IMPLEMENTATION OF CLASSICAL, MOLECULAR BIOLOGICAL AND IMMUNOENZYMATIC METHODS IN ISOLATION AND DETECTION OF *Listeria monocytogenes* IN FOOD

Brankica Z. Lakićević¹, Aleksandra B. Martinović², Tatjana M. Baltić¹, Slobodan V. Lilić¹, Branka R. Borović¹, Predrag M. Ikonić³, Vesna V. Janković¹

¹Institute of Meat Hygiene and Technology, Kaćanskog 13, Belgrade, Serbia
²Faculty of Food Technology, Food Safety and Ecology, Donja Gorica, 81 000 Podgorica
³University of Novi Sad, Institute of Food Technology, Bulevar cara Lazara 1, 21000 Novi Sad, Serbia

*Corresponding author:
Phone: +381112650655
Fax: +381112651825
E-mail address: brankica@inmesbgd.com

ABSTRACT: Food borne pathogens and spoilage bacteria are influencing the safety and quality of food and animal feed and can result in serious adverse effects to human and animal health as well as to the food quality. Consequently, microbiological quality control in the food industry has become the priority of the food producers and it aims towards minimizing the risks related to food pathogens and spoilage bacteria. One of the most important pathogens in the food industry is *Listeria monocytogenes*. Listeriosis has a significant public health and economic impact because of its high hospitalization and mortality rate. Most people infected with *Listeria* are hospitalized and mortality is approximately 30%. Therefore it is necessary to undertake efficient control measures, especially concerning the necessity of a rapid and accurate detection of this pathogen in the food industry as well as in retail food samples. Recently used conventional methods are often time consuming and require intensive work. However, obtained results can often be false considering the presence of viable but not cultivable microorganisms. Advances in biotechnology and bioinformatics have resulted in the development of novel testing technologies that enable tracking, more reliable and faster detection of food pathogens. Furthermore, molecular-biology methods, although still not applied routinely in everyday practice, are the promising alternative which can replace current reference methods in this area.

Key words: *Listeria monocytogenes*, laboratory detection methods, rapid tests.

INTRODUCTION

*Listeria monocytogenes*, a facultative intracellular Gram positive bacterium, can cause serious illness in humans and animals. It is associated with abortion, meningoencephalitis, and septicaemia in humans, especially in individuals at risk including pregnant women, newborn babies, the elderly and the immunocompromised (McLauchlin, 1997). In recent years, several studies reported outbreaks of listeriosis caused by *L. monocytogenes* associated with consumption of different foods, particularly ready-to-eat products like soft cheese (Bannister, 1987), cooked and chilled poultry products (Barners et al., 1989), deli meat products (Jacquet et al., 1995) or hot dogs. Also, some epidemiologically monitored outbreaks have been reported (Kathariou, 2000). The increasing incidence of *L. monocytogenes* in food-borne outbreaks has led to the urgent
need for a rapid detection system for testing food samples. Conventional microbiological methods for routine detection of *L. monocytogenes* include culture enrichment in liquid medium, (usually Fraser broth), plating on selective agar medium, biotyping and even serotyping. This procedure takes from 5 to 7 days. In the last 10 years, several PCR detection methods have been proposed to replace the time-consuming culture based conventional techniques. The most frequently chosen target gene is the *hlyA*, encoding listeriolysin O, and other targeted genes include the *iap* (invasion-associated protein), the *dth-18* (delayed type hypersensitivity), the 16S rRNA, an aminopeptidase coding gene and *inlA* (internalin operon) (Aznar and Alarcon, 2003).

**Conventional microbiological methods**

Conventional microbiological methods used for detecting microorganisms in food are easily adaptable, simple, practical, generally inexpensive and they can be used in quantitative and qualitative research. However, there are some limitations using conventional microbiological methods, especially when applied to the detection of pathogens. These methods are based on the growth of the target microorganisms in one or more culture media, visual detection of microbial growth and confirmation of the presence of the pathogen which usually included combination of biochemical and serological tests. The main disadvantages of conventional microbiological methods are: extremely challenging phases of analysis, subjective interpretation of results and for some pathogens long time required to carry out the analyses.

The most commonly used reference methods for the detection of *Listeria* spp. in food are ISO 11290 standards (ISO, 1996; EC, 1999). The U.S. Food and Drug Administration (FDA) has developed a protocol for the isolation of *Listeria* spp. from dairy products, seafood and vegetables (Hitchins, 2003), and the U.S. Department of Agriculture (USDA) developed a method for the isolation of these bacteria from meat and poultry products as well as from the environmental samples (USDA, 2002). Conventional method remains the “gold standard” with which other method is compared. They are usually very sensitive. These methods use selective agents and enrichment procedure to reduce the number of contaminating microorganisms and allow multiplication of *L. monocytogenes*.

