne pathogens, because of its high discriminatory power and reproducibility. In this study, the combination of these methodologies with computer-assisted data analysis was applied to an isolate set representing considerable diversity (eighty isolates belonging to the twelve serovars representatives of the three genetic lineages of L. monocytogenes).

By using MPCR serogrouping, the 80 isolates were assigned to five groups (1–4 and L) (Fig. 1) in accordance with previous results from conventional serotyping. The exception was the strain EGDe (serovar 1/2a) that showed a profile corresponding to MPCR serogroup 2 (1/2c–3c) (Fig. 1) in agreement with a previous report of its unusual behaviour in this respect [5]. In accordance with this, the AscI and ApaI PFGE patterns of this strain were closer to the ones displayed by two culture collection strains from serovar 1/2c (0.79 similarity) than to the other patterns displayed by the serovar 1/2a isolates analysed (Fig. 1).

Strain 3980 given to us as serovar 3, displayed a multiplex pattern for group 3 (1/2b–3b–7), suggesting to be a serovar 3b isolate. This result was confirmed by the location of this isolate in the dendrogram (Fig. 1).

The results of multiplex-PCR of strains from the rare serovars 4a and 4c (group L) showed a negative influence of biomass excess in the cell lysate. In fact, in addition to the expected 370 bp fragment, a faint band of about 471 bp was present in the first amplification profiles of these strains (results not shown). This fact could lead into an incorrect assignment of these strains to multiplex group 3 (serovars 1/2b–3b–7). Nightingale et al. [17] also reported the misclassification of strains from serovars 4a and 4c in multiplex group 3. In the last trials using a lysate with less biomass, this faint band disappeared pointing out the need to work in stringent compliance with the protocol.

PFGE of the DNAs of the 80 isolates digested with AscI showed 7–16 fragments ranging from approximately 30 to 1130 kb in size, while 11–16 fragments of 30–510 kb were obtained following digestion using ApaI. The genomic DNA from two out of the three strains from serovar 4a (strain CECT 934 and isolate 3977) and serovar 4c (strain CIP78.39. ATCC19116 (serovar 4c) equivalent to our strain CIP78.39) showed 7–16 fragments ranging from approximately 30 to 1130 kb in size, while 11–16 fragments of 30–510 kb were obtained following digestion using ApaI. The genomic DNA from two out of the three strains from serovar 4a (strain CECT 934 and isolate 3977) and serovar 4c (strain CIP78.39. ATCC19116 (serovar 4c) equivalent to our strain CIP78.39).

The relationships among L. monocytogenes strains based on their combined ApaI and AscI PFGE profiles are shown in the dendrogram displayed in Fig. 1, and are supported by a cophenetic correlation coefficient (r) equal to 0.92. This value explained the goodness of fit of the clustering to the original data [21] and usually ranges between 0.60 and 0.95 and values higher than 0.80 are considered reasonable [18].

Based on data obtained from the duplicate experiments, a cut-off level of 0.95 similarity was established for the definition of a PFGE pattern. PFGE of AscI macrorestriction fragments differentiated the 80 isolates into 33 different patterns. The PFGE of ApaI macrorestriction fragments of the 77 typeable isolates produced 44 different patterns. A higher SID value was achieved with ApaI (0.91) compared to AscI (0.85). The combination of patterns produced by both enzymes yielded 62 combined ApaI and AscI PFGE types (pulsotypes) increasing the discriminatory power up to 0.97 (Fig. 1).

At about 0.42 similarity level L. monocytogenes isolates could be divided into two major clusters. The
smaller cluster corresponded to lineage III (three isolates from serovars 4a and 4c typed only by AseI) well separated from the bigger cluster that gathered at 0.50 of similarity two subgroups. In one of these subgroups the 19 isolates from serovars 1/2a, 1/2c, 3a and 3c (lineage II) were clustered. The other subgroup gathered the 58 isolates from serovars 1/2b, 3b, 4b, 4d, 4e and 7 (lineage I). These results are in agreement with Dounith et al. [4] that used a DNA macroarray-based subtyping method to divide the three lineages of L. monocytogenes into five phylogenetic groups, each correlated with serovars. In fact, clustering of the isolates according to their pulsotypes is in accordance with MPCR serogroups.

