Obtaining nitrite from vegetables sources by fermentative process using nitrate-reducing bacteria *Staphylococcus carnosus* and *S. xylosus*

María Carolina Rodríguez-Daza, Diego A. Restrepo-Molina & Mario Evelio Arias-Zabala

*Facultad de Ciencias Agrarias, Universidad Nacional de Colombia, Medellín, Colombia macrodrigueza@unal.edu.co, darestre@unal.edu.co*

Received: January 21st, 2019. Received in revised form: June 28th, 2019. Accepted: August 12th, 2019.

**Abstract**

Sodium nitrate is an essential food additive widely used in cured meats. The increased unfavorable perceptions about chemicals in foods and their repercussions on health have positioned nitrites as undesirable compounds in meat products. Natural and organic ingredients have taken an important place within the consumer preferences. Here, obtaining nitrite from natural sources was achieved using nitrate-reducing bacteria *Staphylococcus carnosus* and *S. xylosus*. Pre-incubation strains into a celery-based medium allowed the stimulation of nitrate reductase activity. The increased values of substrate, the oxygen condition and pH influenced the nitrite generation significantly (p<0.05). The reduction rate from nitrate to nitrite was 50.85%, with a value of 320.2 mg L⁻¹ of nitrite. This method presents characteristics comparable to those traditionally applied in the generation of conventional sodium nitrite leading the food industry to take advantage of fermentation processes to supply needs in meats and supplementary food products.

**Keywords**: sodium nitrite; nitrate; nitrite; nitrate-reductase; *Staphylococcus carnosus*; *S. xylosus*.

**1. Introduction**

In the last few years consumer preferences have been aimed at demanding health-friendly, high quality, nutrient-rich natural products. Research on consumption behavior and market trends has provided evidence on individuals’ unfavorable perceptions of food additives, preservatives, and chemicals, in general [1].

Nitrites are among the most widely used compounds for the preservation of meat products. The food industry values...
their use due to their great contribution to meats’ organoleptic characteristics and antimicrobial effects.

Within curing processes, nitrite is responsible for the red pigment production of cured meats, participating in the development of flavor and in slowing oxidative rancidity. In addition, it contributes to growth inhibition of pathogenic bacteria such as Clostridium botulinum [2]. Nevertheless, over the years great concerns have been expressed regarding the exposure of consumers to certain harmful products that may be formed in meat and meat products during and after curing [3]. The use of nitrates for meat curing became a major concern in the 1960s when it was discovered that nitrates have the potential to form carcinogenic nitrosamines when combined with secondary amines in a mildly acidic environment [4,5].

Although they have been considered to be negative in foods due to toxicological reasons, there is no strong epidemiological evidence for a correlation between nitrosamine formation and the incidence of gastrointestinal types of cancer. A recent meta-analysis of published papers (systematic literature searching of PubMed) provided a quantitative assessment of their relationships with gastric cancer, explaining that whereas human diet is a potentially modifiable exposure, it remains difficult to attribute the cancer etiology to a single nutrient [6].

In addition to nitrite and nitrate’s essential functions such as food additives, those compounds have caused a deep impact on many physiological effects. Dietary consumption of nitrate has been demonstrated in clinical studies to have numerous health benefits, especially related to improving cardiovascular function and blood pressure [7]. Likewise, potential effects have been associated to the inorganic nitrate (NO3-) inclusion on nutritional supplementation products in order to enhance muscle efficiency, tolerance and exercise resistance [8,9].

As we have seen, those compounds harbor relevant effects in different contexts. In the sense of the decreased consumer intentions toward conventionally produced and processed foods, studies have been conducted to evaluate new alternatives and develop innovative products that may support the essential nitrite and nitrate functions. In order to effectively respond to modern consumer needs, the food industry has focused research towards the inclusion of natural and organic ingredients that would maintain the quality and microbiological safety of the product. Natural curing processes and antimicrobial natural sources are examples of actual trends, decreasing components and chemical additives in meat products [10]. Nitrate/nitrites can be indirectly added to meat products without losing the common sensory effects of traditionally cured ones. Ingredients that naturally contain nitrates reflect the current trends [11].

