AN OPTIMISED QUANTITATIVE REAL-TIME PCR ASSAY FOR LISTERIA MONOCYTGENES INCLUDING AND INTERNAL AMPLIFICATION CONTROL

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ABSTRACT
The assessment of results obtained by PCR is a critical issue for its implementation as a routine tool in food microbiology diagnostics. As an analytical technique, PCR is exposed to inhibitors that can produce false negative results or underestimation in quantitative analysis. In this study, we illustrate the design, development and optimization of a duplex real-time PCR (RTi-PCR) assay for the quantitative detection of Listeria monocytogenes based on the co-amplification of a L. monocytogenes-specific gene (hly) sequence (Rodríguez-Lázaro et al., 2004a) and an internal amplification control (IAC) to evaluate PCR performance. A PCR IAC is a non-target DNA that is co-amplified with the same set of primers that target sequence. The assay was 100% specific as it allowed unambiguously detection of 49 L. monocytogenes isolates of different serotypes and sources and 104 strains of other species were negative. The detection and quantification limits were 8 and 30 cfu respectively, and PCR linearity and efficiency were R^2=0.997 and 0.80 respectively. Interestingly, the assay was capable of detecting underestimation of L. monocytogenes in reactions containing inhibitory culture media (e.g. Fraser and Half Fraser media) and foodstuffs (e.g. raw salmon or raw pork meat). Specific L. monocytogenes detection could also be performed using agar-grown colonies and by conventional PCR. In conclusion, we present a simple and rapid duplex RTi-PCR assay for the quantitative detection of L. monocytogenes, which can also assess potential underestimations or false negative results.

KEYWORDS
Quantitative detection; Real-time PCR; Internal Amplification Control; Listeria monocytogenes.

INTRODUCTION
Listeria monocytogenes is a human pathogen that can produce a severe disease often associated with the ingestion of contaminated ready-to-eat products (Kathariou, 2000; 2002; Ryser, 1999; Vázquez-Boland et al., 2001). Consequently, the application of microbiological control within the quality assessment programs in the food industry is a premise to minimize the risk of infection with this pathogen for the consumer. In this scenario, the current methodological approach is based on traditional microbiological methods, which are labour-intensive and time consuming and produce presumptive results that require additional biochemical tests for the definitive confirmation of this pathogen (Donnelly, 1999). These
drawbacks are overcome by real-time PCR (RTi-PCR), a technology that may allow the rapid and quantitative detection of infectious agents with extremely high specificity and high sensitivity (Klein, 2002; Mackay, 2004; Rijpens and Herman, 2002; Walker, 2002). Furthermore, the absence of post-PCR analytical steps reduces cross-contamination risks and allows high throughput and automation (Klein, 2002; Mackay, 2004). However, many components of food products, culture media and nucleic acids extraction reagents can act as PCR inhibitors (Rådström et al., 2003; Rossen et al., 1992; Wilson, 1997), and their presence in the reaction may cause a dramatic decrease in sensitivity or even block amplification reaction, consequently generating false negative results or an underestimation of the bacterial load (Hill, 1996; Scheu et al., 1998; Rådström et al., 2003). Therefore, the assessment of the PCR efficiency in every assay is a critical issue for the implementation of PCR as a routinely analytical tool in food microbiology diagnostics. An important analytical control for use with molecular amplification-based methods is the inclusion, in each reaction, of an internal amplification control (IAC) (Hoorfar et al, 2003; Rodríguez Lázaro et al., 2004b). A PCR IAC is a non-target DNA which is co-amplified simultaneously with the target sequence (Ballagi–Pordány and Beláé, 1996; Cone et al., 1992; ), thus, with negative results for the PCR target, the absence of a positive IAC signal indicates that amplification has failed (Hoorfar et al., 2004). A standard IAC for RTi-PCR assays should include the sequences of the primers for the target sequence flanking an internal region totally unrelated where a fluorescence probe for the IAC detection should be directed. Thus, The simultaneous use in a single reaction of two differently labelled fluorescent probes (for the target and IAC) allows both the detection of the target and assessment of the PCR efficiency in every reaction. Although a number of RTi-PCR assays have been developed for food-borne pathogens detection, adequate IAC have rarely been optimized and never used with quantitative purposes, even when the current consensus among the scientific community is that an IAC should be mandatory for PCR-based diagnostic tests (Hoorfar et al., 2003). Here, we report the development and optimization of a duplex RTi-PCR assay for the specific quantitative detection of L. monocytogenes, which includes an IAC designed to evaluate the PCR performance of every reaction. This assay combines the L. monocytogenes-specific hly-based assay and the amplification of a chimeric IAC DNA. We discuss technical aspects that were critical for optimization of the duplex reaction and provide data of experiments to determine the inhibitory effect of some food products and different culture broths for L. monocytogenes quantitative detection by RTi-PCR.