All above mentioned protocols included homogenization and incubation of pre-enrichment and enrichment culture for 24 - 72 h at 30 - 37 °C. Furthermore, before the 1990s, all commercially available media for the isolation of *L. monocytogenes* had low selectivity with the lack of the ability to distinguish *L. monocytogenes* from non-pathogenic species. Therefore, there is a need for the development of new and more selective media. Chromogenic media is the most used culture confirmation method because of its ease preparation and interpretation. Using chromogenic agars (ALOA, Rapid‘L. mono), the presumptive identification of *L. monocytogenes* is possible after 24 h, compared with 3-4 days using Oxford and other conventional agars (Greenwood et al., 2005).

**Immunoenzymatic methods**

There are various methods based on antibodies specific to *Listeria* available as commercial kits that have been applied in food testing for many years. However, only a few are available for the specific detection of *L. monocytogenes*. The enzyme linked immunosorbent assay is the most common antibody assay format used for pathogen detection in foods.

The enzyme immunoassays may contribute to accelerate and simplify the detection of various types of pathogens in foods and are the most commonly used because they possess several advantages: simplicity, rapidity, sensitivity, specificity and convenience as a screening method (Ky et al., 2004). ELISA methods that use an antibody immobilized to a microtitre well for antigen capture in combination with a secondary antibody coupled to an enzyme (or another
label) to detect the capture antigen, are the most widely applied methods because they combine ease of use with the generation of rapid test results. It is easily applied, generates rapid test results and allows the use of difficult sample matrix. The success of ELISA method for detection of *L. monocytogenes* depends on the specificity of the antibody. Using hybridoma technology, it became possible to develop monoclonal antibodies that react only with *L. monocytogenes* (Janzten et al., 2006). There is a test which uses monoclonal antibodies that recognize the protein p60 (invasion–associated protein) encoded by the *iap* gene for identification of *L. monocytogenes* (Ky et al., 2004). By combining monoclonal antibodies recognized by this protein, it was possible to develop an ELISA system which can specifically identify *L. monocytogenes* or simply detect *Listeria* species. The Transia Plate *Listeria monocytogenes* (Diffchamb AB) method, which combine a two-step enrichment protocol and a specific *Listeria monocytogenes* immunoassay present specific, effective and reliable method for *Listeria monocytogenes* detection in food. The enrichment protocol was applicable to various food matrices, and microplate format of ELISA could be manipulated manually, or automatically. These method characteristics allow implementation in any food analysis laboratory. Another commercially available test is RIDASCREEN Listeria, sandwich enzyme immune assay for the qualitative detection of *Listeria* spp. from pre–enrichment cultures of food samples and swab samples from the production environment.

Automated Mini-VIDAS system, (BioMérieux, France), based on ELFA technology (The enzyme linked with fluorescent assay) combines ELISA and fluorescence detection. The VIDAS® LMO II was validated in 2002 (USDA, 2002) as a rapid method for all food products and environmental samples. It has been successfully used for the detection of stable antigens of virulent *L. monocytogenes*. In addition, there are other commercially available mini VIDAS tests such as VIDAS Listeria DUO for simultaneous detection of *L. monocytogenes* and other bacteria of the genus *Listeria* (Janzten et al., 2006) and LMX for the qualitative detection of *L. monocytogenes*.

A low quantity of culture is divided by monocal pipette, regulated to 500 µL, in the strip for reactive, the samples are absorbed and disposed from cone with a specified time period.

Figure 1. Mini-VIDAS®
Listeria antigens that are present in the sample will interlock with anti-Listeria antibodies, which are disposed on cone walls. Unattached components of the cone are removed during washing steps. Monoclonal antibodies marked with alkaline phosphatase (conjugate) are absorbed and eliminated from the cone, coupled with any captured antigen of Listeria on cone walls.

A final washing step removes the unbound conjugate. During the final stage of detection, the substrate, 4-methyl-phosphate umbelliferol is vacuumed and removed from the cone. Enzymatic conjugate has a catalytic action on hydrolysis of this substrate with a fluorescent product (4-methyl-umbelliferone) whose fluorescence is measured at 450 nm. When the test has been finished, the results are analyzed automatically by software. Resulting values are compared with a set of references and each sample could be positive or negative (BioMérieux, France).