Vegetables offer a great potential for introducing nitrates/nitrites in foods [12]. Examples of them are the families of Brassicaceae (rocket, radish, mustard), Chenopodiaceae (spinach, beets, spinach) Amaranthaceae, Asteraceae (lettuce) and Apiaceae (celery, parsley) [12]. The broad interest highlighted on the organic and natural products intake have positioned those natural sources as potential alternatives to the replacement or the reduction of synthetic nitrates and/or nitrites. Here, the reduction of nitrate to nitrite is a key step. This reaction may be represented by the use of chemistry-reducing agents, enzymatic processes, or by microorganism nitrate-reducers. Obtaining nitrate as substrate for nitrite generation through the action of nitrite-reducing bacteria, recognized as safe, represents a promising approach. The coagulase-negative staphylococci group (CNS) are well known, not only for presenting the enzymatic repertoire for this process, but for its compatibility with foodstuffs [13,14].

Considering the biological variability and metabolic behavior of the nitrate-reducing bacteria, it results important to study the appropriated conditions of nitrate-reductase activation. The expression of this enzyme in staphylococci, for example, may be induced by anaerobic growth under the nitrate presence, with the maximum level of activity during the exponential growth phase [15].

In line with this, the present study aimed to evaluate the nitrite generation from natural sources by fermentative process using nitrate-reducing bacteria. This alternative would allow enlarging new processed meat and supplementary food products with added natural compounds and a reduced level of nitrite to increase.

2. Methods

2.1. Nitrate extractions

Vegetable samples (broccoli and celery) were acquired from greengroceries of Medellin, Colombia. After transmission of the samples to the laboratory, they were washed with distilled water. Initially, different extraction methods were tested. 1) Disinfection/Maceration, 2) Scalding and 3) Maceration/autoclaving. They all were green extractions, based on the use of water, heat and mechanical disruptions in order to allow the derived product to be applied on foods.

For the first method, similar vegetables were mixed together and disinfected under submersion into ethanol 70% for exposition intervals of 5-10 minutes. Then, the samples were washed at least three times with sterile distilled water and led to dry at laboratory temperature for subsequent blender-based maceration. The scalding was applied to the same type of washed vegetables using a ratio water: substrate for nitrite generation through the action of nitrite-reducing bacteria, recognized as safe, represents a promising approach. The coagulase-negative staphylococci group (CNS) are well known, not only for presenting the enzymatic repertoire for this process, but for its compatibility with foodstuffs [13,14].

Considering the biological variability and metabolic behavior of the nitrate-reducing bacteria, it results important to study the appropriated conditions of nitrate-reductase activation. The expression of this enzyme in staphylococci, for example, may be induced by anaerobic growth under the nitrate presence, with the maximum level of activity during the exponential growth phase [15].

In line with this, the present study aimed to evaluate the nitrite generation from natural sources by fermentative process using nitrate-reducing bacteria. This alternative would allow enlarging new processed meat and supplementary food products with added natural compounds and a reduced level of nitrite to increase.

2.2. Nitrate-reducing strains

Lyophilized strains Bactoferm C-P-77-S and S-SX composed by Staphylococcus carnosus, Staphylococcus xylosus and Lactobacillus pentosus were used (Chr Hansen Hanssen Inc., Milwaukee, Wis., U.S.A). A qualitative adaptive process was performed in order make conditions favorable for the microorganism tolerance to the new vegetable-based conditions of the fermentable media. Two types of culture media were assayed, a commercial Brain
heart infusion Broth (BHI OxoidTM) (37g/L) and a celery-based homemade broth (10% w/v) supplemented with yeast extract (5 g/L) and glucose (1 g/L). The culture media were autoclaved at 121°C, 15 psi, for 20 min.