**MATERIAL AND METHODS**

**Bacterial strains, grown conditions, DNA isolation and quantification.** In this study, the stains (100 Listeria strains -49 L. monocytogenes, 17 L. innocua, 7 L. grayi, 10 L. seeligeri, 5 L. welshimeri and 12 L. ivanovii isolates- and 45 non-Listeria species), growth conditions, DNA isolation and quantification reported in a previous study (Rodríguez–Lázaro et al., 2004a) were used.

**IAC preparation.** The IAC DNA consisted of a 104 bp fragment of the acetyl-coenzyme A carboxylase gene (GenBank acc. no. X77576) from rapeseed (Brassica napus) flanked by hlyQF/R primers (Rodríguez–Lázaro et al., 2004a). This chimerical IAC DNA was constructed by a two-step PCR as previously described (Figure 1). Briefly, 100 ng of Brassica napus DNA were amplified using primers hlyccAF (5’ catggcaccaaccagcatctggagtgtgatataatc 3’) and hlyccAR (5’ atcgcgtttttcttgagcgctc 3’)

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using conditions previously described (Hoorfar et al., 2000). The amplicon was subsequently diluted 1:1,000 in double-distilled water and used as template in a second PCR reaction performed with the hly-specific primers hlyQF/R in the same conditions. The PCR product was purified using the QIAEX®II Gel Extraction kit (Qiagen) according to the recommendations of the manufacturer. The eluate was quantified and diluted in double-distilled water in the presence of 5 ng/µl tRNA down to the working concentration.

**Figure 1:**

**A**

Non-target genomic DNA (*Brassica napus*)

hly-accF

hlyQF

acc1

1st PCR

hlyQF

hly-accR

2nd PCR

**B**

Chimeric IAC DNA

hlyQF

IAC

hlyQP

hlyQF

hlyQP

hlyQF

L. monocytogenes DNA

hlyQF

hlyQP

hlyQR

**Fig. 1.** Schematic representation of the construction of the IAC (A) and of the templates and hybridizing sites for the fluorescent probes in the duplex hly-IAC RTi-PCR (B). In (A), acc1 and acc2 refer to the 3’ *B. napus* BnACCg8 gene-specific primer sequences. **RTi-PCR.** Reactions were performed using TaqMan® PCR core reagents kit (Applied Biosystems-Roche) in a 20 µl reaction volume containing 1× PCR TaqMan® buffer A (including ROX as a passive reference dye), 6 mM of MgCl2, 200 µM of each dATP, dCTP, dGTP, 400 of µM dUTP, 50 nM of hlyQF/R primers, 100 nM of hlyQP probe (FAM and TAMRA double labelled hly-specific probe) [(Rodríguez-Lázaro et al., 2004a) and 100 nM of accp (VIC and and TAMRA double labelled IAC-specific probe) (Hernández et al., 2001), 1 unit of AmpliTaq Gold® DNA polymerase, 0.2 units of AmpErase® uracil N-glycosylase (UNG), 100 molecules of IAC DNA (unless specifically stated) and 1 µl of the target DNA solution. Reactions were run and analysed as described in Rodriguez-Lázaro et al. (2004a). Conventional (c-)PCR were performed in the same conditions as RTi-PCR.
except for PCR buffer II. PCR products were detected in analytical 3% agarose gels stained with ethidium bromide.

