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1	Protective cultures against foodborne pathogens in a nitrite reduced fermented meat
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4	Ivana Nikodinoska ^{a,b} , Loredana Baffoni ^a , Diana Di Gioia ^a , Beatriz Manso ^b , Lourdes García-
5	Sánchez ^b , Beatriz Melero ^{b*} , Jordi Rovira ^b
6	
7	^a Department of Agricultural and Food Sciences (DISTAL), University of Bologna, viale
8	Fanin 42, 40136 Bologna, Italy
9	^b Department of Biotechnology and Food Science, University of Burgos, Plaza Misael
LO	Bañuelos s/n, 09001 Burgos, Spain
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L3	*Corresponding Author: bmelero@ubu.es
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Abstract

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In the present work, a combined hurdle approach for fermented meat preservation was investigated. Challenge tests were performed in *Chorizo* sausage model using the maximum allowed NaNO₂ amount (150mg/kg), a reduced amount (75 mg/kg) and no nitrite, with and without protective cultures inoculation. Cocktail strains of L. monocytogenes and Salmonella spp. were used as indicator strains. In a nitrite reduced sausage model, L. monocytogenes growing trend did not significantly change (p>0.05) when compared with that containing higher nitrite concentration (150 mg/kg NaNO₂). The addition of L. plantarum PSC20 significantly lowered L. monocytogenes growth when compared with control batches without PCS20 (p<0.05), obtaining 3.84 log cfu/g and 2.62 log cfu/g lower counts in the batches with 150mg/kg NaNO₂ and 75mg/kg NaNO₂ respectively. None of the protective cultures demonstrated in situ antagonistic activity against Salmonella spp. This work pointed out that the reduction of nitrites with the combined use of a protective culture could be a feasible approach to control L. monocytogenes growth in fermented meat foods. **Keywords:** Protective cultures; nitrite reduction; *Listeria monocytogenes*; *Salmonella* spp.; fermented pork meat

1. Introduction

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In the era where demand for ready to eat and preservative free products is constantly 38 growing, the microbiological food safety has to be guaranteed, proportionally with this 39 40 ongoing trend. In the recently published European Food Safety Authority (EFSA) foodborne outbreak report, referred to 2016, Salmonella spp. human infections had the same high level 41 of the previous year (94.530 confirmed cases), whereas human listeriosis, caused mainly by 42 Listeria monocytogenes, showed a 9.3% increase (2.536 confirmed cases) (EFSA, 2017). 43 Despite the relatively low incidence of listeriosis, compared with the number of 44 45 campylobacteriosis and salmonellosis cases, its importance is due to the severity of the disease and the higher case-fatality rate (Baffoni et al. 2017; D'Ostuni et al., 2016; EFSA, 46 2017). 47 48 Curing with nitrite is the most used approach to control foodborne pathogens in the meat (Honikel, 2008). Nitrites have additional functions in the meat, as they help to prevent lipid 49 oxidation and rancidity, guarantee a bright red color and a typical "cured" flavor (Sebranek & 50 51 Bacus, 2007). Although nitrites are widely used in the meat industry, they are classified by International Agency for Cancer Research as potentially carcinogenic agents (IARC, 2010), 52 due to their ability to react with amines in the gastrointestinal tract, resulting in N-53 nitrosamines formation. Nitrites, hitherto, are the most effective solution against C. botulinum 54 55 growth in meat products (EFSA 2003; Hospital, Hierro & Fernández, 2014; Hospital, Hierro, 56 Stringer & Fernández, 2016). Therefore, 150 mg/kg NaNO₂ and 300 mg/kg NaNO₃ were authorized as maximum added levels in meat in Europe until May 2018 (EFSA, 2003; 57 European Commission, 2011). Starting from May 2018, a new regulation, proposed by the 58 59 Danish authorities in 2015, was approved and the maximum accepted nitrite level in fermented salami is now 100 mg/kg (European Commission, 2018). Additionally, the EC 60 Regulation N° 889/2008 for organic meat products, establishes 80 mg/kg for added nitrite and 61

