ABSTRACT

Storage stability of cooked meat sausages with 50 g marine oil/kg and two salt combinations: (1) 14.00 g NaCl/kg and 2.0 g sodium tripolyphosphate (TPP)/kg, (2) sodium reduced formulation with 6.08 g NaCl/kg, 4.92 g KCl/kg and 5.00 g TPP/kg were studied. In addition, effect of BHA or tocopherols as antioxidants was tested. Changes in process yield, purge loss, texture, color, microbial growth and pH during vacuum refrigerated storage were monitored. Partial substitution of sodium did not affect matrix stability, maintaining high process yields and low purge losses (≤5.5%). The products with marine oil used as fat source resulted in: high PUFA levels and lower risks indicators associated with cardiovascular events. Tocopherols prevented the oxidation process; n-6/n-3 ratio remained unchanged throughout the storage, establishing a natural alternative to BHA. Moreover, the consumption of 15–18 g of this product would cover the recommended daily intake of EPA + DHA.

PRACTICAL APPLICATIONS

In previous works, we developed formulations replacing the beef fat with pre-emulsified and deodorized marine oil. We also study an alternative formulation with low sodium content. These characteristics are a necessity for the consumers who are demanding better nutritional quality products, and the producers must attend that demand. Other authors have studied different low fat and/or low sodium meat systems or meat emulsions with different fat sources to enhance the nutritional quality. Nevertheless there is not much knowledge of the stability of these new meat systems, containing more water, and more PUFA. Thus, the aim of this research was to study the storage stability of different cooked meat sausages with fish oil from different approaches (microbial, physicochemical and oxidative). Assuring the stability of these products is essential to the producers to maximize the shelf-life.

INTRODUCTION

Meat products reformulation is one of the strategies that have been studied in order to develop meat-based functional foods, generally based on animal fat replacement with other lipids such as plant and/or marine oils (Berasategi et al. 2014). The high polyunsaturated fatty acids (PUFA) present in marine oils have numerous beneficial health effects associated with its consumption (Funahashi et al. 2006; Coates et al. 2009), particularly of eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA). WHO and USDA (WHO 2008; USDA 2010) recommend a dairy intake of 250 mg of long-chain n-3 PUFA in persons with and without cardiovascular diseases.

Muscle foods are susceptible to oxidation. Meat processing operations that increase surface area, addition of potential
PUFA, and heat treatments decrease oxidative stability (Lee et al. 2005). The use of antioxidants could prevent the oxidative spoilage of n-3 PUFA enriched foods, however, similar antioxidants show different effects on same food matrix (Jacobsen et al. 2008). Previous works have shown that it is possible to develop new PUFA enriched meat products with preemulsified oils and antioxidants to improve its nutritional properties (Asuming-Bediako et al. 2014; Berasategi et al. 2014; Marchetti et al. 2015).

Synthetic antioxidants (BHT, BHA, PG, TBHQ) are widely used in food industry. However, in several studies they have been related with tumors development and other negative effects (Amadasi et al. 2008; Gharavi and El-Kadi 2005). Nowadays consumers encourage food manufacture from natural sources and with the so-called green technologies (Valenzuela et al. 2011). Natural antioxidants are polyphenolic compounds that can be found in herbs, spices and other vegetables. In 2013, the World Health Assembly (WHO 2013) agreed nine global voluntary targets for the prevention and control of Noncommunicable Diseases, which include a 30% relative reduction in the intake of salt by 2025. According to WHO (2013) reducing salt intake has been identified as one of the most cost-effective measures that countries can take to improve population health outcomes. However, in meat products, NaCl promotes the solubilization of myofibrillar proteins, increasing the hydration and water retention capacity, thus reducing cooking and exudate losses. If the NaCl content of the formulation is reduced, it might adversely affect such properties. Potassium chloride (KCl) is the most commonly used substitute in low/reduced sodium foods. Feltrin et al. (2015) reported that KCl was the only salt replacer that showed temporal sensory profile similar to NaCl. However, at blends over 50:50 potassium chloride/sodium chloride in solution, a significant increase in bitterness and loss of saltiness was observed (Desmond 2006; Soglia et al. 2014). Both, fat and salt play an important role in this product so alternatives must be carefully chosen to reduce both components.

Cooling, vacuum packaging and edible coating are common techniques to maintain the quality of agri-food products. For cooked meats, cooling is also a very important process to ensure product safety before consumption (Feng and Sun 2013). During vacuum refrigerated storage of cooked meat emulsions changes in their quality parameters (weight loss, texture, color, microbial growth and fatty acid profile) that may limit shelf-life may occur. Andrés et al. (2009) found that low-fat chicken sausages containing squid oil with synthetic vitamin E had good stability and quality attributes during the storage. In a previous work we studied low-fat meat emulsions with preemulsified fish oil with different hydrocolloids added, optimized the carrageenan and milk proteins levels (Marchetti et al. 2014), and then optimized the formulation in order to reduce its sodium content (Marchetti et al. 2014, 2015). Although they contained 46 and 71% less sodium than a commercial sausage, both formulations presented good sensory scores, but it is still necessary to study their storage stability, particularly the inhibition of lipid oxidation that keep n-3 PUFA unaltered, the possibility of larger exudates values when less Na is present in the system.

In the present paper, the objective was to study changes in physicochemical characteristics (purge loss, color, textural), microbial counts and pH during 45 days of vacuum refrigerated storage (4C) of two low-fat sausage formulations, where deodorized marine oil has been used for replacing saturated animal fat, and containing milk protein concentrate and k/i carrageenans. In one of the formulations a partial NaCl replacement with KCl and sodium tripolyphosphate (TPP) was carried out. The experimental design included different levels of natural tocopherols or BHA to prevent lipid oxidation and assure an adequate shelf-life. Changes in their fatty acids (FA) profile and lipid oxidation were also studied, and its effect on different health related indexes.

MATERIALS AND METHODS

Materials

Low-fat sausages were prepared using fresh lean beef meat (adductor femoris and semimembranosus muscles) obtained from local market (pH: 5.48 ± 0.01, fat content: 13 ± 1.7 g/kg). Meat (18 kg, from eight different carcasses for each batch of experiments) without visible fat and connective tissue was passed through a grinder with a 0.95 cm plate (Meifa 32, Buenos Aires, Argentina). Thirty-six lots of 500 g was vacuum packed in Cryovac BB4L bags (PO2: 0.35 cm3/m2/d/kPa at 23C, Sealed Air Co., Buenos Aires, Argentina), frozen and stored at −20C until used (no more than 3 weeks).

