Meat Pet Snacks By Containing Encapsulated Saccharomyces Boulardii

Sheila Baroncello, Nathalia Turkot Candiago, Jane Mary Lafayette Neves Gelinski, Vinicius Caliari and César Milton Baratto

Abstract— The global trend is to minimize losses by using by-products from slaughterhouses as part of sustainable food production, both for human and animal consumption. Another growing trend is the development of pet feed with properties beneficial to the consumer’s health. Another trend is the development of feeds with beneficial properties for the health of animals. Thus, the objective of this study was to use mechanically separated pork meat and by-products (trachea) to produce a pet snack containing the encapsulated probiotic yeast Saccharomyces boulardii. The experimental design followed in three main stages: 1) development of a pet pork snack (trachea snack); 2) microencapsulation of the probiotic and insertion in the pet snack; 3) Shelf life evaluation and palatability of the pet snack. S. boulardii was microencapsulated by extrusion technique and remained viable after 120 days of shelf life. The acceptability index of the probiotic pet snack was 77.8%. We concluded that S. boulardii provides beneficial probiotics characteristics to pet snack whose composition has high levels of protein (57.50%). Thus, the novel pet product has a good appeal to the functional products market.

Index Terms — Encapsulation, Extrusion, Pet feed market, Probiotic, Yeast.

I. INTRODUCTION

The pet food market is booming. The U.S. is by far the largest pet food market [38]. However, with 54.2 million dogs and 23.9 million cats, Brazil is the second largest market for pet products in the world, reaching 20.3 billion in 2018 [1]. The nutrition of dogs and cats has for some time been equated with human nutrition, with the incorporation of functional substances in food [10]. The market has been adapting to the needs and creating new specific products and services of the most varied formats and prices, all to keep the animal within the family reality [40]. In this sense, important trends [55], [8] are established in the sector of products aimed at domestic animals:

1) segmentation: product that meets specific needs by: stage of life (puppy, adult, senior), specific diets, size of the animal (small, medium, large), breed, among others;
2) premium market: increasingly noble and specific products;
3) humanization: attention to the close relationship between dog and owner;
4) practicality of the food: pet owners have an arduous work routine and seek dry, practical foods, beneficial for the consumption and health of their animals;
5) market distribution: the ease of access, the increase in logistics networks allows for a closer relationship between suppliers, distributors and consumers, due to the increasing insertion of products in regional markets.
6) health and well-being: product with functionality: prebiotics, probiotics and symbiotics.

Probiotics are beneficial microorganisms, supporting animal health that provide greater longevity and decrease the risk of developing diseases throughout life; they are also alternatives to the treatment and prevention of diarrhoea [52], [6], [33], [19].

Within the group of beneficial microorganisms, the probiotic yeast Saccharomyces boulardii, has been widely used for the treatment of diarrhoea and inflammatory conditions of the gastrointestinal tract [50], [30]. Clinical trials of S. boulardii have shown probiotic activity of this yeast, as it is considered a probiotic supplement to support gastrointestinal health, since it prevents intestinal infections caused by different pathogens [59].

In addition to being effective against infections caused by pathogenic bacteria, S. boulardii has an anti-secretory and anti-inflammatory effect [47], antibacterial and antimutagenic action [56], [36]. When ingested, yeast is not established in the gastrointestinal tract, however, it is very well absorbed by the organism, remaining viable during treatment [44], [54]. Briefly, the mechanisms of action of S. boulardii can be classified into three points: luminal action, trophic action, and anti-inflammatory effects on the intestinal mucosa [41].

Various studies [4], [21], [42] have shown that to maintain the probiotic potential of beneficial microorganisms in the host organism, encapsulation is an alternative of multiple applications. It is a technique in which the encapsulated is protected from the action of the gastrointestinal tract, with release in a controlled manner and under conditions specific [29].

In the present study, S. boulardii was encapsulated to form a snack because the target audience consists of dogs and cats whose stomach pH is around 2.5 - 3 and between 5.7 - 6.4 in the intestine [20]. Microencapsulation is a way...
of ensuring that the yeast during consumption reaches the intestine in a viable way and tolerates sudden changes in pH, without undergoing major changes [2]. Therefore, in the present study, by-products of swine slaughterhouses were used to develop a meat snack for dogs and cats containing encapsulated S. boulardii. In addition, we seek to contribute to the sustainable development of the food production chain by providing a product with probiotic properties and aimed at the pet market. Thus, the aim was to use mechanically separated pork meat and by-products (trachea) to produce a pet snack containing the encapsulated probiotic yeast S. boulardii.

II. MATERIAL AND METHODS

A. Probiotic yeast viability tests

The viability of the yeast Saccharomyces boulardii obtained in lyophilized form (COANAn-Import and Export Ltd) was initially assessed by counting in a Neubauer chamber (New Optik) from dilution (1/10) of the sample in 0.85% saline solution, and analysis by optical microscopy (Nikon Eclipse E100, obj.400x). Differentiation of live cells from dead cells was carried out with the 2% Trypan blue dye (Laborclin Brazil). The number of viable cells per mL was obtained with equation 1:

\[ n^\circ \text{of cell per mL} = \frac{\text{total number of cells}}{\text{number of quadrant counted}} \times \text{dilution factor} \times 10000 \quad (1) \]

B. Evaluation of the yeast under different growth conditions

1. Temperature and oxygen

To assess the best temperature and condition of yeast development, it was inoculated in Yeast Extract and Glucose (YEG) broth by using 5 g / L of yeast extract (Bacto™) and 20 g / L of glucose (Synth Brazil). The incubation was carried out at different temperatures: 25°C, 28°C, 35°C and 37°C, for 48 hours in aerobic and anaerobic conditions.