A hydration/activation step of the strains was carried out into the prepared broths at 37°C for 24h. Succeeding the activations, the strains were cultured in selective agar plates with the purpose to obtain isolated colony forming units (CFU). Here, the Mannitol Salt agar (MSA) was used to isolate the Staphylococcus from L. pentosus, considering that only the Sthaphylococcus strains have the enzymatic potential to reduce nitrate to nitrite. Once, obtained the characteristic morphology of the target microorganisms S. carnosus and S. xylosus, those were transferred to plate count agar (PCA) and incubated at 37°C for 24h. Finally, the colonies were used to prepare the inoculums (the Celery-agar (PCA) and incubated at 37˚C for 24h). Finally, the colonies were used to prepare the inoculums (the Celery-agar (PCA) and incubated at 37°C for 24h). Finally, the colonies were used to prepare the inoculums (the Celery-agar (PCA) and incubated at 37°C for 24h). Finally, the colonies were used to prepare the inoculums (the Celery-agar (PCA) and incubated at 37°C for 24h). Finally, the colonies were used to prepare the inoculums (the Celery-agar (PCA) and incubated at 37°C for 24h). Finally, the colonies were used to prepare the inoculums (the Celery-agar (PCA) and incubated at 37°C for 24h). Finally, the colonies were used to prepare the inoculums (the Celery-agar (PCA) and incubated at 37°C for 24h).

### 2.3. Evaluation of fermentation conditions

Different fermentation processes were performed to look for the best conditions for an appropriated nitrate-reductase activity stimulation. Among them, the use of pre-adapted inoculums, the incubation under different oxygen conditions, pH values, fermentable culture supplementations, and the use of dried or fresh vegetal material were considered. Then, combinations of above mentioned conditions were examined. Two experimental replications and two technique replications were carried out for each condition. The use of synthetic sodium nitrate was used as control of fermentable substrate source.

#### 2.3.1. Strain’s pre-adaptation step and conditions of incubation

Initially, the inoculums activated either using the celery-based homemade broth were used separately to evaluate the effects of the strain’s pre-adaptation step to the fermentable vegetal conditions over the nitrate-reductase activity. Since, the blended fermentable media submitted to autoclaving process (121°C, 15 psi, for 20 min) showed to favor the nitrate extraction, no application of preceding extraction methods was needed.

Considering that the presence of oxygen may be also an influential factor on the stimulation of the nitrate and nitrite reductase activity of *S. carnosus* and *S xylosus*, the fermentation process was carried out under both aerobic and anaerobic conditions. The anaerobic atmosphere was achieved by pumping nitrogen gas into the Erlenmeyer flasks for 3 minutes (dimensions 140 x 83 mm, height x widest diameter) and sealing it with rubber stopples. Anaerobic indicators (BD, BBL Dry anaerobic Indicator strips, Becton Dickinson and Company) were introduced in each flask. The experiment was incubated in a covered shaker water bath (SW23, Julabo) at 30°C, 150 rpm for 24h.

#### 2.3.2. Influence of pH values

Once found the anaerobic atmosphere and strains pre-adaptation as determining features to prompt the bacteria enzymatic action, variations in the pH values of the fermentable media were assessed. For this purpose, only the base of fermentable media, mainly composed by celery 10% w/v, yeast extract 5 g/L, and dextrose 1g/L were used. The pH values ranged from 7 to 10. The inoculum adapted on the nitrate-rich celery-based broth (10% v/v) was aseptically added to the fermentable media under a biological laminal flow chamber. The media without pH adjusting was took like negative control. The experiments were incubated in a covered shaker water bath (SW23, Julabo) at 30°C, 150 rpm for 24h. Experiments were performed in duplicate.

#### 2.3.3. Supplementation of celery-based fermentable media

The fermentable media was mainly composed of 10% w/v celery, yeast extract 5 g/L, and dextrose 1 g/L. But, they were either enriched with BHI 10 g/L, potassium caseinate 10 g/L, NaCl 2 g/L, K2 HPO4 13 g/L. The addition of those compounds to the media varied as shown in the Table 1. Consequently, three different media compositions were evaluated.