Fat source was commercial deodorized marine oil (Omega Sur S.A., Mar del Plata, Argentina). As stabilizer or emulsifier agents food-grade commercial preparations of milk proteins concentrate (802 g/kg proteins (casein-s + whey proteins, solubility 97.3 ± 0.4%; Milkaut, Santa Fe, Argentina) and synergistic 2:1 k/i carrageenans mixture (ADAMA S.A., Buenos Aires, Argentina) were used (Marchetti et al. 2014). Cold distilled water was used in all formulations (4C). Mixed phytosterols (Advasterol 90, AOM S.A., Buenos Aires, Argentina) were included. Analytical grade sodium chloride (NaCl), nitrite (NaNO2), erythorbate and tripolyphosphate (TPP) salts were employed. Sodium nitrite concentration was selected according to the level permitted by Argentinean food law (0.15 g/kg, Código Alimentario Argentino (1999)).

The following components were included to prepare 1 kg of uncooked meat batter: meat (666.5 g), water (250 g), deodorized
The experiment included two different salts combinations levels, which corresponded to the optimized systems studied by Marchetti et al. (2014, 2015). Formulations were codified as Na (14.00 g NaCl + 2.00 g TPP/kg), and partially NaCl replaced (Na/K: 6.08 g NaCl/kg + 4.92 KCl g/kg + 5.00 g TPP/kg). In any case, the total amount of these salts was 16.00 g/kg, a content lower than traditional products (Desmond 2006).

Two levels of natural tocopherols (T, Tocomin 70, AOM SA, Buenos Aires, Argentina, with d-γ-/d-β-tocopherol 43.81%, d-δ-tocopherol 19.31% and d-α-tocopherol 7.40%) were evaluated. Formulations without antioxidants were included as controls for both salt combinations (Table 1). One formulation of Na sausages with butylated hydroxyanisole (BHA, Fagron S.A., Madrid, Spain) at maximum permitted level (0.5 mg/100 g product, Cypres 2016) was also included in the design. The sample size of each formulation was 80–100 links (28–33 g per sausage) and the study was run in duplicate.

### Microbial Analysis

Bacterial counts were determined using the pour plate method at different times during refrigerated storage according to Andrés et al. (2009). The methodology of Brennan and Bourne (1994) was followed to determined Texture Profile Analysis (TPA), analyzing 10 replicates per point. Color was determined at room temperature on the surface of transversally slices, recently cut, according to Marchetti et al. (2015). Five measures were taken for each data point. Finally, pH of the samples was measuring in triplicate using a spear tip glass electrode with Ag/AgCl reference (Phoenix 557-3512, AZ) on a pHmeter (EC30, Hacht, Loveland, CO).

### Physicochemical Determinations

Process yield and purge loss were performed by triplicate according to Andrés et al. (2009). The methodology of Brennan and Bourne (1994) was followed to determined Texture Profile Analysis (TPA), analyzing 10 replicates per point. Color was determined at room temperature on the surface of transversally slices, recently cut, according to Marchetti et al. (2015). Five measures were taken for each data point. Finally, pH of the samples was measuring in triplicate using a spear tip glass electrode with Ag/AgCl reference (Phoenix 557-3512, AZ) on a pHmeter (EC30, Hacht, Loveland, CO).

### Product Manufacture

Elaboration of the sausages was according to Marchetti et al. (2014, 2015). Briefly, 500 g grounded meat was homogenized in a commercial food processor (Universo, Rowenta, Germany, 14 cm blade) with Na or Na/K mixture according to the design (Table 1). Carrageenans, milk proteins, sodium nitrite and erythorbate were dissolved in cold water and then homogenized with the deodorized marine oil using a hand-held food processor (Braun, Buenos Aires, Argentina) during 2 min to form a coarse emulsion. The obtained emulsion was added to ground meat, processing all ingredients during 5 min afterward. Final temperature of batter varied between 12 and 15C. Samples were stuffed (vertical piston stuffer, Santini s.n.c., Marostica, Italy; into cellulose casing 22 mm diameter, Farmesa, Buenos Aires, Argentina), thermally treated in a hot water bath (80C) until the center reached 74C, cooled, vacuum packaged in Cryovac BB4L bags and stored at 4C during 45 days (typical shelf-life of commercial products).

| Code   | Sodium chloride (g/kg) | Potassium chloride (g/kg) | Sodium tripolyphosphate (g/kg) | Tocopherols (mg/kg) | BHA (mg/kg) |
|--------|------------------------|---------------------------|-------------------------------|---------------------|-------------|
| Na-C   | 14.00                  | –                         | 2.00                          | –                   | –           |
| Na-BHA | 14.00                  | –                         | 2.00                          | –                   | –           |
| Na-T1  | 14.00                  | –                         | 2.00                          | 37.5                | –           |
| Na-T2  | 14.00                  | –                         | 2.00                          | 50.0                | –           |
| Na/K-C | 6.08                   | 4.92                      | 5.00                          | –                   | –           |
| Na/K-T1| 6.08                   | 4.92                      | 5.00                          | 37.5                | –           |
| Na/K-T2| 6.08                   | 4.92                      | 5.00                          | 50.0                | –           |

* Units are expressed per kg of raw meat batter.

Codes: Na = sodium formulations, Na/K = partial Na replaced formulations, C = control without antioxidant, BHA = butylated hydroxyanisole, T1-2 = tocopherols levels.
were also tested for total coliform counts using the most probable number method (MPN) according to AOAC (AOAC 1984) method 46016, and sulfite-reducing Clostridium were enumerated in Tryptone Sulfite Neomycin Agar (TNS agar, Oxoid) (incubated at 30°C for 2 d). Data were expressed as log colony-forming units per gram of sample.

**Lipid Oxidation and Fatty Acids Profile Determination**

TBARS values were determined by quadruplicate according to Pennisi Forell et al. (2010) to evaluate the lipid oxidation in the sausages. Results were expressed as mg malonaldehyde (MDA)/kg product.