**RESULTS AND DISCUSSION**

**Design and construction of the IAC and Optimization of the RTi-PCR assay.** The IAC was constructed following the indications shown in the materials and methods. It did not exhibit significant homology to any DNA sequence deposited in DNA databases (different to BnACCg8 positions 9651-9755) when analyzed using the bioinformatics tool BLAST-N. It should be amplified using the hly-specific primers (Rodríguez-Lázaro et al., 2004a) as shows their sequence at 5’- and 3’- ends, respectively, whereas it should not be detected by the hly-specific probe (FAM-labelled) but with the specific-IAC probe (Hernández et al., 2001) (VIC-labelled). This should facilitate simultaneous detection of *L. monocytogenes* and IAC in a single reaction by means of each of specific fluorescent probes. In addition, the IAC length is 143-bp, 79-bp longer than the hly-specific amplicon (64-bp), and thus it is also possible to distinguish both amplicon by gel-electrophoresis using c-PCR. In conclusion, we constructed an IAC DNA that did nor reveal homology to any other DNA sequence found in nature (to avoid any cross-reaction) and longer than the hly-specific amplicon (suitable for gel-electrophoresis analysis), and thus fulfilling the sequence and the size recommendations previously described (Ballagi-Pordány and Belák, 1996; Hoorfar et al., 2004).

Subsequently, we optimised the IAC (accep) probe concentration using decreasing accep probe concentrations (from 250 to 25 nM) in the conditions previously described for the hly-specific assay (Rodriguez-Lázaro, 2004a) with 1,000 IAC molecules and in the absence of *L. monocytogenes* DNA, and selecting the minimum probe concentration that did not result in a C_T increment (i.e. high sensitivity) (Applied Biosystems, 1998; Rodriguez-Lázaro et al., 2004c). Optimal conditions are those described in Materials and Methods. The determination of the optimal concentration of IAC in the PCR assay is a critical step. The IAC concentration should be should be kept as low as possible to avoid negative effect or even inhibition of the target-specific reaction. However, the IAC concentration should not be so low as to make it difficult to obtain reproducible IAC amplification signal. Thus, we determined the detection limit of the IAC using decreasing concentrations of IAC (1,000, 300, 100, 30 and 10 molecules per reaction) in the absence of *L. monocytogenes* DNA. The IAC could be consistently detected to 10 molecules, but VIC C_T values varied among replicates. In parallel, we also evaluated the IAC inhibition capacity, using reactions that contained increasing amounts of IAC (10, 30, 100, 300, 1,000, and 10,000) and approximately 30 *L. monocytogenes* cells. We observed that 100 IAC molecules was the highest amount of IAC that did not inhibit the *L. monocytogenes* signal. Considering these results and that we reproducibly detected always 100 IAC molecules by c-PCR (not shown), we established this amount as the IAC concentration per reaction. This number is within the range recommended by Ballagi-Pordány and Belák (50-500 copies) (Ballagi-Pordány and Belák, 1996), and is similar to those used in some previously PCR methods, e.g. 300 (Malorny et al., 2003) or 1,000 (Lambertz et al., 2000) and smaller than those described in other real-time PCR detection assays (Hoorfar et al., 2000; Klerks et al., 2004).

**Specificity and sensitivity of the duplex RTi-PCR assay.** The specificity of the assay was evaluated using 1 ng of genomic DNA purified from 49 *L. monocytogenes* strains (including representative strains of the different serovars) and 96 non-target bacteria (51 *Listeria* strains -17 *L. innocua*, 7 *L. grayi*, 10 *L. seeligeri*, 5 *L. welshimeri* and 12 *L. ivanovii-* and 45 non-
Listeria strains) (see table 1 and 2 of Rodríguez-Lázaro et al., 2004a). In addition, a colony of each strain directly transferred from an agar plate was also tested. The duplex RTi-PCR assay unequivocally distinguished L. monocytogenes isolates, accordingly with previously reported data for hly-specific RTi-PCR assays targeting the same (Rodríguez-Lázaro et al., 2004a) or neighbouring sequences (Nogva et al., 2000; Hough et al., 2002). In addition, all duplex reactions showed a positive (VIC) IAC signal, indicating thus that the negative (FAM) hly-signal obtained from non-L. monocytogenes isolates was not caused by RTi-PCR inhibition.