62 50 mg/kg for residual nitrite (European Commission, 2008). The U.S. FDA accepts a maximum level of 200 mg/kg NaNO₂ and 500 mg/kg NaNO₃ in meat finished products 63 (CFR, 2018). Although outbreaks regarding food poisoning by nitrite derived from meat 64 products are not described in the literature, unintentional poisoning has been reported upon 65 eating homemade sausages (Cvetković, Živković, Lukić, & Nikolić, 2018). 66 Therefore, meat industries are challenged to employ healthier and safer approaches for meat 67 68 preservation. In the attempt of finding alternatives to nitrites for fermented food preservation, several authors suggested the use of lower nitrite levels in combination with other compounds 69 70 or processing technologies, in a way that antimicrobial properties against the common foodborne pathogens could be guaranteed without alteration of sensory qualities (Alahakoon, 71 72 Jayasena, Ramachandra & Jo, 2015; Cavalheiro et al., 2015). Lactic acid bacteria (LAB) with 73 demonstrated *in vitro* antimicrobial activity against a wide spectrum of foodborne pathogens 74 (Leroy, Geyzen, Janssens, De Vuyst & Scholliers, 2013) as well as the addition of natural extracts or phytochemicals are the mostly studied approaches for the development of 75 76 innovative processed meat products (Alahakoon et al., 2015; Gaggia, Di Gioia, Baffoni & Biavati, 2011; Oliveira, Ferreira, Magalhães & Teixeira, 2018). However, several natural 77 extracts may contain even more than the allowed nitrate amount, thus the nitrosamine 78 formation is questioned (Bedale, Sindelar, & Milkowski, 2016). LAB strains with 79 demonstrated sensorial or health promoting properties are approved by FDA as Generally 80 81 Recognized as Safe (GRAS) and by EFSA with the Qualified Presumption of Safety (QPS) status (EFSA, 2018; FDA, 2018). 82 In the present work, we studied the effectiveness of a combined hurdle approach, i.e. a 50% 83 84 reduction of nitrites plus the addition of previously characterized *Lactobacillus* strains (Lactobacillus plantarum PCS20 or Lactobacillus delbrueckii DSM 20074), against common 85 foodborne pathogens in Chorizo, a dry fermented sausage produced in Spain. 86

2. Material and methods

88 2.1 Bacterial strains

- 89 L. plantarum PCS20 (MSCL P977) and L. delbrueckii DSM 20074 were used as protective
- 90 cultures for their demonstrated anti-microbial activity against several pathogens (Di Gioia et
- al., 2016; Savino et al., 2011). They were grown in de Man Rogosa Sharpe medium (MRS,
- 92 Oxoid Ltd., Basingstoke, England) in anaerobic conditions (Anaerogen, AN0025A, Oxoid),
- 93 at 37 °C for 48 h.
- A cocktail of *Listeria monocytogenes* strains has been used: *L. monocytogenes* CECT 5366
- 95 (serovar 4b, source: human), CECT 934 (serovar 4a, sourse: brain of sheep with circling
- disease), CECT 4032 (serovar 4b, source: associated with case of meningitis after eating soft
- 97 cheese) and LTA0020 (isolated from poultry minced meat in Burgos, Spain), already used in
- 98 similar studies (Melero, Diez, Rajkovic, Jaime, & Rovira, 2012; Melero, Vinuesa, Diez,
- 99 Jaime, & Rovira, 2013). The strains were grown at 37°C in Brain Heart Infusion Broth (BHI,
- Oxoid). For evaluation of viable cell population Chromogenic *Listeria* agar (Oxoid)
- supplemented with OCLA (ISO) Selective Supplement (SR 0226E, Oxoid) and Brillance
- 102 Listeria Differential Supplement (SR 0228E, Oxoid) was used.
- Four Salmonella strains were also employed in the challenge tests. All strains were isolated
- from meat and cheese products in Burgos. Bacterial strains were grown at 37°C in BHI.
- Brilliance Salmonella agar (Oxoid) supplemented with Salmonella Selective Supplement (SR
- 106 0194, Oxoid) was used for the evaluation of viable cell population.
- 107 2.2 Study design
- 108 Two Challenge tests in sausage prototypes were designed, referred to as 1 and 2. Challenge
- test 1 aimed at studying the effect of L. plantarum PCS20 against L. monocytogenes and
- Salmonella spp. in fermented sausages, both without nitrite addition and with 150 mg/kg of
- 111 nitrite. Challenge test 2 was focused on the effects of two protective cultures, L. plantarum