For fatty acid (FA) analysis of Na-T2, Na/K-T2, Na-BHA and Na/K-C formulations at initial and final storage time, total lipids were extracted using chloroform-methanol mix (2:1, v/v) according to Folch et al. (1957) procedure, and were methylated with 100 g/kg boron trifluoride methanol complex in methanolic solution. FA composition was determined at the Instituto Nacional de Investigación y Desarrollo Pesquero (INIDEP, Mar del Plata), following (Pennisi Forell et al., 2010), in a Shimadzu 2010 gas chromatograph (Hewlett-Packard, Palo Alto, CA) equipped with capillary column Omegawax 320 (30 m/0.32 mm id/0.25 μm) and mass detector. FA profiles were obtained by comparison of the retention times with a standard of 37 fatty acids (Supelco 37 Component FAME Mix, Cat. No. 18919-1 AMP, Sigma-Aldrich) previously analyzed in same conditions. Fatty acids were identified by comparison of the retention times.

**Changes in Health Lipid Indexes During Storage**

Based on the FA results the atherogenic index (AI, Eq. (1)) and the thrombogenic index (TI, Eq. (2)) were calculated according to Ulbricht and Southgate (1991) to assess the nutritional quality of the products, as a measure of the propensity of the product consumption influence the incidence of coronary heart disease:

\[
AI = \frac{C_{12:0} + 4 \times C_{14:0} + C_{16:0}}{[\text{PUFA}_{n-6} + \text{PUFA}_{n-3} + \text{MUFA}]} \tag{1}
\]

\[
TI = \frac{\left[\frac{1}{2} \text{PUFA}_{n-6} + \frac{3}{2} \text{PUFA}_{n-3} + \frac{1}{2} \text{MUFA} + \frac{1}{2} \frac{\text{PUFA}_{n-3}}{\text{PUFA}_{n-6}}\right]}{\frac{C_{12:0} + C_{14:0} + C_{16:0}}{2}} \tag{2}
\]

where \(C_{n:0}\) corresponds to each fatty acid content expressed as % FA.

Also the nutritional fat index (NFI = PUFA + MUFA)/SFA) was calculated (Amine et al. 2002).

**Statistical Analysis**

Analysis of variance (ANOVA, SYSTAT, Inc., Evanston, IL) was carried out to test the significance of independent variables. Experimental data were reported as mean values ± the corresponding standard error of the mean (SEM) when appropriate. For simultaneous pairwise comparisons, least significance differences (LSD) test was chosen. Differences in means and F tests were considered significant when \(P < 0.05\).

**RESULTS AND DISCUSSION**

**Physicochemical Properties**

Process yield was not affected by salt contents or antioxidants. Formulations exhibited an average value of 985 ± 3 g/kg (\(P > 0.05\)), indicating high liquid retention of the matrix during the thermal treatment, even in Na/K formulations. These results were in agreement with Triki et al. (2013), who found no differences in process yields between merguez sausage formulation with 50% of NaCl replacement. Purge losses could be a serious problem, besides the fact of an unpleasant aspect of the product, by stimulating the microbial growth resulting in a lower shelf-life (López-López et al. 2009). Purge loss varied between 12 ± 1 g/kg at the beginning of storage for both Na content, and 43 ± 2 or 53 ± 2 g/kg for Na or Na/K formulations, respectively, for the final storage time (Fig. 1). These values were similar to those reported by other authors for lean sausages (Candogan and Kolsarici 2003; Andrés et al. 2009). Sodium reduced and nonreduced formulations showed different behavior (Fig. 1). Up to 14 days, purge loss exhibited a sharp increase and no significant differences among formulations. After 20 days, the effect of sodium replacement becomes significant. Those formulations with KCl added, released more liquid than the formulations without Na replacement that remained fairly constant. Low NaCl level could decrease the concentration of extracted/solubilized proteins involved in the formation of the emulsified gel. Low purge losses could be related with high liquid retention by the matrix throughout the storage, which was not modified by the antioxidant added to the product. Similar results have been reported by Colmenero et al. (2005), who studied the effect of NaCl, KCl, and transglutaminase in low-fat sausages formulations and found that the partial NaCl replacement decreased water binding properties.

Texture profile could reflect the possible changes that if noticed by the consumers may impact in their acceptance of the product. In Fig. 2, the obtained results of textural parameters, hardness, chewiness and resilience of formulated sausages during refrigerated storage are showed. Hardness was significantly affected by sausage formulation and
storage time (Fig. 2a). Initial hardness of reduced sodium sausages (Na/K) was lower than nonreplaced ones. Literature shows diverse texture results depending on meat system, type and salt level used as NaCl partial replacer. Horita et al. (2011) found similar variations in emulsified meat products texture when NaCl was 50% reduced, with a hardness decrease when NaCl was reduced up to 75%. Besides, Marchetti et al. (2015) working with sodium-reduced lean sausages with fish oil found that for a given KCl level, hardness increased with TPP fraction, probably because changes in ionic strength and protein solubility affected meat texture.

Both sets of formulations increased its hardness with storage time, and after 30 days, no significant differences between formulations were observed; thus, there was a marked hardness increase when potassium chloride and TPP were added (28.3%) with respect to formulations without KCl (11.1%). This could be explained by the differences observed in purge loss, partially replaced sodium sausages (Na/K) lost more liquid and increased their hardness more rapidly than the nonreplaced formulations (Na), resulting in less water available to act as matrix plasticizer. Therefore, a possible relationship between hardness and purge loss was investigated (Fig. 3), finding a significant correlation ($P < 0.05$) between both parameters for each salt mixture (sodium-replaced and nonreplaced). Nevertheless, hardness values (9–12 N) were similar to those measured for Argentinian commercial products containing 20% fat. These results agree with other authors who had informed increases in hardness during refrigerated storage of cooked meat emulsions (Estévez et al. 2005; Hassaballa et al. 2009).

Chewiness showed a similar tendency to hardness (Fig. 2b). On the other hand, cohesiveness and springiness were not significantly altered by storage time or formulation. The obtained mean values were 0.873 ± 0.007 (mm/mm) for springiness and 0.573 ± 0.004 for cohesiveness (J/J).

Color is one of the main factors that affect the acceptability of a meat product by consumers. Chromaticity parameters ($a^*$ and $b^*$) showed neither changes during storage nor between formulations ($P > 0.05$); the obtained mean values for $a^*$ and $b^*$ were approximately 0 and 15, respectively.

FIG. 1. PURGE LOSS (g/kg) OF MEAT SAUSAGES FORMULATED WITH MARINE OIL DURING REFRIGERATED VACUUM STORAGE

Codes: (1) Na formulations (14.00 g NaCl + 2.00 g TPP/kg product): (●), 37.5 g tocopherols/kg (Na-T1); (●), 50 g tocopherols/kg (Na-T2); (▲), 5 g BHA/kg (Na-BHA); (■), control without antioxidant (Na-C); (2) Na/K formulations (6.08 g NaCl + 4.92 g KCl + 5.00 g TPP/kg product): (○), 37.5 g tocopherols/kg (Na-T2); (○), 50 g tocopherols/kg (Na-T2); (□), control without antioxidant (Na/K-C). Error bars indicate SEM.