Prior studies showed an increased nitrite concentration in the fermentable media during the first hours of incubation. In this sense, here, unlike the previous fermentations, the duplicated flasks were incubated until 6 hours under the same aerobic and anaerobic conditions as above explained.

#### 2.3.4. Use of dried or fresh vegetal

The nitrate level in vegetable products is highly variable. It depends on the biological properties of the vegetal culture, such as the plant size, soil composition, density growth, plant maturity, duration of growth, harvest time, nitrogen source, time and storage conditions [16]. In this study, we intended to standardize the fermentable medium using a laboratory-made normalized celery powder. Then, both the fresh celery based medium (10% w/v) and the celery powder based medium (2.5% w/v) were fermented. The media composition was BHI 10 g/L, potassium caseinate 10 g/L, NaCl 2 g/L, K2 HPO4 13 g/L and 10% inoculum v/v. The

---

**Table 1. Fermentable culture media compositions.**

<table>
<thead>
<tr>
<th>Component</th>
<th>Media 1</th>
<th>Media 2</th>
<th>Media 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Celery powder (25 g/L)</td>
<td>5 / 20 g</td>
<td>5 / 20 g</td>
<td>5 / 20 g</td>
</tr>
<tr>
<td>Yeast extract (5 g/L)</td>
<td>1 g</td>
<td>1 g</td>
<td>1 g</td>
</tr>
<tr>
<td>Dextrose (1 g/L)</td>
<td>0.2 g</td>
<td>0.2 g</td>
<td>0.2 g</td>
</tr>
<tr>
<td>NaCl (2 g/L)</td>
<td>0.4 g</td>
<td>0.4 g</td>
<td>—</td>
</tr>
<tr>
<td>K2 HPO4 (13 g/L)</td>
<td>2.6 g</td>
<td>2.6 g</td>
<td>—</td>
</tr>
<tr>
<td>Brain heart infusion-BHI (10 g/L)</td>
<td>2 g</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Potassium caseinate (10 g/L)</td>
<td>—</td>
<td>2 g</td>
<td>—</td>
</tr>
<tr>
<td>Distilled water (mL)</td>
<td>200</td>
<td>200</td>
<td>200</td>
</tr>
<tr>
<td>pH value</td>
<td>9.0</td>
<td>9.0</td>
<td>9.0</td>
</tr>
</tbody>
</table>

Source: The Authors.
fermentations were carried out at 30°C, in constant agitation (150 rpm) for 6h, under anaerobiosis.

2.4. Analysis of nitrate/nitrite concentrations

2.4.1. Samples preparation

As nitrites were extracted by the autoclaving process of each celery-based fermentable media, aliquots were taken before and after fermentation to determine nitrate/nitrite contents. Samples were centrifuged at 3000 x g for 15 min (Botofix 32a, Hettich Zentrifugen); the supernatants collected were filter-purified (0.2μm Syringe filter, membrane NYL) to eliminate bacterial cells, and were then, transferred to 1 mL dark micro-vials. All samples were immediately analyzed within 1 hr. after sample preparation.

An Agilent 1260 infinity - high performance liquid chromatography system, equipped with photodiode-array UV detector was utilized in this study. A C18 column (5 μm, 250 4.6 mm I. D.) was employed for separation.

2.4.2. Preparation of mobile phase and standard solutions

Aqueous solutions consisted of Potassium dihydrogen phosphate (KH₂PO₄ Merck 99.5 per cent pure) and Sodium hypophosphite (Na₂HPO₂ Merck Darmstalt, Germany 99% Purity), prepared in deionized Milli-Q water (Millipore). The pH values were monitored using pH-meter HandyLab (Schott instruments) and Analytics pH electrode Blueline. The adjustment of pH value to 3.0 was done by adding phosphoric acid.

Standard solutions containing 200 mg L⁻¹ of sodium nitrate and sodium nitrite were prepared, adjusted pH to 6.15. The pH values were monitored using pH-meter HandyLab (Schott instruments) and Analytics pH electrode Blueline. The solutions were freshly prepared for every running HPLC chromatograms. The calculated standard curve and correlation coefficients was indicative of the linearity within the tested range of concentrations.