- PCS20 and L. delbrueckii DSM 20074, against L. monocytogenes strains in pork meat batters
- treated with 75 mg/kg and 150 mg/kg of nitrite. Challenge test protocols are detailed below
- 114 (2.3 and 2.4).
- 115 *2.3 Inocula preparation*
- 116 2.3.1 Pathogen strains
- Each L. monocytogenes and Salmonella spp. strain was grown at 37°C overnight in BHI
- broth up to 9 log cfu/ml. Cells were washed and suspended in sterile Ringer solution (Oxoid).
- For Challenge test 1, dilutions were performed in order to obtain a final concentration of 4.5
- log cfu/g in the meat batter (Figure 1), whereas for Challenge test 2, meat batter was
- inoculated with L. monocytogenes cocktail strains in order to obtain the final concentration of
- 122 3 log cfu/g (Figure 2).
- 123 *2.3.2 Protective cultures*
- 124 L. plantarum PCS20 and L. delbrueckii DSM 20074 were grown at 37°C overnight in MRS
- broth up to 9.5-10 log cfu/ml. Cells were washed and suspended to a final concentration of 6-
- 126 7 log cfu/g (Figure 2).
- 127 *2.4 Challenge tests*
- The batter was composed of ground pork meat and fat (70% and 30%, respectively) supplied
- by a meat processing company in Burgos (Spain). Spices were not used not to interfere with
- the results obtained.
- For Challenge test 1, the ground meat (4 kg) was divided in 2 trays, each containing 2 kg. In
- one tray, 2% NaCl was added whereas, in the other tray, meat was supplemented with 2%
- NaCl plus 150 mg/kg NaNO₂ (Figure 1). After homogenization in a vacuum mixer, each 2 kg
- portion was splitted in two: 1 kg was inoculated with L. plantarum PCS20 whereas the other
- kg was not inoculated with any protective culture. Subsequently, each kg was divided in 3
- batches (333 g), one inoculated with the cocktail of Salmonella strains, the second one with

- the *L. monocytogenes* strains and the last one was not inoculated with any pathogen (control).
- The 12 treatments and the relative acronyms are shown in Fig. 1.
- For Challenge test 2, the ground meat (4 kg) was divided in 2 trays of 2 kg meat each. 2 kg
- were amended with 2% of NaCl, 0.5% dextrose and 75 mg/kg NaNO₂ and 2 kg with 2% of
- NaCl, 0.5% dextrose and 150 mg/kg NaNO₂. Each tray was divided in two (1 kg each): one
- kg was inoculated with L. monocytogenes and the other kg was not inoculated with L.
- monocytogenes. Then each kg of meat was divided in three batches (333 g each) and
- submitted to different treatments: inoculated with PCS 20, with DSM 20074 and not
- inoculated with protective cultures. The 12 treatments and the relative acronyms are shown in
- Fig. 2. Each batch containing 333 g of meat batter was used to produce two sausages (two
- replicates per treatment). Sausages were then stuffed in collagen casings (45 mm diameter)
- 148 (Viscofan, Navarra, Spain). For Challenge test 1, the fermentation was performed for 2 days
- at 23°C, 95% humidity, followed by a short ripening of 6 days at 15°C and lower humidity
- 150 (80-75%). pH evaluation and microbiological analyses were performed at the following days:
- D0, D1, D2, D4, D6 and D8. Differently, for Challenge test 2, the fermentation was studied
- for 2 days followed by 5 days of short ripening in the same conditions as for the Challenge
- test 1. pH evaluation and microbiological analyses were performed at the following days: D0,
- 154 D3, D5 and D7.
- 155 *2.5 pH analysis*
- pH was measured with a pin electrode of a pHmeter (micropH2001, Crison, Barcelona,
- Spain) inserted directly 3 times into the sample.
- 158 *2.6 Microbiological Analysis*
- Meat samples (10 g per sampling point) were aseptically removed from each *Chorizo* (two
- sausages per treatment) and homogenized in 90 ml of Buffered Peptone Water (BPW; AES
- Laboratoire, Combourg, France) for 2 min in a sterile plastic bag using a Smasher (AES

Laboratoire). For cell counts, decimal dilutions (1:10 in BPW) of the meat homogenate were prepared and aliquot of 100 µl were inoculated onto selective solid agar plates for, both, lactic acid bacteria and for pathogens growth. The counts were performed in triplicate. Lactic acid bacteria were counted on MRS agar plates, incubated anaerobically for 48 h at 37°C. Randomly picked colonies were subjected to morphological and PCR analysis with LAB specific primers (data not shown). Previously described selective solid medias were used for L. monocytogenes and Salmonella spp. counts determination. Then, plates were incubated for 24h and for 48h, respectively, at 37°C. ISO protocols were used for the detection of natural contamination in not artificially inoculated batches: ISO 11290-1:1996 (ISO, 1996) and ISO 6579:2002 (ISO, 2002) for L. monocytogenes and Salmonella spp., respectively. 2.7 Statistical analysis The results of microbiological analysis, for each sampling point, were obtained from two chorizo replicates per treatment; for each replicate counts were performed in triplicate. Data were subjected to one-way ANOVA analysis. Differences among means were tested by Duncan's multiple range test (significance p<0.05). All the analyses obtained from the Challenge tests were performed using the Statistica 8.0 (StatSoftInc., USA). Results of statistical analysis are presented as mean value \pm standard deviation. 3. Results 3.1 Challenge test 1 3.1.1 pH analysis No differences in pH were observed during the fermentation and short ripening process (data not shown). Considering the slight decrease of pH observed, 0.5% of dextrose was added in