FIG. 2. EFFECT OF REFRIGERATED VACUUM STORAGE TIME ON TEXTURE PROFILE ANALYSIS PARAMETERS OF MEAT SAUSAGES FORMULATED WITH MARINE OIL

(a) Hardness, (b) chewiness. Codes: (1) Na formulations (14.00 g NaCl + 2.00 g TPP/kg product): (●), 37.5 g tocopherols/kg (Na-T1); (●), 50 g tocopherols/kg (Na-T2); (▲), 5 g BHA/kg (Na-BHA); (■), control without antioxidant (Na-C); (2) Na/K formulations (6.08 g NaCl + 4.92 g KCl + 5.00 g TPP/kg product): (○), 37.5 g tocopherols/kg (Na-T1); (○), 50 g tocopherols/kg (Na-T2); (□), control without antioxidant (Na/K-C). Error bars indicate SEM.
Luminosity (L*) changes microbiological traits (total aerobic mesophilic and psychrotrophic and LAB) in vacuum-packaged rabbit meat. Dominant flora in these products was psychrotrophic lactic acid bacteria, in concordance with other authors (Nychas and Drosinos 1999; Andrés et al. 2009). This spoilage might significantly affect product quality due to the acidification in anaerobic conditions. Table 2 shows average microbial counts of the formulations analyzed at different storage time. All formulations presented low initial microbial counts for total mesophilic and psychrotrophic microorganisms, and lactic acid bacteria (LAB), in consequence of the adequate thermal treatment done in their production. At the end of storage, total mesophilic levels were lower than 5 log cfu/g, maximum level permitted by Argentinean regulations (Código Alimentario Argentino 1999). No lag phase was observed for the microbial growths for mesophilic, psychrotrophic and LAB, Feng et al. (2014) reported similar trends in refrigerated Irish sausages. Regarding the pH evolution of the samples during storage pH decreased from 5.82 to 5.34 between initial and final time, related to LAB development (Table 2). Cayre et al. (2005) proposed that the vacuum storage of meat products limited the growth of Pseudomonas spp., resulting in lactic acid bacteria as the main component of the flora. In these products, its development and metabolism depend of different factor (pH, temperature, atmospheric composition within package, substrate availability) (Yan et al. 2008).

Enterobacteriaceae and yeast and molds counts were below the detection limit of the technique (2 log cfu/g) during the refrigerated storage of all the analyzed formulations. Total coliforms counts were <2 MPN/g in all formulations at the end of storage. These results were in accordance to Argentinean regulations (Código Alimentario Argentino 1999). In addition, no sulfite-reducing Clostridium was noted in the sausages during the storage period, indicating safe sanitary conditions, and related to the inclusion of NaNO2, which is a key component to avoid Clostridium spp. growth (Christiansen et al. 1975).

![FIG. 3. CORRELATION BETWEEN HARDNESS AND PURGE LOSS](Image)

Codes: (1) Na formulations (14.00 g NaCl + 2.00 g TPP/kg product); (●), 37.5 g tocopherols/kg (Na-T1); (●●), 50 g tocopherols/kg (Na-T2); (▲), 5 g BHA/kg (Na-BHA); (■) control without antioxidant (Na-C); (2) Na/K formulations (6.08 g NaCl + 4.92 g KCl + 5.00 g TPP/kg product); (○), 37.5 g tocopherols/kg (Na-T1); (○○), 50 g tocopherols/kg (Na-T2); (□) control without antioxidant (Na/K-C). Error bars indicate SEM.

were 10.3 ± 0.9 and 13.3 ± 0.8 for a* and b*, respectively. These color parameters result in agreement with those reported by García-García and Totosaus (2008). However, luminosity of all formulations significantly decreased after 20 days of storage (P < 0.05), as shown in Table 2. These changes could be related to the higher solid content of the product as a result of liquid lost as purge.

### Microbial Quality

Sodium reduction did not significantly affect the microbial growth, because KCl has shown similar antimicrobial effect than NaCl (Bidlas and Lambert 2008). Soglia et al. (2014) informed that replacing up to 30% of NaCl by KCl did not change microbiological traits (total aerobic mesophilic and lactic LAB counts) in vacuum-packaged rabbit meat. Dominant flora in these products was psychrotrophic lactic acid bacteria, in concordance with other authors (Nychas and Drosinos 1999; Andrés et al. 2009). This spoilage might significantly affect product quality due to the acidification in anaerobic conditions. Table 2 shows average microbial counts of the formulations analyzed at different storage time. All formulations presented low initial microbial counts for total mesophilic and psychrotrophic microorganisms, and lactic acid bacteria (LAB), in consequence of the adequate thermal treatment done in their production. At the end of storage, total mesophilic levels were lower than 5 log cfu/g, maximum level permitted by Argentinean regulations (Código Alimentario Argentino 1999). No lag phase was observed for the microbial growths for mesophilic, psychrotrophic and LAB, Feng et al. (2014) reported similar trends in refrigerated Irish sausages. Regarding the pH evolution of the samples during storage pH decreased from 5.82 to 5.34 between initial and final time, related to LAB development (Table 2). Cayre et al. (2005) proposed that the vacuum storage of meat products limited the growth of *Pseudomonas* spp., resulting in lactic acid bacteria as the main component of the flora. In these products, its development and metabolism depend of different factor (pH, temperature, atmospheric composition within package, substrate availability) (Yan et al. 2008).

Enterobacteriaceae and yeast and molds counts were below the detection limit of the technique (2 log cfu/g) during the refrigerated storage of all the analyzed formulations. Total coliforms counts were <2 MPN/g in all formulations at the end of storage. These results were in accordance to Argentinean regulations (Código Alimentario Argentino 1999). In addition, no sulfite-reducing *Clostridium* was noted in the sausages during the storage period, indicating safe sanitary conditions, and related to the inclusion of NaNO2, which is a key component to avoid *Clostridium* spp. growth (Christiansen et al. 1975).