The L. monocytogenes-specific signal should not be inhibited by the simultaneous co-amplification of IAC, especially when small amounts of target molecules are expected. This is of particular relevance for L. monocytogenes, which is often present in low numbers (Ryser et al., 1999). We determined the sensitivity of the duplex RTi-PCR assay in the presence of 100 molecules of IAC using decreasing amounts of genomic DNA from L. monocytogenes UdG 1010 (equivalent to approximately 30, 15, 8, 4 and 1 cells). Table 1 shows FAM (hly) and VIC (IAC) C_{T} and ΔR_{n} values obtained in a total of 9 replicates performed in 3 independent experiments. The duplex RTi-PCR assay consistently detect down to 8 L. monocytogenes DNA molecules and detected at least in a 44.44% of the replicates corresponding to 4 and 1 target molecules. These results were the same that as the hly-specific assay (Rodriguez-Lázaro et al., 2004a), and similar to other hly-specific RTi-PCR assays (Hough 20002; Nogva et al., 2000). As expected, co-amplification of IAC was positive in all reactions, with overall C_{T} values of 33.59 ± 0.68 and ΔR_{n} values of 0.66 ± 0.11. The sensitivity of the duplex c-PCR assay was evaluated using the same experimental design. We consistently detected down to 15 target molecules, i.e. the sensitivity was slightly poorer in c-PCR than in RTi-PCR assays (data not shown). In conclusion, addition of 100 initial IAC molecules did not result in any detectable decrease in the sensitivity of the assay.

Table 1. Detection and quantification limits of the duplex RTi-PCR assay. 100 IAC molecules were included in each reaction. Lower and upper limit confidence intervals were calculated for the expected template molecules in each dilution at 95% confidence level. C_{T} is the cycle number at which the fluorescence intensity equals a fixed threshold and ΔR_{n} represents the R^{n} (reporter emission intensity/passive reference emission intensity) compared to R_{n} (background reporter emission intensity/passive reference emission intensity - calculated in no template controls-) (Applied Biosystems, 1998).

<table>
<thead>
<tr>
<th>Approx. no of L. monocytogenes molecules</th>
<th>Confidence interval</th>
<th>Signal ratio</th>
<th>hly system (FAM)</th>
<th>IAC system (VIC)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Approx. no of L. monocytogenes molecules</td>
<td>Lower</td>
<td>Upper</td>
<td>Limit</td>
<td>Signal ratio</td>
</tr>
<tr>
<td>3×10^{4}</td>
<td>29661</td>
<td>30340</td>
<td>9</td>
<td>22.38</td>
</tr>
<tr>
<td>3×10^{3}</td>
<td>2893</td>
<td>3108</td>
<td>9</td>
<td>25.91</td>
</tr>
<tr>
<td>3×10^{2}</td>
<td>267</td>
<td>334</td>
<td>9</td>
<td>30.17</td>
</tr>
<tr>
<td>60</td>
<td>45</td>
<td>76</td>
<td>9</td>
<td>32.49</td>
</tr>
<tr>
<td>30</td>
<td>20</td>
<td>41</td>
<td>9</td>
<td>34.25</td>
</tr>
<tr>
<td>15</td>
<td>8</td>
<td>23</td>
<td>9</td>
<td>35.57</td>
</tr>
<tr>
<td>8</td>
<td>3</td>
<td>13</td>
<td>9</td>
<td>36.07</td>
</tr>
<tr>
<td>4</td>
<td>1</td>
<td>8</td>
<td>5</td>
<td>35.56</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>3</td>
<td>4</td>
<td>35.56</td>
</tr>
</tbody>
</table>

Calculated for the expected number of template molecules at each dilution at the 95% confidence level.

FAM C_{T} values were calculated with a prefixed threshold at 0.035, and a baseline from cycles 3 to 15.

VIC C_{T} values were calculated with a prefixed threshold at 0.035, and a baseline from cycles 3 to 23.

hly negative amplifications were excluded from Mean and SD calculations.