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186 pork meat batter in Challenge test 2 with the aim of stimulating the Lactobacillus growth and acidification. 187 3.1.2 Microbiological analysis 188 189 The growing trend of L. monocytogenes and Salmonella spp. in Challenge test 1 is shown in Figure 3. Both pathogens demonstrated ability to survive and colonize the pork meat in the 190 sausage model. 191 Regarding L. monocytogenes growth, a significantly lower counts (p<0.05) of 0.95 and 2.78 192 log cfu/g, were observed at day 4 and 6, respectively, in the batch with 150 mg/kg NaNO₂ 193 194 and PCS20 (NLP) when compared with the batch containing nitrite but without PCS20 (NL) (Figure 3A). Moreover, considering the initial inoculum, in the NL batch, an increase of 3.55 195 log cfu/g of L. monocytogenes counts was observed, whereas this increase was of 1.96 log 196 197 cfu/g in the NL+ batch P (Figure 3A) at the last sampling time (D8). Comparing control batches without nitrate addition, P+L and L, significantly (p<0.05) lower L. monocytogenes 198 counts of 0.60 and 0.52 log cfu/g, were observed at day 4 and 6, respectively, whereas no 199 200 significant differences were observed at D8. Lower L. monocytogenes growth was observed in batches where NaNO₂ was added 201 (NL/NL+P) in comparison with batches without additives (L/P+L). At the last sampling day 202 (D8), significant (p<0.05) decrease of L. monocytogenes counts of 2.37 log cfu/g was 203 observed when comparing NL+P and P+L batches, whereas significant (p<0.05) decrease of 204 205 0.58 log cfu/g was observed when comparing batches NL Ctr and L. Salmonella spp. counts within the study period are shown in Figure 3B. L. plantarum PCS20 206 did not show antimicrobial activity against Salmonella spp. growth. However, nitrites 207 208 demonstrated a significant decrease (p<0.05) of Salmonella spp. growth (1.23 log cfu/g) in

N+S batch in comparison with batch S at D8.

- 210 Initial counts of LAB in the meat without protective culture were between 3-4.5 log cfu/g.
- The level of PCS20 inoculum was 5.6-5.9 log cfu/g. After 3 days, when the fermentation
- 212 conditions were settled, LAB counts increased in all batches of 2.5-3.5 log cfu/g, reaching
- values in the range 7-9 log cfu/g in batches with protective culture and 7-8 log cfu/g in
- uninoculated batches, at the end of the study (data not shown).
- 215 3.2 Challenge test 2
- 216 *3.2.1 pH analysis*
- pH trend in the meat subjected to different treatments is shown in Table 1. As expected, the
- 218 addition of 0.5% dextrose caused a significant pH reduction at D7 (from 5.80 to 5.05;
- p<0.05), in all batches where L. plantarum PCS20 was inoculated. Differently, the addition
- of *L. delbrueckii* DSM 20074 did not lead to a significant pH reduction (p>0.05).
- *3.2.2 Microbiological analysis*
- Figure 4 shows the trend of L. monocytogenes inoculated at 3 log cfu/g in all batches.
- 223 Comparing batches containing 75 mg/kg NaNO₂, with and without PSC20 (batches ½NL+P
- and ½NL Ctr, respectively, Fig. 4A), a significantly lower counts (p<0.05) of 2.20 and 2.62
- log cfu/g of the inoculated L. monocytogenes were observed at day 3 and 5, respectively, in
- 226 the batch where PCS20 was inoculated (½NL+P); this reduction was maintained until D7.
- Interestingly, considering the initial inoculum, the pathogen counts increase of only 1.61 log
- cfu/g in the batch ½NL+P compared with a 3.99 log cfu/g increase in the batch ½NL Ctr, at
- D7. On the other hand, in batches with higher nitrites concentration a significantly lower
- counts of *L. monocytogenes* of 3.93 log cfu/g were observed at D5, in batch containing
- PCS20 as protective culture (NL+P) in comparison with batch without PCS20 (NL Ctr), with
- a final decrease of L. monocytogenes of 3.84 log cfu/g at D7. In summary, at the end of the
- study, the pathogen growing trend was not statistically different (p>0.05) when compared
- batches with 75 or 150 mg/kg of nitrites (½NL Ctr and NL Ctr), while, in batches with