2.4.3. HPLC analysis

In this study, the evaluation of the nitrate-reductase activity of S. carnosus and S. xylosus viable cells was based on the measurements of residual nitrate and the generated nitrite in the fermented celery-composed media. The mobile phase solution was allowed to pass through the HPLC column until a stable baseline signal was equilibrated. The flow rate was 1.0 mL/min and the detecting UV wavelength was 214 nm. When the injections of the nitrate/nitrite standard solutions gave stable and reproducible retention times and peak areas, each filtered sample was then injected for analysis. The injection volume was 5.0 μL. The peaks of the samples were identified by comparison to the fitting peaks of the standards. When the curve of the peak areas was larger than that of the maximum amount from the standard curve, the test solution was diluted to appropriate concentrations. The amounts of nitrate and nitrite in the test solution were calculated from the peak areas by using linear regression equations of nitrate and nitrite standard curves. At the end of the analysis, the HPLC column was washed.

2.5. Statistical analysis

The statistical significance of the differences in the measured parameters in each fermentable medium type were tested by the analysis of variance (ANOVA) test. Here, the generation rate of nitrite was compared. A student-t test was further performed to compare the mean of nitrate/nitrite levels, variations in the vegetal type and pH values. The results shown in the figures are the mean values ± SD. Mean values are significantly different at P < 0.05, as indicated by asterisks (*) or letters in the figures. Graphpad Prism 7.0 software was used.

3. Results and discussion

The concerns about the use of nitrites are extensively linked to the formation of toxicological compounds in cured meat products. However, less is known about the traditional methods used for obtaining synthetic nitrate/nitrite at industrial level, which may embrace process highly intervened by chemical agents. Here, we presented an alternative approach to obtain nitrite from natural sources attuned with the current consumers preferences. Natural nitrite differs from synthetic nitrite in the way that they are produced. Synthetic nitrites are linked to chemical processes, as are typically created by the use of C (carbon), Fe (iron) or Pb (lead) for nitrate reduction at elevated temperatures. The industrial manufactured NaN₃ has been described as a result of bubbling a mixture, comprising NO and NO₂ in a hydroxide solution by alkali metal nitrates [17].

A variety of vegetables have shown high nitrate contents. In the present study, we evaluated food-compatible nitrate extractions of two vegetables with textural differences, broccoli and celery. The application of three extraction techniques resulted in obtaining nitrate levels from 795.87 mg L⁻¹ to 1045.62 mg L⁻¹ for celery samples and from 189.48 mg L⁻¹ to 261.74 mg L⁻¹ for broccoli. The nitrate quantities exhibited variations according to the applied extraction method and vegetable type. The application of maceration/autoclaving technique registered the high nitrate value for both celery and broccoli, with 1045.62 mg L⁻¹ and 261.74 mg L⁻¹ respectively. Significant differences were found when compared to the other applied techniques (p<0.05). On the other hand, scalding and disinfection/maceration procedures showed similar extracting abilities for both vegetables, resulting in 852.48 and 795.87 mg L⁻¹ in the case of celery and 199.13 and 189.48 mg L⁻¹ nitrate for broccoli. Those nitrate levels are comparable with previous studies. Iammarrino et al. (2004) reported extraction values from 35.0-1800 mg L⁻¹ of nitrate using a homogenization-based method at 80°C for 5 min, and values so high as 2978.1 and 2291.8 mg L⁻¹ of nitrate obtained from frizzed and fresh spinaches [18]. The results here exposed reveal the importance of heat application during the extraction phase. Despite variations on the design of the techniques reported in several studies, including differences in the vegetable type and time treatment, all have as common element the use of hot water preceded by proper vegetable homogenization [19, 20]. It was perceived that when extractions were performed at bench temperature or lower than 80°C (the case of
disinfection/maceration and scalding), the obtained nitrate concentration was reduced. Consequently, the maceration/autoclaving was kept as a combined method of nitrite extraction and sterilizations, which at time, guarantees an aseptic media for the strains metabolic activity.