### Table 2. Changes in Average Luminosity (L*), pH and Microbial Counts During Refrigerated Storage of Low-Fat Meat Emulsions Prepared with Marine Oil

| Time (days) | Luminosity (L*) | pH | Total mesophilic counts (log cfu/g) | Total psychrotrophic counts (log cfu/g) | Lactic acid bacteria (log cfu/g) |
|------------|----------------|----|-------------------------------|--------------------------------------|---------------------------------|
| 1          | 61.8 ± 0.2a    | 5.82 ± 0.01a | 2.98 ± 0.08e | 1.87 ± 0.05g | 2.03 ± 0.1f |
| 7          | 61.5 ± 0.3ab   | 5.79 ± 0.02ab | 3.33 ± 0.07e | 2.22 ± 0.3f | 2.44 ± 0.2e |
| 14         | 60.9 ± 0.2b    | 5.74 ± 0.02bc | 3.70 ± 0.1d | 2.61 ± 0.07e | 2.81 ± 0.09e |
| 22         | 60.5 ± 0.2bc   | 5.69 ± 0.01d | 3.98 ± 0.06dc | 2.99 ± 0.08d | 3.27 ± 0.3d |
| 28         | 60.1 ± 0.2cd   | 5.61 ± 0.01d | 4.26 ± 0.1bc | 3.10 ± 0.1cd | 3.4 ± 0.07cd |
| 34         | 60.0 ± 0.3cde  | 5.51 ± 0.03e | 4.51 ± 0.1ab | 3.45 ± 0.1bc | 3.71 ± 0.2bc |
| 41         | 59.9 ± 0.1de   | 5.42 ± 0.02f | 4.69 ± 0.2a | 3.58 ± 0.08ab | 3.89 ± 0.1b |
| 45         | 59.6 ± 0.2e    | 5.34 ± 0.01g | 4.78 ± 0.08a | 3.88 ± 0.03a | 4.29 ± 0.9a |

Average values ± standard error of the mean (SEM), different superscripts within the same column indicate that average values differ significantly (P < 0.05).
TBARS for meat systems with fat sources composed of an O/W emulsion with algae oil (Crypthecodinium cohnii). Several physicochemical or sensory TBARS limits in meat products or systems have been reported. Campo et al. (2006) informed that levels > 2 mg MDA/kg are not accepted in bovine meat. Otherwise, Georgantas et al. (2007) established a maximum limit of 0.6 mg MDA/kg over which it is detectable a rancid flavor in meat products. Therefore, according to the obtained results formulations with natural tocopherols or BHA presented TBARS values lower than even the strictest limits suggested in the literature during the 45 days of storage. However, it was necessary to add at least 37.5 and 50 mg tocopherols/kg to Na and Na/K formulations, respectively, to achieve the inhibition obtained with BHA in sausages containing 14 Na/kg.

These results agree with those reported by Kim (2012) who obtained a reduction of TBARS and improved color stability of a meat emulsion system by using 67 and 134 mg tocopherols/kg product. Also it has been reported that the addition of 50 and 100 mg tocopherols/kg to stuffed cooked meat product reduced the peroxide value, free fatty acids and TBARS number (Aksu 2007). Cáceres et al. (2008) reported low lipid oxidation (TBARS 0.37–0.52 mg MDA/kg) during cooling of bologna made with commercial fish oil with α-tocopherol, resulting in similar values to those obtained in this work.

### Fatty Acid Profile

The results of fatty acid composition are consistent with the type of ingredients used in the formulation. Table 3 shows the obtained fatty acids profiles from the lipid phases of several formulations (sodium reduced or not) made with marine oil with different antioxidants (BHA or tocopherols) at the initial and end (45 days) of the storage period. In addition, it was included a FA profile of a reduced sodium formulation without antioxidants (control) and a traditional product with animal fat (USDA 2015).

The obtained FA profiles are within the current diet recommendations, due to marine oil incorporation. In addition to considerations of individual fatty acids, scientific evidence suggests that ratios such as PUFA/SFA (recommended > 0.4) and n-6/n-3 PUFA (recommended < 4) are the main parameters currently used to assess the nutritional quality of the lipid fraction of foods. In 45 g (1 commercial sausage link) of the products studied in this work, saturated or 50 mg tocopherols/kg (Na-T2 and Na/K-T2), with a slight increase in TBARS at the end of the storage (< 0.4 mg MDA/kg product), without significant differences between both antioxidants (P > 0.05). This implies an adequate inhibition of lipid oxidation in the studied meat systems, showing that the synthetic antioxidant could be replaced with a natural one with similar results.
(SFA) and monounsaturated (MUFA) fatty acids were lower than those corresponding to a traditional formulation (659 mg versus 4219 mg, and 959 mg versus 3939 mg, respectively). In addition, one serving (45 g) of low-fat sausages with marine oil contained 820 mg PUFA, providing 241 mg of EPA and 419 mg of DHA, contrasting with the traditional product with pork fat, which presents 313 mg of PUFA per 45 g sausage (USDA 2015), with no EPA or DHA.

The FA profile of the reformulated products results in a significantly lower n-6/n-3 ratio. Furthermore, the PUFA/SFA ratio was always >1.2, thus replacement of pork or beef fat by marine oil with antioxidants, significantly increased this ratio from the commonly found for these products (Delgado-Pando et al. 2011) (about 0.34, Table 3).

EFSA dietary recommendations (EFSA 2012) for EPA and DHA based on cardiovascular diseases risk considerations for adults are between 250 and 500 mg/d. This product could easily sum up for the daily intake of EPA and DHA; an intake of one serving of this product would greatly exceed the minimum 250 mg required.

The formulation without antioxidant (Na/K-C) showed a noteworthy decrease (P < 0.05) of EPA, DHA, and total PUFA (21.3, 26.6 and 27.7% reduction, respectively), also, in oleic, linoleic and linolenic acid contents at 45 days of storage. With the antioxidants addition, the oxidation of the last fatty acids was inhibited, while EPA and DHA oxidation was reduced. The n-6/n-3 ratio of the products remained unchanged throughout the storage period (range: 0.09–0.16).

FA profiles and their changes at the end of vacuum-packaged refrigerated storage are in agreement with the results obtained in the TBARS assay, where inclusion of tocopherols in the formulation were able to delay lipid oxidation, establishing a natural alternative to BHA.

Average values of AI and TI for sausages manufactured with marine oil were 0.40 and 0.17, respectively, significantly lower than the traditional product indexes, in agreement with the literature reports (Ulbricht and Southgate 1991; Higgs 2000; Senso et al. 2007; Afonso et al. 2013), indicating less risk of cardiovascular event. Moreover, all cooked sausages achieved the World Health Organization’s recommendation (Amine et al. 2002) on the nutritional fat index ((NFI = PUFA + MUFA)/SFA ≥ 2) which is very relevant to the development of healthier formulations since the calculated values ranged between 2.26 and 3.15. Besides three indexes remained unchanged during storage when antioxidants were added (formulations Na-BHA, Na-T2 and Na/K-T2).