PCS20, L. monocytogenes counts were higher in ½NL+P compared with NL+P (difference of 235 1.49 log cfu/g). 236 Figure 4B shows the *L. monocytogenes* growth in pork meat batter with 150 mg/kg or 75 237 mg/kg NaNO₂ with or without L. delbrueckii DSM 20074 inoculum. At the end of the study, 238 no significant differences in L. monocytogenes growth were observed among batches. 239 Counts of LAB growth were under the detection limit (<2 log cfu/g) in the control batches 240 without protective culture inoculum at D0; whereas LAB counts were in the range 6-7 log 241 cfu/g in the batches inoculated with PCS20 at D0 (Table 2). At the end of the study, LAB 242 243 counts reached 7-8 log cfu/g in batches without PCS20, and 8-9.2 log cfu/g in batches with PCS20. Batches inoculated with DSM 20074 did not reach the same LAB count level as 244 PCS20. In particular, 5.89 log cfu/g were obtained in the control batch with 150 mg/kg 245 246 NaNO₂ and 6.36 log cfu/g in that with 75mg/kg NaNO₂, at D7. These counts are almost 3 log lower than those obtained for PCS20. 247 Similarly to the previous experiment, significant differences (p<0.05) were observed between 248 D1 and D3, i.e. in the final part of the fermentation period (3rd day). At the end of the short 249 ripening period, LAB reached counts in the range 7-9 log cfu/g. 250 251 4. Discussion 252

The aim of the present work was to evaluate the possibility of using protective cultures to eliminate or reduce nitrite amount in fermented meat products. For this purpose, the biopreservative activity of previously characterized LAB strains, *L. plantarum* PCS20 and *L. delbrueckii* DSM 20074, was studied against *L. monocytogenes* and *Salmonella* spp. in a dry fermented sausage model without nitrite, with half (75 mg/kg) and maximum (150 mg/kg) allowed nitrite amount considering the maximum amounts allowed in Europe Until May 2018.

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The results showed that the addition of L. plantarum PCS20 as protective culture in nitritefree sausages, artificially contaminated with pathogen, is capable of significantly reducing the pathogen load after 4 and 6 days from the beginning of the fermentation, although the same effect was not observed at D8. On the contrary, the antimicrobial activity of PCS20 was not observed against the cocktail of Salmonella strains, whereas their growth was significantly (p<0.05) reduced in the presence of 150 mg/kg nitrites. Interestingly, Hospital et al. (2014) obtained complete Salmonella inactivation using a halved nitrite amount (75 mg/kg) in fermented sausages at the end of the storage period. Other works showed the ineffectiveness of commercial protective cultures, as well as of meat-isolated Lactobacillus strains, against Salmonella spp., when inoculated in different meat models (Dias, Duarte, Ramos, Martins Santos & Schwan, 2013; Kotzekidou & Bloukas, 1998). The outcomes of this study support the Hugas (1998) consideration on the hurdle effect strategy. Our study also shows that it is possible to reduce *Listeria* counts by inoculating the meat with L. plantarum PCS20 and a halved amount of nitrite (75 mg/kg). This result is particularly important considering the EC decision of adopting more stringent criteria for potential carcinogenic additives. Therefore, the combination of a protective culture with a reduced nitrite amount is an effective hurdle approach in fermented sausage production that may allow both to reduce pathogen load and to have the known positive effects of nitrites, such as the bright color. The anti-Listeria activity observed is in agreement with a recent work (Giello, La Storia, De Filippis, Ercolini & Villani, 2018) that showed the effectiveness of the bacteriocin-producing Lactobacillus curvatus 54M16 strain in fermented sausages. Several authors pointed out that bacteriocin action can be hindered in carnis by bacteriocin binding to food matrixes or degradation by proteases or their production can be prevented by nitrites (Galvez, Abriouel, Lopez, & Ben, 2007; Kouakou et al., 2009). Therefore, non-bacteriocin producing strains