Quantification capacity of L. monocytogenes by duplex RTi-PCR assay. We assessed the suitability of the developed duplex RTi-PCR assay to accurately quantify L. monocytogenes using decreasing amounts of L. monocytogenes strain UdG 1010 genomic DNA (equivalent to 3×10^{4}, 3×10^{3}, 3×10^{2}, 60 and 30 target DNA molecules per reaction). Table 1 shows FAM (hly) and VIC (IAC) C_{T} and ΔR_{n} values from a total of 9 replicates of 3 independent experiments. As expected, there was an excellent linearity between initial L. monocytogenes
DNA molecules and FAM CT values down to 30 target molecules, as indicated by the square regression coefficient ($R^2 = 0.997$). This, together with the small SD obtained in the replicates and in independent experiments, indicated that quantification of \textit{L. monocytogenes} can be achieved by the developed duplex RTi-PCR assay. The quantification limit of the duplex assay, further confirmed by theoretical statistical analyses, can be established at 30 cells, the same as that previously reported for the corresponding uniplex assay (Rodríguez-Lázaro et al., 2004a) and similar to most of the quantitative RTi-PCR systems published to date (e.g. Bach et al., 2003; Hernández et al., 2003).

The PCR amplification efficiency can be calculated through the equation $E = 10^{1/s-1}$, where $s$ is the slope of the linear regression curve (Klein et al., 1999). When the efficiency is optimal (1.00), the slope is -3.322 (Knutsson et al., 2002). The slope of the duplex PCR assay (-3.916) denoted a 0.800 efficiency, slightly below the efficiency of the corresponding \textit{hly} assay (0.916) which signifies an 87.38% of the uniplex PCR efficiency. Comparison of \textit{hly} and IAC amplification profiles (Table1) evidenced that the \textit{hly}-specific reaction predominates above that for IAC. \textit{hly} (FAM) $\Delta R_n$ values were above 0.74 in all reactions (overall, $\Delta R_n$ 0.84 $\pm$ 0.09), whereas IAC (VIC) $\Delta R_n$ values gradually decreased from 0.73 in reactions containing 4 \textit{L. monocytogenes} DNA molecules down to 0.10 when 30,000 \textit{L. monocytogenes} DNA molecules were added to the reaction. This was in contrast with VIC CT values that, as expected, were highly stable (overall CT value = 33.56 $\pm$ 0.69).

**Use of the duplex RTi-PCR assay for the quantitative detection of \textit{L. monocytogenes} in the presence of different culture broths.** We tested the applicability of our duplex RTi-PCR assay to quantitative detect \textit{L. monocytogenes} in the presence of four different culture media typically used for the microbiological detection or quantification of this species: Fraser, half-Fraser, BHI and buffered peptone water. We performed duplex RTi-PCR using as template 300 copies of genomic DNA from \textit{L. monocytogenes} strain UdG 1010 and 100 molecules of IAC, and adding to the PCR mix 1 µl of either media or double-distilled water (as control). Both FAM and VIC CT values obtained in the presence of BHI broth, buffered peptone water or double-distilled water were not statistically different (p<0.05). Based on FAM values, 283 $\pm$ 25 \textit{L. monocytogenes} DNA molecules were calculated in each sample, which corresponded to 94.00 $\pm$ 8.00 % of the real value. VIC CT values proved the lack of inhibition due to BHI and particularly, buffered peptone water, specified for use in the \textit{L. monocytogenes} enumeration standard (Anonymous, 1998). In contrast, reactions containing Fraser or half-Fraser media (specified for use in the \textit{L. monocytogenes} detection standard, Anonymous, 1996) showed \textit{hly} and IAC positive amplification, although with significantly higher FAM and VIC CT values (above 40 i.e. below the quantification limit). This confirmed that these media act as a potent PCR inhibitors. Several of its components e.g. LiCl, nalidixic acid, acriflavin and excucin have also been shown to inhibit PCR (Rossen et al., 1992).