In previous works sensory assays showed that neither the deodorized fish oil inclusion nor the partial substitution of

### TABLE 3. FATTY ACID (FA) PROFILES OF DIFFERENT SAUSAGES FORMULATED WITH MARINE OIL AT INITIAL OR END OF STORAGE. TP DENOTES A TRADITIONAL PRODUCT ACCORDING TO USDA (2015)

| Fatty acid (% of total FA) | (14 g NaCl + 2 g TPP)/kg | (6.08 g NaCl + 4.92 g KCl + 5 g TPP)/kg |
|---------------------------|--------------------------|----------------------------------------|
|                           | Na-BHA (5 mg BHA/kg)    | Na-T2 (50 mg T/kg)                    | Na/K-C (no antioxidant) | Na/K-T2 (50 mg T/kg) |
|                           | 0 days | 45 days | 0 days | 45 days | 0 days | 45 days | 0 days | 45 days | 0 days | 45 days |
| Lauric C12:0              | N.D.   | N.D.    | N.D.   | N.D.    | N.D.   | N.D.    | N.D.   | N.D.    | N.D.   | 2.7     |
| Myristic C14:0            | 3.8b   | 4.2ab   | 4.0b   | 4.0b    | 0.9c   | 1.1c    | 1.0c   | 1.1c    | 1.0c   | 4.4     |
| Palmitic C16:0            | 16.9d  | 18.1b   | 17.0c  | 17.2cd  | 17.5c  | 18.6b   | 15.9e  | 16.3e   | 15.9e  | 20.6    |
| Palmitoleic C16:1 n-7     | 5.2c   | 5.5c    | 5.2c   | 5.2c    | 7a     | 6.4b    | 7.3a   | 6.8ab   | 6.0b   | 4.8     |
| Stearic C18:0             | 4.8c   | 4.8c    | 4.8c   | 4.9c    | 6.1b   | 5.9b    | 6.0b   | 6.2b    | 6.0b   | 22.1    |
| Oleic C18:1 n-9 cis       | 27.1c  | 27.2c   | 26.6cd | 27.1c   | 28.6e  | 26.1de  | 28.6b  | 26.1de  | 28.6b  | 41.1    |
| Linoleic C18:2 n-6        | 2.7b   | 2.8b    | 2.8b   | 2.8b    | 2.5b   | 1.6c    | 2.5b   | 2.5b    | 2.5b   | 3.3     |
| Linolenic C18:3 n-3       | 2.1a   | 2.1a    | 2.2a   | 2.1a    | 2.4a   | 0.7b    | 2.4a   | 2.2a    | 2.4a   | 0.4     |
| C20:1 (undefined)         | 5.2a   | 5.2a    | 5.3a   | 5.2a    | 3.5b   | 3.4b    | 3.4b   | 3.6b    | 3.4b   | 0.6     |
| Arachidonic C20:4 n-6     | 1.6a   | 1.6a    | 1.6a   | 1.4a    | 1.4a   | 0.7b    | 1.5a   | 1.4a    | N.D.   |         |
| Eicosapentaenoic C20:5 n-3| 10.9a  | 9.9b    | 10.8a  | 9.8b    | 8.9c   | 7.0e    | 8.8cd  | 8.4d    | N.D.   |         |
| Docosahexaenoic C22:6 n-3 | 17b    | 16.2c   | 16.9b  | 16.2c   | 17.7a  | 13.0a   | 17.6a  | 16.7b   | N.D.   |         |
| SFA                       | 25.5c  | 27.1b   | 25.8c  | 26.1bc  | 24.5c  | 25.6bc  | 22.9c  | 23.6c   | 24.5c  | 49.8    |
| MUFA                      | 37.5c  | 37.9bc  | 37.1c  | 37.5bc  | 36.3cd | 34.1d   | 39.3b  | 36.5c   | 36.3cd | 46.5    |
| PUFA                      | 34.3a  | 32.4ab  | 34.3a  | 32.1bc  | 32.9ab | 23.8d   | 32.8ab | 31.2c   | 32.8ab | 3.7     |
| n-6/n-3                   | 0.14b  | 0.09b   | 0.10b  | 0.09b   | 0.15b  | 0.12b   | 0.15b  | 0.16b   | 0.15b  | 8.33    |
| NFI                       | 2.82a  | 2.59ab  | 2.77a  | 2.67ab  | 2.82a  | 2.26b   | 2.15a  | 2.87a   | 2.15a  | 1.01    |
| PUFA/SFA                  | 1.35ab | 1.20c   | 1.33a  | 1.23c   | 1.34b  | 0.93d   | 1.43a  | 1.32b   | 0.70   | 0.07    |
| Aterogenicity Index       | 0.45b  | 0.50b   | 0.46b  | 0.48b   | 0.30c  | 0.40b   | 0.28c  | 0.31c   | 0.31c  | 0.81    |
| Trombogenicity Index      | 0.18b  | 0.19b   | 0.17b  | 0.18b   | 0.16b  | 0.22b   | 0.15b  | 0.16b   | 0.16b  | 1.06    |

N.D. = Not detected. Different superscripts within the same row f indicate that average values differ significantly (P < 0.05).
NaCl had a negative impact over the flavor, color, texture and overall acceptability (Marchetti et al. 2014, 2015). It may be concluded that these products would present good storage stability if natural tocopherols were added in at least 50 mg/kg.

CONCLUSIONS

A significant reduction of sodium content did not alter process high yields (985 g/kg) and low purge losses (≤5.5%). Reducing Na content initially produced harder sausages, but hardness increased during storage at a different rate that depended on Na content, reaching similar values at the end of the 45 days period, within the commercial products hardness range. Sodium replacement significantly affected the oxidative stability of the products, although 50 mg natural tocopherol/kg successfully prevented rancidity in products with and without NaCl partial replacement. The resulting fatty acid profile was associated with a reduction in risks of different cardiovascular diseases (lower TI and AI).

Thus, it is possible to obtain cooked meat emulsions (sausages) with low sodium, low saturated fat, and high amounts of n-3 PUFA by applying a combination of carrageenans, milk proteins concentrate and preemulsified marine oil, without significant adverse effects over the quality of the products for at least 45 days of refrigerated storage.