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showing anti-listerial activity can be of great importance in fermented meat production, in particular in the presence of nitrites. This is the case of L. plantarum PCS20 strain, that does not produce bacteriocins (Cho, G.S., Huch, M., Hanak, A., Holzapfel, W.H., & Franz, C.M.A.P. 2010) and exerts anti-microbial activity in the presence of a reduced amount of nitrites. Its anti-microbial activity against L. monocytogenes can be attributed to cell-to-cell contact mechanisms or the production of organic acidic metabolites. An additional strength of our study is the use of four different L. monocytogenes strains, belonging to different serovars (Lianou & Koutsoumanis 2013; Scott et al. 2005). Moreover, our work confirmed that dextrose is an important pH lowering agent, allowing to reach pH values between 4.5 and 5.5, a range in which nitrite is mainly in the undissociated state, possessing the greatest antibacterial activity. Moreover, a rapid pH drop below 5.1 is considered as a desirable acidification rate for protective cultures in fermented meat products (Ammor and Mayo 2007). On the other hand, the inability of L. delbrueckii DSM 20074 strain to demonstrate a significant pH lowering, resulted in an antagonistic failure against L. monocytogenes at the end of the study, even when 150 mg/kg of NaNO2 were added. Our study supports the outcomes of a recent survey (Hung et al. 2016), in which meat industry stakeholders expressed interest in the development of innovative and healthier processed meat products but asked the scientific community to provide additional evidences of the microbiological safety of developed approaches. Consumers are important players in industrial innovation shaping, thus the taste and the microbiological safety are the most important criteria for the novel food formulations (Bedale et al. 2016, Hung et al. 2016).

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5. Conclusions

This work pointed out that a combined approach based on half of the allowed nitrite amount and of protective culture may be effective in a dry-fermented meat product (*chorizo*) to

reduce the growth of *L. monocytogenes*, a pathogen with high case fatality incidence and causing severe diseases. This study has also shown that the effectiveness of nitrites against this pathogen is not related to their amount; the inoculation with lactic acid bacteria contributing to pH lowering and to reach the effective dissociation state of nitrite is probably a crucial factor for their effectiveness. However, further studies aimed at better elucidating the anti-microbial mechanisms against pathogens in food matrix need to be pursued.

In conclusion, the results obtained from this study will provide additional scientific evidence in the evaluation of microbiological and preservative risks/benefits in fermented meat products. The proposed combined hurdle approach (a reduced amount of nitrite plus the inoculation of a protective culture) is promising for innovative fermented meat products development.

Conflict of interest

The authors declare that they have no conflict of interest.

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465 Figure 1 Study design of the 12 treatments related to Challenge test 1. Legend: Ctr=meat batter; 466 Ctr-N=150 mg/kg NaNO₂ added; Ctr-P=PCS20 added; Ctr-NP=PCS20+150 mg/kg NaNO₂ added; 467 468 L=L. monocytogenes added; N+L=150 mg/kg NaNO₂+ L. monocytogenes added; P+L=PCS20+L. monocytogenes added; NP+L=150 mg/kg NaNO₂+ PCS20+L. monocytogenes added; S=Salmonella 469 spp. added; N+S=150 mg/kg NaNO₂ + Salmonella spp. added; P+S=PCS20+Salmonella spp. added; 470 NP+S=150 mg/kg NaNO₂+PCS20+Salmonella spp. added. For each condition two sausages were 471 472 prepared and processed. 473 Figure 2 Study design Challenge test 2. Legend: N Ctr=meat batter added with 150mg/kg NaNO₂; 474 475 ¹/₂N Ctr=75 mg/kg NaNO₂ added; N+P=150mg/kg NaNO₂+PCS20 added; ¹/₂N+P=75mg/kg NaNO₂+PCS20 added; N+D=150mg/kg NaNO₂+DSM 20074 added; ½N+D=75mg/kg NaNO₂+DSM 476 20074 added; NL Ctr=150mg/kg NaNO₂+L. monocytogenes added; ½NL Ctr=75mg/kg NaNO₂+L. 477 monocytogenes added; NL+P=150mg/kg NaNO₂+L. monocytogenes+PCS20 added; NL+P=75mg/kg 478 479 NaNO₂+L. monocytogenes+ PCS20 added; NL+D=150mg/kg NaNO₂+L. monocytogenes+DSM 480 20074 added; ½NL+D=75mg/kg NaNO₂+DSM 20074 added. For each condition two sausages were 481 prepared and processed. 482 483 Figure 3 Antimicrobial activity of L. plantarum PCS20 against L. monocytogenes and Salmonella spp. in dry fermented sausage with and without 150 mg/kg NaNO₂. 484 485 A) L. monocytogenes counts within the ripening period. L=L. monocytogenes added; N+L=150 mg/kg NaNO₂+ L. monocytogenes added; P+L=PCS20+L. monocytogenes added; NP+L=150 mg/kg 486 NaNO₂+ PCS20+L. monocytogenes added; B) Salmonella spp. counts within the ripening period. 487 S=Salmonella spp. added; N+S=150 mg/kg NaNO₂+ Salmonella spp. added; 488 489 P+S=PCS20+Salmonella spp. added; NP+S=150 mg/kg NaNO₂+PCS20+Salmonella spp. added.