The advantages of this method are: automatic printing of results at the end of the analysis, reduced time for enrichment step and detection, standardization of work (reproducibility and reliability), flexibility in operation, high sensitivity and specificity.

Molecular biological methods

Molecular biological methods have become an important support to conventional microbiological methods. Characteristic of molecular methods is rapidity, repeatability and reliability. In addition, molecular methods are independent of the variable conditions of the growth of microorganisms. Methods based on DNA analysis are simple, reliable and more cost-effective methods for identification and classification of microorganisms.

Polymerase Chain Reaction (PCR) is a method based on the selective amplification of specific fragments of DNA. The first studies of sensitivity of PCR were carried out on pure cultures (Border et al., 1990; Wang et al., 1992) and artificially contaminated food samples (Furrer et al., 1991; Wang et al., 1992). Validation with naturally contaminated samples has also been performed in a wide range of foods (Rossen et al., 1991; Bohnert et al., 1992; Niederhauser et al., 1992, 1993). Results of these studies are not comparable with the results obtained by different procedures, in different food matrices and PCR conditions.

The most frequently chosen target gene is the hlyA, encoding listeriolysin O, (Furrer et al., 1991; Bohnert et al., 1992; Niederhauser et al., 1992; Ericsson and Stålhandske, 1997; Lakicevic et al., 2010), and other target genes include the iap (invasion-associated protein) (Köhler et al., 1990; Niederhauser et al., 1992; Wang and Hong, 1999; Lakicevic et al., 2010), the dth-18 (delayed type hypersensitivity) (Wernars et al., 1991), the 16S rRNA (Wang et al., 1992) while int/A (internalin operon) (Almeida and Almeida, 2000).

Complex sample preparation and detection of PCR products using agarose gels in the final step (when a large part of the resulting PCR products are affected by degradation processes) are marked as the main disadvantages of conventional PCR. In addition, conventional PCR methods only detect the presence of pathogens (Churchill et al., 2006) and are not able to make a difference between live and dead bacterial cells (O’Connor, 2003).

Multiplex PCR is a variation of the traditional PCR. This method uses multiple sets of primers to amplify a number of genes or gene fragments simultaneously. For instance, Bhagwat (2003) artificially contaminated fresh produce and used multiplex PCR to simultaneously detect L. monocytogenes, S. typhimurium, and E. coli O157:H7. Although identification procedure is effective, analysis the PCR products by agarose gel electrophoresis does not make this method suitable to high-throughput screening. Real-Time PCR detection is preceded by a single-step or a two-step enrichment using media of different selectivity.

These methods use different enrichment and DNA preparation approaches to reach the goal of increasing the target live cell numbers to a detectable level and dilution of dead L. monocytogenes.
Real Time PCR method detects the accumulation of amplicon during the PCR reaction. The data is then measured at the exponential phase of the reaction when the amplification DNA process is going fast, and the reaction in this phase is highly specific and precise. The principle of the method is based on detection and quantification of reporter fluorescent dye, whose emission is directly proportional to the amount of PCR product. Samples with an increasing fluorescence signal were considered positive, regardless of the internal control amplification. Samples with no increasing fluorescence signal but with amplified internal control were considered negative. Samples with no fluorescence signal for both specific target and internal control were considered inhibited. Advantages of Real Time PCR methods are analytical and diagnostic accuracy, high sensitivity, quick performance analysis, universal sample preparation for analysis, the possibility of quantitative analysis, and many others (Braguta, 2008).

Guilbaud et al. (2005) have developed a quantitative Real Time PCR method for determination of *L. monocytogenes* in an artificially made biofilms using a non-specific SYBR Green dye. Their results showed that the number of *L. monocytogenes* cells, which can be detected in biofilms is $6 \times 10^2$ CFU/cm$^2$. Rantsiou et al. (2008) developed quantitative real time PCR method for the detection, quantification and determination of the viability of *L. monocytogenes* in milk, meat, soft cheese, fermented sausages, ham and salad. Limit of quantification of method is $10^3$-$10^4$ cfu / g or ml, while after overnight incubation at 37 ° C it was possible to detect a 10 cfu / ml or g. In addition, in the case of the mixed culture, if *L. innocua* is present at level that is 2.5 to 3 times higher than *L. monocytogenes*, the identification of blue green colonies with halo, characteristic of *L. monocytogenes*, was not possible.