ACKNOWLEDGMENTS

This research was supported by Consejo Nacional de Investigaciones Científicas y Tecnológicas (CONICET, Argentina), Agencia Nacional de Promoción Científica y Tecnológica (Argentina) and Universidade Nacional de La Plata. The authors thanks Dr. Carolina Pennisi Forell (Instituto Nacional de Investigación y Desarrollo Pesquero, INIDEP, Mar del Plata, Argentina) for the fatty acid profile analysis, and Milkaút S.A., Arla Foods Ingredients S.A., Omega Sur S.A., and AOM S.A. for the materials supplied.

REFERENCES

AFONSO, C., CARDOSO, C., LOURENÇO, H., ANACLETO, P., BANDARRA, N., CARVALHO, M., CASTRO, M. and NUNES, M. 2013. Evaluation of hazards and benefits associated with the consumption of six fish species from the Portuguese coast. J. Food Compos. Anal. 32, 59–67.
AKSU, M.I. 2007. The effect of α-tocopherol, storage time and storage temperature on peroxide value, free fatty acids and pH of kavurma, a cooked meat product. J. Muscle Foods 18, 370–379.
AMADASI, A., MOZZARELLI, A., MEDA, C., MAGGI, A. and COZZINI, P. 2008. Identification of xenoestrogens in food additives by an integrated in silico and in vitro approach. Chem. Res. Toxicol. 22, 52–63.
AMINE, E., Baba, N., Belhadj, M., Deurenbery-Yap, M., Diazayery, A., Forrester, T., Galuska, D., Herman, S., James, W. and Mbuyamba, J. 2002. Diet, Nutrition and the Prevention of Chronic Diseases: Report of a Joint WHO/FAO Expert Consultation, World Health Organization, Geneva, Switzerland.
ANDRÉS, S.C., Zaritzky, N. and Califano, A. 2009. Innovations in the development of healthier chicken sausages formulated with different lipid sources. Poult. Sci. 88, 1755–1764.
AOAC. 1984. Official Methods of Analysis, 14th ed., Assoc. of Official Analytical Chemists, Washington, D.C.
ASUMING-BEDIAKO, N., Jaspal, M., Hallett, K., Bayntun, J., Baker, A. and Sheard, P. 2014. Effects of replacing pork backfat with emulsified vegetable oil on fatty acid composition and quality of UK-style sausages. Meat Sci. 96, 187–194.
Berasategi, I., Navarro-Blasco, Í., Calvo, M.I., Cavero, R.Y., Astiasaran, I. and Ansorena, D. 2014. Healthy reduced-fat Bologna sausages enriched in ALA and DHA and stabilized with Melissa officinalis extract. Meat Sci. 96, 1185–1190.
Bidlás, E. and Lambert, R.J. 2008. Comparing the antimicrobial effectiveness of NaCl and KCl with a view to salt/sodium replacement. Int. J. Food Microbiol. 124, 98–102.
Brennan, J. and Bourne, M. 1994. Effect of lubrication on the compression behaviour of cheese and frankfurters. J. Texture Studies 25, 139–150.
Cáceres, E., García, M.L. and Selgas, M.D. 2008. Effect of pre-emulsified fish oil – as source of PUFA n-3 – on microstructure and sensory properties of mortadella, a Spanish bologna-type sausage. Meat Sci. 80, 183–193.
Campos, M., Nute, G., Hughes, S., Enser, M., Wood, J. and Richardson, R. 2006. Flavour perception of oxidation in beef. Meat Sci. 72, 303–311.
Candogan, K. and Kolsarici, N. 2003. The effects of carrageenan and pectin on some quality characteristics of low-fat beef frankfurters. Meat Sci. 64, 199–206.
Cayré, M., Garro, O. and Vignolo, G. 2005. Effect of storage temperature and gas permeability of packaging film on the growth of lactic acid bacteria and Brochothrix thermosphacta in cooked meat emulsions. Food Microbiol. 22, 505–512.
Coates, A.M., Sioutis, S., Buckley, I.D. and Howe, P.R. 2009. Regular consumption of α-tocopherol-enriched pork modifies cardiovascular risk factors. Brit. J. Nutr. 101, 592–597.
Código Alimentario Argentino. 1999. de la Canal y Asociados, Buenos Aires, Argentina.
Colmenero, F., Ay, O. and Carballo, J. 2005. Physicochemical properties of low sodium frankfurter with added walnut: Effect of transglutaminase combined with caseinate, KCl and dietary fibre as salt replacers. Meat Sci. 69, 781–788.
DESMOND, E. 2006. Reducing salt: A challenge for the meat industry. Meat Sci. 40, 488–490.

EFSA-EUROPEAN FOOD SAFETY AUTHORITY. 2012. Scientific opinion on the tolerable upper intake level of eicosapentaenoic acid (EPA), docosahexaenoic acid (DHA) and docosapentaenoic acid (DPA). EFSA Panel on dietetic products, nutrition and allergies (NDA). EFSA J. 10(7), 2815.

ESTÉVEZ, M., VENTANAS, S. and CAVA, R. 2005. Physicochemical properties and oxidative stability of liver paté as affected by fat content. Food Chem. 92, 449–457.

FELTRIN, A.C., SOUSA, V.R., SARAIVA, C.G., NUNES, C.A. and PINHEIRO, A.C.M. 2015. Sensory study of different sodium chloride substitutes in aqueous solution. Int. J. Food Sci. Technol., 50, 730–735.

FENG, C.H. and SUN, D.W. 2014. Optimisation of immersion vacuum cooling operation and quality of Irish cooked sausages by using response surface methodology. Int. J. Food Sci. Technol. 49, 1850–1858.

FENG, C.H., DRUMMOND, L. and SUN, D.W. 2014. Modelling the growth parameters of lactic acid bacteria and total viable count in vacuum-packaged Irish cooked sausages cooled by different methods. Int. J. Food Sci. Technol. 49, 2659–2667.

FOLCH, J., LEES, M. and SLOANE-STANLEY, G. 1957. A simple method for the isolation and purification of total lipids from animal tissues. J. Biol. Chem. 226, 497–509.

FUNAHASHI, H., SATAKE, M., HASAN, S., SAWAI, H., REBER, H., HINES, O. and EIBL, G. 2006. The n-3 polyunsaturated fatty acid EPA decreases pancreatic cancer cell growth in vitro. Pancreas 33, 462.