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Figure Captions

Figure 4 Antimicrobial activity of L. plantarum PCS20 (A) and L. delbrueckii DSM20074 (B) against 491 492 L. monocytogenes in dry fermented sausage added with 75 or 150 mg/kg NaNO₂. A) L. 493 monocytogenes counts in batches inoculated with or without L. plantarum PCS20. Legend: NL 494 Ctr=150mg/kg NaNO₂+L. monocytogenes added; ½NL Ctr=75mg/kg NaNO₂+L. monocytogenes added; NL+P=150mg/kg NaNO₂+L. monocytogenes+PCS20 added; ½NL +P=75mg/kg NaNO₂+L. 495 496 monocytogenes+PCS20 added. B) L. monocytogenes counts in batches inoculated with L. delbrueckii 497 DSM 20074. Legend: N+D=150mg/kg NaNO₂+DSM 20074 added; ½N+D=75mg/kg NaNO₂+DSM 498 20074 added; NL Ctr=150mg/kg NaNO₂+L. monocytogenes added; ½NL Ctr=75mg/kg NaNO₂+L. monocytogenes added; NL+P=150mg/kg NaNO₂+L. monocytogenes+PCS20 added; NL+P=75mg/kg 499 500 NaNO₂+L. monocytogenes+PCS20 added; NL+D=150mg/kg NaNO₂+L. monocytogenes+DSM 501 20074 added; ½NL+D=75mg/kg NaNO₂+L. monocytogenes+DSM 20074 added.

Table 1. Challenge test 2. The trend of pH during the fermentation and ripening period

Batches**	Days *	Days *				
	0	3	5	7		
N Ctr	$5.96 \pm\! 0.03^{\rm B}$	$5.78 \pm 0.03^{\rm C}$	$6.12 \pm \! 0.06^{\rm A}$	6.03 ± 0.06^{B}		
½N Ctr	$5.90 \; {\pm} 0.09^{\mathrm{B}}$	$5.99 \; {\pm}0.03^{\rm B}$	6.11 ± 0.04^{A}	$5.90 \; {\pm} 0.01^{\rm B}$		
½NL Ctr	5.77 ± 0.08^{B}	$5.89 \pm\! 0.02^{\rm A}$	5.92 ± 0.02^{A}	5.91 ± 0.02^{A}		
NL Ctr	$5.85 \pm 0.03^{\mathrm{B}}$	5.99 ± 0.01^{A}	$6.10 \pm \! 0.04^{\rm A}$	$5.85 \; {\pm}0.02^{\rm B}$		
N+P	$5.86 \pm\! 0.06^{A}$	$5.44 \; {\pm}0.02^{\rm B}$	5.23 ± 0.02^{C}	5.02 ± 0.04^{D}		
½N+P	$5.85 \; {\pm}0.07^{A}$	$5.28 \pm\! 0.06^{B}$	5.21 ± 0.02^{B}	5.05 ± 0.02^{C}		
NL+P	5.77 ± 0.06^{A}	5.31 ± 0.01^{B}	5.14 ± 0.04^{C}	5.09 ± 0.04^{C}		
½NL+P	5.83 ± 0.01^{A}	$5.30 \pm\! 0.01^{\rm B}$	5.16 ± 0.01^{C}	5.02 ± 0.01^{D}		
N+D	$5.87 \; {\pm}0.03^{\rm A}$	$5.93 \pm 0.04^{\rm A}$	5.93 ± 0.05^{A}	$5.89 \pm\! 0.04^{\rm A}$		
½N+D	5.80 ± 0.04^{B}	5.93 ± 0.02^{A}	5.93 ± 0.02^{A}	5.89 ± 0.05^{A}		
NL+D	6.05 ± 0.01^{A}	$5.94 \; {\pm}0.03^{\rm B}$	6.04 ± 0.05^{A}	$5.97 \; {\pm}0.03^{\mathrm{B}}$		
½NL+D	6.17 ± 0.04^{A}	5.91 ± 0.01^{B}	5.94 ± 0.04^{B}	5.80 ± 0.04^{C}		

Data are expressed as mean of n=3 measurements.