Since the celery vegetable showed higher substrate levels, and exhibited a reduced pigmentation in the obtained juice, it was chosen as the appropriate source to further fermentations. Nitrate reduction to nitrite was improved when used the inoculum pre-adapted on the nitrate-rich celery-based broth. The HPLC chromatograms (Fig. 1) showed an elevated nitrite level of 392.035 mg L\(^{-1}\) obtained from celery-based fermented media for 24h and did not registered nitrite levels in the synthetic nitrate-based fermentable control media. While the strains were pre-incubated in the commercial BHI broth, the nitrite obtained from the nitrate-rich celery-based broth was diminished, resulting in 49.497 mg L\(^{-1}\) of nitrate.

In general, there was no generated nitrite, when using the strains not adapted; a reduced nitrate was found after 24h of the anaerobic fermentation process, compared to the initial content. Here, the substrate level fell 26.6% compared to the non fermentable media. All the experimentations carried out under aerobic conditions did not show nitrite generation. On the contrary, the celery-based fermentations submitted under anaerobic atmosphere presented a mean nitrite level of 392.035 mg L\(^{-1}\) ± 5.045, indicating a 73.27% of reduction from nitrate to nitrite (535 mg L\(^{-1}\) initial nitrate and 392.035 mg L\(^{-1}\) final nitrite).

The earlier strains exposition to the vegetal material favor the enzymatic activity; therefore, all accumulations of nitrite was the result of the action of adapted viable cells. The induction of the nitrate-reductase activity on strain of *S. carnosus* have been reported elevated when the substrate was added to the growth culture medium. The nitrate-reducing rate was 8 to 10 times higher in anoxic conditions [15]. These notions are clearly supported by the results here obtained considering that the pre-adaptation step to the fermentable vegetable and the anaerobic atmosphere were key factors in reflecting improved nitrate-reducing rate. It was noticed that the stimulation of nitrate-reductase activity was proportional to the increased nitrate concentration in the fermentable media, suggesting that nitrate reduction is not attached to the sources type, but to enough substrate availability. In this context, authors have explained that the NO\(_2^-\) liberation to the media by the bacteria take place if the NO\(_3^-\) level in the media is higher than 5 mM [21]. This allows the bacteria using the nitrate produced intracellularly as nitrogen source for the cellular metabolism, and hence, it is not secreted to the external media.

During respiration, the bacteria *S. carnosus* and *S. xylosus* use nitrate as electron receptor when the oxygen is not available. Thus, the nitrate reduction is coupled to the generation of a driving force of protons which is directly taken as energy source [15]. Associated mechanisms to the activation of the nitrate-reductase activity still need to be elucidated. However, prior research has reported a regulator gene named *NreABC*, in charge of the detection of anoxic conditions, as well as, the nitrate presence, resulting in the nitrate-reducing activation system on the bacteria [22].

The first goal of this study was to find out the appropriate conditions of the fermentation process that guaranteed at time, an elevated staphylococci enzymatic action. Regarding to the evaluated pH values, better nitrate-reduction was registered at pH between 8.0 to 9.0, with 255.25 and 281.39 mg L\(^{-1}\) of nitrite respectively (Table 2.). The ordinary one-way ANOVA analysis showed significant difference between the treatments p <0.05. The media which pH value was not adjusted, presented acid pH ranged between 4.0 to 5.0. Multiple comparisons test against unadjusted experimentations were statistical significant p<0.05. Once compared by *student-t* test the two higher registered levels of nitrite (pH 8.0 and 9.0), a difference was observed (p 0.0089). It’s possible that the pH decrease during the fermentation was the result of glucose degradation, and hence, the appearance of derived metabolites such as acetate and pyruvate [21]. Here, the use of buffers in the fermentable media are

![Figure 1. HPLC chromatograms of fermented samples for 24h. A-B) Nitrate-rich celery-based media C) Nitrate/nitrite sodium standards. Source: The Authors.](image-url)
Table 2.
Peaks area of nitrate and nitrite obtained on running chromatograms from samples at different pHs.