Nevertheless, PFGE subtyping could not differentiate serovars within MPCR serogroups. Moreover, within serogroups identical pulsotypes were shared by isolates belonging to different serovars: serogroup 3 (serovars 1/2b and 3b, isolates 3024 and 3026) and serogroup 4 (serovars 4e and 4d, isolates 4056, 4057 and 4062) (Fig. 1). Interestingly, two isolates belonging to MPCR serogroup 4 (serovars 4b-4d-4e) had significantly different levels of pathogenicity (evaluated both in vivo and in vitro) [16]; human isolate 3846 (4b) was significantly more virulent than food isolate 442 (4d/4e) although their pulsotypes had about 0.94% similarity.

For some isolates, similarity clustering analysis of PFGE patterns suggested association between molecular types and specific sources or Portuguese origin, in accordance with previous reports [3,7]. At 0.53 similarity level, 19 isolates from lineage II were clustered in two subgroups (Fig. 1). In one of these subgroups seven out of eight isolates were from Portugal and collected from food (chicken meat and milk). There was no evidence that the isolates from milk were related or not and there was no acknowledged association between the chicken meat and the milk. In the other subgroup, the eleven isolates were mainly from out of Portugal and from diverse sources.

At 0.73 similarity level (Fig. 1) the isolates from lineage I were grouped in three major subgroups. One of these subgroups gathered nine L. monocytogenes (serogroup 4) Portuguese isolates from milk, cheese, dairy environment and from human cases of listeriosis. Two of these isolates (3183 and 3972) collected in 2002 from raw milk, and in 2004 from placenta, respectively, have identical pulsotypes. In this cluster, there are isolates from milk and cheese (3077 and 3143) persistently collected in two different cheese dairies.

At the same similarity level, the two other major clusters with 18 and 14 isolates (serogroups 3 and 4, respectively) were diverse in what geographical origin (from Portugal and from out of Portugal) and source (type of food, humans, environment and animal) are concerned. The wide distribution of some of these pulsotypes was confirmed by querying the Pathogen-Tracker database for 22 selected isolates. Isolate 3845 collected in 2000 in Portugal from an infected person shared the same pulsotype (designated in Fugett et al. [6] by pulsotype 7) with 17 isolates in PT database: bovine (2), human (5), water (4), food (2), caprine (2), environment (2). Among this group were isolates involved in two outbreaks related to cheese consumption, in Los Angeles (LA) and in USA (1985) and Switzerland (1983–1987) [6]. Portuguese isolates 3857 and 3858 collected from two infected persons in 1997 and 1998, respectively, had a pulsotype indistinguishable from pulsotype 20 displayed by isolates involved in a human epidemic case related to pâté consumption in 1988–1990, in the UK [6].

Computer-assisted data analysis of PFGE data simplifies the processing of a large number of samples while allowing data sharing if a stringent compliance to standard protocols is achieved. This standardization includes not only a rigorous accomplishment of the Pulse-Net standardized protocol by Graves and Swaminathan [9] but also standardization of procedures in computer-assisted data analysis for pattern normalization and transfers of information intra and inter laboratories.

PulseNet Europe [13] recommends settings of both optimization and position tolerance at 1.5% for band comparison in accordance with Pulsnet USA. Although complying with these settings, strains visually indistinguishable may still be considered different according to the clustering analysis [11]. Aiming to minimize this, the determination of the experimental variation between duplicate experiments was evaluated in this work, allowing an establishment of a cut-off value of similarity of 0.95 for typing identical PFGE patterns. In spite of this, visual PFGE matches were found different by computer analysis (e.g. serovar 4b isolates 3102 (cheese), 3183 (milk) and 3972 (human)). Owing to this, a causal relationship between food and human isolates cannot be deduced without an additional subtyping method, in preference amenable to automation and based on DNA sequence. Nevertheless, to improve correct interpretation of subtyping data, the availability of a large and diverse PFGE type database is needed.

To our knowledge, this is the first report on clustering analysis of PFGE patterns of a diversity set of strains from twelve out of the thirteen serovars representative of all three genetically different L. monocytogenes lineages.

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References