^{**}Batch: N Ctr=meat batter added with 150mg/kg NaNO₂; ½N Ctr=75 mg/kg NaNO₂ added; ½NL Ctr=75mg/kg NaNO₂+L.monocytogenes added; NL Ctr=150mg/kg NaNO₂+L.monocytogenes added; N+P=150mg/kg NaNO₂+PCS20 added; ½N+P=75mg/kg NaNO₂+PCS20 added; NL+P=150mg/kg NaNO₂+L.monocytogenes+PCS20 added; ½NL+P=75mg/kg NaNO₂+L.monocytogenes+ PCS20 added; N+D=150mg/kg NaNO₂+DSM 20074 added; ½N+D=75mg/kg NaNO₂+DSM 20074 added; NL+D=150mg/kg NaNO₂+L.monocytogenes+DSM 20074 added; $\frac{1}{2}$ NL+D=75mg/kg NaNO₂+DSM 20074 added. ****A,B,C: Mean values in the same row (corresponding to the same batch) differ significantly (p < 0.05).

Table 2. Challenge test 2. LAB counts (log cfu/g) within the 7 days of fermentation and ripening period

Batches**	Days *			
	0	3	5	7
N Ctr	$<2 \pm 0.00^{D}$	6.63 ± 0.25^{C}	7.82 ± 0.12^{B}	7.89 ± 0.12^{A}
½N Ctr	$<2 \pm 0.00^{B}$	7.18 ± 0.46^{A}	7.60 ± 0.17^{A}	7.09 ± 0.34^{A}
NL Ctr	$<2 \pm 0.00^{D}$	6.59 ± 0.28^{C}	8.15 ± 0.07^{A}	$7.38 \pm\! 0.29^{B}$
½NL Ctr	$<2 \pm 0.00^{B}$	6.89 ± 0.27^{A}	7.15 ± 0.51^{A}	7.43 ± 0.28^{A}
N+P	6.34 ± 0.15^{C}	8.87 ± 0.20^{B}	9.09 ± 0.12^{AB}	9.13 ± 0.10^{A}
¹/₂N+P	6.34 ± 0.13^{D}	9.06 ± 0.13^{B}	9.26 ± 0.06^{A}	8.03 ± 0.13^{C}
NL+P	6.59 ± 0.17^{C}	$8.86 \pm\! 0.12^{B}$	9.10 ± 0.07^{A}	9.14 ± 0.09^{A}
¹/₂NL+P	6.61 ± 0.13^{B}	$9,04 \pm 0.07^{A}$	9.04 ± 0.08^{A}	9.17 ± 0.08^{A}
N+D	$5.71 \pm 0.26^{\circ}$	6.49 ± 0.22^{B}	7.29 ± 0.38^{A}	5.89 ± 0.37^{C}
¹/₂N+D	5.84 ± 0.11^{D}	$6.98 \pm\! 0.19^{B}$	7.44 ± 0.15^{A}	6.36 ± 0.13^{C}
NL+D	5.98 ± 0.11^{D}	6.52 ± 0.12^{C}	7.69 ± 0.28^{A}	$7.19 \; {\pm} 0.08^{\rm B}$
¹/2NL+D	6.09 ± 0.15^{C}	6.35 ± 0.04^{B}	7.44 ± 0.15^{A}	7.58 ± 0.13^{A}

^{*} Data are expressed as mean of n=3 measurements.

^{**}Batch N Ctr=meat batter added with 150mg/kg NaNO₂; ½N Ctr=75 mg/kg NaNO₂ added; N+P=150mg/kg NaNO₂+PCS20 added; ½N+P=75mg/kg NaNO₂+PCS20 added; N+D=150mg/kg NaNO₂+DSM 20074 added; ½N+D=75mg/kg NaNO₂+DSM 20074 added; NL Ctr=150mg/kg NaNO₂+L.monocytogenes added; ½NL Ctr=75mg/kg NaNO₂+L.monocytogenes added; NL+P=150mg/kg NaNO₂+L.monocytogenes+PCS20 added; NL+P=75mg/kg NaNO₂+L.monocytogenes+PCS20 added; NL+D=150mg/kg NaNO₂+L.monocytogenes+DSM 20074 added; ½NL+D=75mg/kg NaNO₂+DSM 20074 added.
****A,B,C: Mean values in the same row (corresponding to the same batch) differ significantly (p < 0.05).