<table>
<thead>
<tr>
<th>pH</th>
<th>Ret. time (min)</th>
<th>Area</th>
<th>mg L⁻¹ nitrate</th>
<th>Ret. time (min)</th>
<th>Area</th>
<th>mg L⁻¹ nitrite</th>
</tr>
</thead>
<tbody>
<tr>
<td>CT</td>
<td>1.359</td>
<td>10697.1</td>
<td>306.61</td>
<td>1.774</td>
<td>414.2</td>
<td>36.84</td>
</tr>
<tr>
<td>7</td>
<td>1.352</td>
<td>10136.9</td>
<td>290.55</td>
<td>1.778</td>
<td>2152.9</td>
<td>191.4°</td>
</tr>
<tr>
<td>8</td>
<td>1.354</td>
<td>11724.9</td>
<td>336.07</td>
<td>1.725</td>
<td>2869.9</td>
<td>255.2°</td>
</tr>
<tr>
<td>9</td>
<td>1.355</td>
<td>11837.9</td>
<td>339.31</td>
<td>1.727</td>
<td>3163.8</td>
<td>281.3°</td>
</tr>
<tr>
<td>10</td>
<td>1.359</td>
<td>10172.0</td>
<td>291.56</td>
<td>1.771</td>
<td>845.3</td>
<td>75.18°</td>
</tr>
<tr>
<td>SD</td>
<td>1.358</td>
<td>6977.66</td>
<td>200</td>
<td>1.721</td>
<td>2248.7</td>
<td>200</td>
</tr>
</tbody>
</table>

*Values significantly different compared to unadjusted pH control (CT), SD (standard) p<0.05. Treatment with distinct letters are statistically different by unpaired t-test p<0.05.

Source: The Authors.

important to maintain a balanced pH and support the growth rate of the Staphylococci strains, which have been reported inhibited at pH 4.5 [23].

The glucose, yeast extract, and the presence of celery-derived nitrate as fermentable substrate played an important role as electron donor (glucose) and electron acceptor (nitrate). When they are added to the fermentable media under anoxic atmosphere, the biomass production and the nitrate reduction may increase. This suggests that there was a diminution on the energetic efficiency of the bacterial anaerobic respiration when incubated under oxygenc condition, and then the glucose could be catabolized to other metabolites [24]. The fermentable media composition found adequate to further fermentation process was then achieved (see Table 1). Its components were fresh celery, yeast extract, dextrose, BHI and NaCl and buffer agents. Despite the use of celery powder allowed to standardize the vegetal source, the addition of 2.5% w/v to the fermentable medium showed decreased solubility, affecting the agitation during the fermentation process. Here, the fermentation kinetic for 6 hours showed better nitrate-reduction rate, registering the higher nitrate-reductase activity between 4-5 hours (Fig. 2).

The enzyme nitrate-reductase on S. carnosus have been described to be attached to the membrane and involved on the conservation of the respiratory energy [15]. The maximum profile expression has been reported during the exponential phase. On the other hand, the enzyme nitrite-reductase may be found in a cytosolic fraction. The action of this enzyme only is activated under anaerobic conditions and is induced by the presence of nitrate or nitrite in the growth media [25]. Here, the indirect regulation of those enzymes may be achieved by modulating the concentrations of the substrates in the fermentable media.

S. carnosus and S. xylosus may reduce nitrate to ammonia in two steps: First, the nitrate is absorbed and reduced to nitrite, and the nitrite is excreted later. Second, after the depletion of the nitrate in the fermentable media, the accumulated nitrite is imported to the interior of the cell, reduced to ammonia, and this one, is accumulated again in media [22,26].

It is possible that the reduced nitrite level in most of the fermentations that took 24 hours was obtained as result of the catabolism of the produced nitrite, once the nutrient and other substrates could be also diminished. Consistent with the above results, short periods of fermentations were performed to catch earlier levels of produced nitrite and prevent the simultaneous degradation of the same secreted nitrite from the media.