GARCÍA-GARCÍA, E. and TOTOSAUS, A. 2008. Low-fat sodium-reduced sausages: Effect of the interaction between locust bean gum, potato starch and α-carrageenan by a mixture design approach. Meat Sci. 78, 406–413.

GEORGANTELIS, D., BLEKAS, G., KATIKOU, P., AMBROSIADIS, I. and FLETOURS, D.J. 2007. Effect of rosemary extract, chitosan and α-tocopherol on lipid oxidation and colour stability during frozen storage of beef burgers. Meat Sci. 75, 256–264.

GHARAVI, N. and EL-KADI, A.O. 2005. tert-Butylhydroquinone is a novel aryl hydrocarbon receptor ligand. Drug Metab. Dispos. 33, 365–372.

HASSABALLA, A., MOHAMED, G., IBRAHIM, H. and ABD EL MAGEED, M. 2009. Frozen cooked catfish burger: Effect of different cooking methods and storage on its quality. Global Vet. 3, 216–226.

HIGGS, J.D. 2000. The changing nature of red meat: 20 years of improving nutritional quality. Trends Food Sci. Technol. 11, 85–95.

HORITA, C.N., MORGANO, M.A., CELEGHINI, R.M.S. and POLLONIO, M.A.R. (2011). Physico-chemical and sensory properties of reduced-fat mortadella prepared with blends of calcium, magnesium and potassium chloride as partial substitutes for sodium chloride. Meat Sci. 89, 426–433.

JACOBSEN, C., LET, M.B., NIelsen, N.S. and MEYER, A.S. 2008. Antioxidant strategies for preventing oxidative flavour deterioration of foods enriched with n-3 polyunsaturated lipids: A comparative evaluation. Trends Food Sci. Technol. 19, 76–93.

JAMORA, J.J. and RHEE, K.S. 2002. Storage stability of extruded products from blends of meat and nonmeat ingredients: Evaluation methods and antioxidative effects of onion, carrot, and oat ingredients. J. Food Sci. 67, 1654–1659.

KIM, Y. 2012. Articles: Utilization of dried garlic powder and α-tocopherol to improve the shelf-life of emulsion-type sausage during refrigerated storage. Korean J. Food Sci. Technol. 32, 725–731.

LANARI, M., SCHAEFER, D., CASSENS, R. and SCHELLER, K. 1995. Atmosphere and blooming time affect color and lipid stability of frozen beef from steers supplemented with vitamin E. Meat Sci. 40, 33–44.

LEE, S., DECKER, E.A., FAUSTMAN, C. and MANCINI, R.A. 2005. The effects of antioxidant combinations on color and lipid oxidation in-3 oil fortified ground beef patties. Meat Sci. 70, 683–689.

LOPEZ-LÓPEZ, I., COFRADIES, S. and JIMÉNEZ-COLMENERO, F. 2009. Low-fat frankfurters enriched with n-3 PUFA and edible seaweed: Effects of olive oil and chilled storage on physicochemical, sensory and microbial characteristics. Meat Sci. 83, 148–154.

MARCHETTI, L., ANDRÉS, S.C. and CALIFANO, A.N. 2014. Low-fat meat sausages with fish oil: Optimization of milk proteins and carrageenan contents using response surface methodology. Meat Sci. 96, 1297–1303.

MARCHETTI, L., ARGEL, N., ANDRÉS, S. and CALIFANO, A. 2015. Sodium-reduced lean sausages with fish oil optimized by a mixture design approach. Meat Sci. 104, 67–77.

NYCHAS, G. and DROSINOS, E. 1999. Meat and Poultry Spoilage. Encyclopedia of Food Microbiology, pp. 1253–1259, Academic Press, San Diego, CA.

PENNISI FORELL, S.C., RANALLI, N., ZARITZKY, N.E., ANDRÉS, S.C. and CALIFANO, A.N. 2010. Effect of type of emulsifiers and antioxidants on oxidative stability, colour and fatty acid profile of low-fat beef burgers enriched with unsaturated fatty acids and phytosterols. Meat Sci. 86, 364–370.
Rhee, K.S. and Myers, C.E. 2004. Sensory properties and lipid oxidation in aerobically refrigerated cooked ground goat meat. Meat Sci. 66, 189–194.

Senso, L., Suárez, M., Ruiz-Cara, T. and García-Gallego, M. 2007. On the possible effects of harvesting season and chilled storage on the fatty acid profile of the fillet of farmed gilthead sea bream (Sparus aurata). Food Chem. 101, 298–307.

Shahidi, F. 1992. Current and novel methods for stability testing of canola oil. Inform 3, 543.

Soiglia, F., Petracci, M., Mudadal, S., Vannini, L., Gozzi, G., Camprini, L. and Cavani, C. 2014. Partial replacement of sodium chloride with potassium chloride in marinated rabbit meat. Int. J. Food Sci. Technol. 49, 2184–2191.

Trikì, M., Herrero, A.M., Jimenez-Colmenero, F. and Ruiz-Capillas, C. 2013. Effect of preformed konjac gels, with and without olive oil, on the technological attributes and storage stability of merguez sausage. Meat Sci. 93, 351–360.

Ulbricht, T. and Southgate, D. 1991. Coronary heart disease: Seven dietary factors. The Lancet 338, 985–992.

USDA. 2010. U.S. Department of Agriculture and U.S. Department of Health and Human Services: Dietary Guidelines for Americans, 7th ed., U.S. Government Printing Office, Washington, D.C.

USDA. 2015. U.S. Department of Agriculture. National Nutrient Database for Standard Reference Release. Available at: http://ndb.nal.usda.gov/ndb/search (accessed November 11, 2014).

Valenzuela, A., Romo, C. and Nieto, M.S. 2011. Tecnologías aplicables a la industrialización de los aceites marinos para permitir su aplicación en la alimentación. Alimentos 20, 1–11.

Yan, P.-M., Xue, W.-T., Tan, S.-S., Zhang, H. and Chang, X.-H. 2008. Effect of inoculating lactic acid bacteria starter cultures on the nitrite concentration of fermenting Chinese paocai. Food Control 19, 50–55.

WHO. 2008. World Economic Forum Report of a Joint Event. Available at: http://www.who.int/entity/dietphysicalactivity/WHOWEF_report_Jan2008_FINAL.pdf (accessed February 25, 2009).

WHO. 2013. The top 10 causes of death, Fact sheet N°310, Updated July 2013, WHO. Available at: http://www.who.int/mediacentre/factsheets/fs310/en/ (accessed November 20, 2013).