Denaturing Gradient Gel Electrophoresis (DGGE) is an electrophoretic method which can detect differences between the DNA fragments of the same size but different nucleotide sequence, based on the different melting points of DNA. This is achieved in a polyacrylamide gel in the presence of denaturing agents urea and formamide. Cocolin et al. (2002) described the direct identification of *Listeria* spp. and *L. monocytogenes* in food samples combining PCR and DGGE. The method is based on the PCR amplification of a fragment of the *iap* gene from the five species belonging to the
genus and on the analysis of the PCR products obtained by denaturing gradient gel electrophoresis. Due to differences in amplified gene sequences, it is possible quickly and easy to distinguish five species of the genus *Listeria* as well as different serotypes within the *L. monocytogenes* species. This method can be used as a quick test to determine the presence of *Listeria* spp. and *L. monocytogenes* in food (Cocolin et al., 2002). In order to interpret data with greater accuracy, electrophoretic techniques were developed to allow better resolution of DNA fragments, such as pulse-field gel electrophoresis (PFGE), which is primarily used in conjunction with restriction enzyme (endonuclease) digests of DNA.

Pulse-Field Gel Electrophoresis (PFGE) enables separation of large DNA fragments in pulsed electric field. Total DNA of analyzed microorganism is cut with restriction enzymes, in the sites which are relatively rare in the genome. It is possible to get from 5 to 50 restriction fragments. This method is highly discriminative with high repeatability and PFGE is considered as method of choice for the molecular typing of *L. monocytogenes*. In 1996, Brosch et al. indicated that PFGE allows typing of *L. monocytogenes* serotype 4b. Furthermore, PFGE has been widely used in industrial plants to investigate the pathways of contamination by bacteria *L. monocytogenes* (Ojeniyi et al., 1996; Unnerstad et al., 1996; Auto et al., 1999; Giovannacci et al. 1999; Miettinen et al., 1999, 2001; Senczek et al., 2000; Fonnesbech Vogel et al., 2001; Berrang et al., 2002).

Multilocus Enzyme Electrophoresis (MEE), based on electrophoretic mobilities of soluble metabolic enzymes, has been applied in a number of epidemiological studies (Farber et al., 1991; Rørvik et al., 1995, 2000; Avery et al., 1996; Nesbakken et al., 1996; Buncic et al., 2001). This method has relatively low discriminatory power (Caugant et al., 1996).

Restriction Endonuclease Analysis (REA) is based on recognizing and cutting the total DNA of the microorganism at specific recognition sequences in the chromosome. As a consequence, restriction endonuclease generated fragments of different lengths. Large number of fragments, which make difficult interpretation of results is marked as the main disadvantage of this method (Gerner Smidt et al., 1996).

Ribotyping is an RNA – based molecular characterization of bacteria. In ribotyping, restriction enzymes (i.e., enzymes which cut DNA at specific positions) are used to cut the genes coding for rRNA into pieces, and gel electrophoresis is used to separate the pieces by size. Genetic probes then visualize locations of different – size fragments of DNA in the gel, which appear as bands. Low power separation of strains can be overcame by using different enzymes (de Cesare et al., 2001). Automated ribotyping was used in a number of epidemiological studies (Tkáčiková et al., 2000; Norton et al., 2001; Berrang et al., 2002; Suikko et al., 2002; Aarnisalo et al., 2003; Hoffman et al., 2003), when has been successfully identified a large number of isolates.

Random Amplification of Polymorphic DNA (RAPD) is a method which uses a short arbitrary primers (Williams et al., 1990) for copying DNA fragments by PCR method in unrestricted conditions. This method is reliable and easy to perform, but the results are not always reproducible (Wernars et al., 1996). RAPD analysis allows discrimination among isolates of the same serotype and also among isolates from a common source though identical profiles were obtained for some isolates belonging to the same serovar (Fonnesbech Vogel et al., 2001).

RAPD has a high power separation of *L. monocytogenes* isolates (Boerlin et al., 1995) and has been used in several epidemiological studies (Lawrence and Gil-mour 1995; Giovannacci et al., 1999; Agudo et al., 2001; Fonnesbech Vogel et al., 2001).

Amplified Fragment Length Polymorphism (AFLP) is a PCR-based fingerprinting method that was originally developed for typing of plants. The AFLP technique involves
three steps: 1) genomic DNA is cleaved with two restriction enzymes, 2) ligation of restriction site-specific adapters occurs and 3) a subset of fragments is amplified by PCR. This method is a “fingerprint” tech-nique that does not require sequence infor-mation of the studied organisms, similarly as RAPD. Reproducibility is higher than RAPD due to stricter conditions for PCR reaction. The initial step involves the digestion of the whole genome by restriction enzymes, followed by ligation and PCR reaction. The fragments obtained by AFLP, are separated on denaturing gel and visualization of the fragments can be fluorescent or autoradiography. AFLP has a high power to distinguish *L. monocytogenes* isolates similarly as PFGE (Fonnesbech Vogel et al., 2001; Keto Timonen et al., 2003). Genotyping which includes macro-and microarray tech-niques are not yet used in routine diagnosis, but initial results indicate the great impor-tance of these methods (Ryser and Marth, 1999).