**Quantitative detection of \textit{L. monocytogenes} by the duplex assay in different foods and assessment of the PCR results by the IAC.** We also assessed the applicability of the duplex RTi-PCR assay for the quantification of \textit{L. monocytogenes} in different food matrixes typically contaminated by this pathogen: meat and meat products (raw pork meat, pork fermented sausage, frankfurter sausage and pork cook ham) and fish and fish products (raw salmon and cold-smoked salmon). These food matrixes were artificially contaminated with decreasing concentrations of of \textit{L. monocytogenes} strain UdG 1010 (approximately from $3\times10^7$, $3\times10^6$ and $3\times10^5$ cfu/g), and 1 µl of a 1:10 dilution (w/v) of the homogenate in buffered peptone water was directly used for the duplex RTi-PCR assay. In parallel, the
standard microbiological procedure (Anonymous, 1998) was performed to ensure that the lack of amplification was not caused by the absence of *L. monocytogenes* cells in the food sample. The results obtained from the meat products (pork fermented sausage, frankfurter sausage and pork cook ham) showed an adequate amplification signals for *L. monocytogenes* both considering each food sample separately and considering the results globally (Table 2). These results indicate that the quantitative detection of *L. monocytogenes* in meat products is achievable since the FAM C<sub>T</sub> results obtained were not statistically different (p<0.05) to those obtained from purified DNA solutions (see Tables 1 and 2). In addition, it was observed non-statistically different results for IAC (p<0.05) to those previously obtained for DNA solutions (overall VIC C<sub>T</sub> values of 33.64 ± 0.57). However, when raw food materials (pork meat and salmon) and cold-smoked salmon were assayed, it was observed a lack of *L. monocytogenes* (FAM) amplification signal in all the cases. In addition, no IAC (VIC) amplification signal was obtained either, indicating that the failure in detecting *L. monocytogenes* was due to inhibition of the duplex RTi-PCR assay in these particular food matrices. This finding emphasizes the importance of including an IAC as it provides a consistent tool to distinguish between real and false negative results.

**TABLE 2.** Detection of *L. monocytogenes* by RTi-PCR assay in different food matrices.

<table>
<thead>
<tr>
<th>Approx. no. of <em>L. monocytogenes</em> cfu/g</th>
<th>Approx. no. of <em>L. monocytogenes</em> cfu/reaction</th>
<th><em>hly</em> C&lt;sub&gt;T&lt;/sub&gt; value&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Pork fermented sausage</td>
<td>Frankfurter sausage</td>
</tr>
<tr>
<td></td>
<td>Mean  SD</td>
<td>Mean  SD</td>
</tr>
<tr>
<td>3×10&lt;sup&gt;7&lt;/sup&gt;</td>
<td>26.05 0.30</td>
<td>25.94 0.23</td>
</tr>
<tr>
<td>3×10&lt;sup&gt;6&lt;/sup&gt;</td>
<td>29.26 0.31</td>
<td>29.68 0.37</td>
</tr>
<tr>
<td>3×10&lt;sup&gt;5&lt;/sup&gt;</td>
<td>33.01 0.62</td>
<td>29.43 0.59</td>
</tr>
<tr>
<td></td>
<td>Pork cook ham</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Mean  SD</td>
<td>Mean  SD</td>
</tr>
<tr>
<td>3×10&lt;sup&gt;7&lt;/sup&gt;</td>
<td>26.05 0.30</td>
<td>26.05 0.30</td>
</tr>
<tr>
<td>3×10&lt;sup&gt;6&lt;/sup&gt;</td>
<td>29.43 0.17</td>
<td>29.46 0.39</td>
</tr>
<tr>
<td>3×10&lt;sup&gt;5&lt;/sup&gt;</td>
<td>32.49 0.57</td>
<td>32.81 0.65</td>
</tr>
<tr>
<td>overall</td>
<td>Mean  SD</td>
<td>Mean  SD</td>
</tr>
<tr>
<td></td>
<td>26.00 0.27</td>
<td>29.46 0.39</td>
</tr>
</tbody>
</table>

<sup>a</sup> FAM C<sub>T</sub> values were calculated with a prefixed threshold at 0.035, and a baseline from cycles 3 to 15.

In conclusion, we have further developed and optimised a previously reported RTi-PCR assay for quantitative detection of *L. monocytogenes* (Rodríguez-Lázaro et al., 2004a) with the inclusion of an IAC. This improved duplex RTi-PCR assay maintains the same specificity, sensitivity and quantification ability characteristics of the original assay whilst providing the capacity to identify false negative results or target underestimation due to the presence of PCR inhibitors in the samples and can be also applied in the sensitive detection of PCR inhibitors in the samples. Finally, using artificially contaminated food matrices, we provided direct evidence that the duplex assay can be a consistent alternative tool for the routine quantitative detection of *L. monocytogenes* in food products.

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