Conditions found to favor nitrate-reductase activity of S. carnosus and S. xylosus consisted of a chain of steps, starting with the pre-adaptation strains to a nitrate-rich celery-based medium, followed by the composition of fermentable media, then the adjustment of pH value was adjusted to 9.0, after which the anoxic condition of incubation was induced. Finally, the medium was exposed to agitation. Additionally, no less important, was the time of fermentation (4 h). After those characteristics favorable for generating nitrite were identified, additional experimentations were performed. Consequently, after 4h of fermentative process of celery-based media, a nitrate-reduction rate of 50.85% was observed, passing from 629.42 to 309.22 mg L⁻¹ of nitrate and generating 302.2 mg L⁻¹ of nitrite. As seen, the entire procedure for obtaining nitrites/nitrate is advantageous to apply to the meat industry. Within the meat industry,
comparable findings have been reported regarding to the residual nitrate contents in manufactured meat products. Prior studies have demonstrated that the use of sodium nitrite in cooked ham results in a residual nitrate level of around 90 mg Kg$^{-1}$ compared to the use of celery juice as a source of nitrate, which results in a residual level of nitrate between 30 mg Kg$^{-1}$ and 60 mg Kg$^{-1}$ [27].

4. Conclusions

Nitrate from vegetable sources were obtained in the present study without using chemical agents. The parameters evaluated led to select processes as simple as the mechanical maceration and sterilization by autoclave, like adequate technique to extract nitrate. Key parameters were identified to run fermentative process with better yield on staphylococci nitrate reduction. Here, the availability of high substrate level as crucial element regulator on the nitrate-reductase activity was underway, as well as the previous adaptation of the bacteria to a nitrate-rich selective environment was identified.

The use of enzymatical agents and nitrate-reducing microorganisms, generally recognized as safe (GRAS), promise to be better alternatives and to reduce the use of chemical compounds in the production of nitrite. These non-toxic processes, are useful, not only to for the meat industry as an agent of curing, but also for the formulation of food supplements, with emphasis in sports performance, and other physiological endeavors, which recently have been associated with benefiting from nitrate consumption. Consumer preferences towards natural products is leading the food industry to seek natural options. The processes outlined in this paper can help supply the market needs for natural products.

Acknowledgements

We thank Tecnas S.A., Medellin, Colombia for the financial support.

Bibliography

parallel operated small-scale bubble columns for microbial process development using Staphylococcus carnosus, J. Biotechnol., 88, pp. 77-84, 2001. DOI: 10.1016/S0168-1656(01)00265-6


M.C. Rodríguez-Daza, received her BSc. in Bacteriology and Clinical Laboratory in 2010 and the MSc. in Food Science and Technology in 2014 of the Universidad Nacional de Colombia, Medellin, Colombia. Currently, she is studying for a PhD. in Food Microbiology in Laval University, Quebec, Canadá. Her research interests include clinical and food microbiology, knowledge in functional food compounds and modulation of the gut microbiota.

ORCID: 0000-0001-6510-1804

D.A. Restrepo-Molina, received his BSc. Eng in Chemical Engineering in 1982, from the Universidad Nacional de Colombia, Medellin, Colombia and his Ms.S in Food Science and Technology in 1998, from the Universidad de la Habana, Cuba. Currently, he is an associate professor in Agricultural and Food Engineering Department, Facultad de Ciencias Agrarias, Universidad Nacional de Colombia, Medellin. His research interests include science and technology of meat, quality control in food and biochemistry of meat.

ORCID: 0000-0002-5623-3319

M.E. Arias-Zabala, received the BSc. Eng. in Chemical Engineer in 1983, MSc. Technology of Chemical and Biochemical Processes in 1991 and PhD. in Chemical Engineering. Currently, he is an associate professor in the Agricultural and Food Engineering Department, Facultad de Ciencias Agrarias, Universidad Nacional de Colombia, Medellin. His research interests include plant biotechnology, biochemistry and biochemical engineering.

ORCID: 0000-0001-8190-7760