### Phenotypic methods

Phenotypic methods include methods of serotyping, phage and bacteriocin typing. Serotyping is based on antigenic differences of *L. monocytogenes* strains (Seeliger and Hohne, 1979). In relation to the somatic and flagellar antigens within the *L. monocytogenes* species, may be 13 different sero-types (Schoenberg et al. 1996) of which only serotypes 1/2a, 1/2b and 4b were isolated from clinical samples and food. Phage typing involves susceptibility *L. monocytogenes* isolates to the specific bacteriophage (Rocourt et al., 1985). Important character-istics of phage typing are greater discrimi-nation power and the identification of num-ber isolates for a relatively short period of time. A disadvantage of this method is that not all *L. monocytogenes* strains can be subjected to typing (McLauchlin et al. 1996; Ojeniyi et al., 1996). For the time being, this method has a low diagnostic value (Radoji-čić et al. 2011).

Bacteriocins typing involves susceptibility *L. monocytogenes* isolates to the different bac-teriocins. This method is rarely used in epi-demiological studies due to poor power to distinguish isolates of *L. monocytogenes* (Curtis and Mitchell, 1992).

### CONCLUSIONS

The current reference methods for the detection of *L. monocytogenes* allow the re-covery of this pathogen from a variety of foods with relative ease. The introduction of chromogenic media efficiently improved the isolation of *L. monocytogenes*. Food producers and distributors have great interest in more rapid methods, which can help to bring about the desired changes in the available technology. Molecular biology, DNA based methods, particularly PCR, can be gradually replaced by the conventional methods for the detection of *L. monocytogenes*. How-ever, there are still many problems to be solved, such as the sample preparation, elimination of the effects caused by non-specific binding and cross hybridization and even greater sensitivity of the overall sys-tem.

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ПРИМЕНА КЛАСИЧНИХ, МОЛЕКУЛАНРНО БИОЛОШКИХ И ИМУНОЭНЗИМСКИХ МЕТОДА У ИЗОЛАЦИЈИ И ДЕТЕКЦИЈИ БАКТЕРИЈЕ Listeria monocytogenes

Бранкица З. Лакићевић1*, Александра Мартиновић2, Татјана М. Балтић1, Слободан В. Лилић1, Бранка Р. Боровић1, Предраг М. Иконић3, Весна В. Јанковић1

1 Институт за хигиену и технологију меса, Каћанског 13, Београд, Србија
2 Факултет за прехрамбене технологије, безбедност хране и екологију, Доња Горица, 81 000 Подгорица
3 Универзитет Нови Сад, Научни институт за прехрамбене технологије у Новом Саду, Булевар цара Лазара бр. 1, 21000 Нови Сад, Србија

Садржај: Патогени хране и бактерије квара утичу на безбедност и квалитет хране и хране за животиње, јер доводе до озбиљних, нежељених ефеката по здравље људи и животиња. Сходно томе, примена микробиолошке контроле у прехрамбеној индустрији је постала приоритет, која има за циљ минимизирање ризика од инфекције патогенима хране и бактеријама квара. Један од најважнијих патогена у прехрамбеној индустрији је Listeria monocytogenes. Листериоза, чији је узрок Listeria monocytogenes, има здравствени и економски значај код људи и животиња, због високе стопе хоспитализације и смртности. Већина људи инфицираних овом бактеријом се хоспитализује, док приближно трећина инфицираних умире. Због тога је неопходно да се предузму ефикасне мере контроле, посебно када је реч о неопходности брзог и прецизног откривања овог патогена у прехрамбеној индустрији.

Конвенционалне микробиолошке методе, захтевају интензиван рад, дуг временски период, а ипак, добијени резултати не морају бити увек поуздани (случај вијабилних микроорганизама који не подлежу култивацији). Напредак у биотехнологији и биоинформатици довели су до развоја нових технологија које омогућавају тестирање, праћење, поузданије и брже откривање патогена хране. Осим тога, молекуларно-биолошке методе, иако се до сада још не примењују у рутинској прaksi, представљају обећавајућу альтернативу и могу заменити или употпуни тренутне референтне методе у овој области.

Кључне речи: Listeria monocytogenes, лабораторијске детекционе методе, брзи тестови

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