Salinity Test. Aliquots of S. boulardii cultures previously grown in YEG broth for 24 hours were subjected to solutions with different concentrations of salt (NaCl): 1%, 1.5%, 2%, 2.5% and 3%, incubating at 25°C for 24 hours.

pH test. To assess the degree of acidity that the probiotic yeast is capable of withstanding, the YEG broth was used by changing the pH with a 7M HCl solution, (Quimis®, São Paulo) to: 1.5; 2.5; 3; 3.5; 4 and 4.5. The test solutions were incubated at 25°C for 72 hours.

2. Growth curve assessment

To evaluate the number of viable cells, a yeast culture was performed in YEG broth and incubated at 25°C for 72 hours in aerobic conditions. After that, an aliquot was inoculated into Erlenmeyer containing YEG broth for a final dilution of 1:100, after transferring it to a shaker incubator (Nova Ética, São Paulo) at 200 rpm, 25°C for 24 hours. At 2-hour intervals, a spectrophotometer reading (Pró-Análise, Brazil) was performed at λ600nm. At the same time, inoculations of each dilution were performed using the spread plate technique in YE agar plates and incubating at 25°C by until 72 hours.

The total dry mass of the sample was evaluated by gravimetric analysis. About 20 mL of sample obtained under the same conditions as above were filtered through a cellulose acetate membrane (0.22 µm) previously dried in a drying oven (Gigante, São Paulo) at 60 °C until constant mass. Then, after filtration, the membrane containing the cell mass was kept at 65°C for 1 hour until an inert weight was obtained.

C. Development of meat snack for pets

1. Feedstock

The raw material of animal origin came from establishments registered with the Federal Inspection System of Brazil. The samples were transported in thermal boxes (maximum temperature of 4 °C) and product temperature monitoring at reception. For the development of the meat product, the following were used: a) swine and gut trachea; b) lyophilized S. boulardii. The storage took place at temperatures of -20 °C until the analysis and release for processing.

2. Standardization and dehydration

After cleaning step to remove excess fats and meat, the trachea was cut into 1 cm pieces. Dehydration was carried out in an oven with air circulation for 7 hours at 68°C.

3. Snack filling

The filling was produced with mechanically separated meat (MSM, 37.0%), wheat flour (34.0%), water (ice, 16.5%), rice flour (5.0%) and animal fat (5.0%), sodium chloride (1.8%), bi-calcium tripolyphosphate (0.5%), ascorbic acid (0.2%), sodium nitrite (0.015%) and Butylated hydroxytoluene (BHT, 0.011%). The percentages were defined based on the human food legislation according to the Health Surveillance Agency and the Ministry of Agriculture, Livestock and Supply of Brazil [13].

4. Processing and cooking

The ingredients were separated and processed in a cutter according to the sequence:

1) beat the MSM made up entirely of chilled kids and half of the curing salt until they were broken up;
2) Add half of the ice (8.6%), beat again;
3) Place the fat and the rest of the spices and beat until the whole mixture is diluted;
4) Add the remaining ice and flour, mix well.

The finished filling was applied inside each trachea unit. The stuffed tracheas were sent to a preheated oven following the cooking times/temperature, as described in Table I.

<p>| TABLE I. Cooking Time Of Meat Snack (Trachea) |</p>
<table>
<thead>
<tr>
<th>Order</th>
<th>Time (min)</th>
<th>Temperature ºC</th>
</tr>
</thead>
<tbody>
<tr>
<td>1ª</td>
<td>30</td>
<td>55</td>
</tr>
<tr>
<td>2ª</td>
<td>45</td>
<td>65</td>
</tr>
<tr>
<td>3ª</td>
<td>45</td>
<td>75</td>
</tr>
<tr>
<td>4ª</td>
<td>15</td>
<td>85</td>
</tr>
<tr>
<td>5ª</td>
<td>15</td>
<td>95</td>
</tr>
<tr>
<td>6ª</td>
<td>30</td>
<td>105</td>
</tr>
<tr>
<td>7ª</td>
<td>30</td>
<td>115</td>
</tr>
</tbody>
</table>

D. Probiotic application – microencapsulation

Three methods of encapsulating S. boulardii were used. The performance of each test was based on the total viable
cells in standard yeast agar (CFU / mL) as well as already described [18].

Methodology 1. *S. boulardii* culture was grown in YEG broth: from a 72-hour culture at 25ºC in YEG broth under aerobicosis. An aliquot was placed in Eppendorf tubes for centrifugation (Quimis®, São Paulo) for 10 minutes, 4ºC, 7590 RCF. Then, the supernatant was discarded and the pellet washed with distilled water. The procedure was performed twice. To the final pellet, 500 µL of glycerine (Dinâmica® São Paulo) and 500 µL of 2% sodium alginate solution (Dinâmica® São Paulo) were added, homogenizing the mixture and transferring to a 1mL sterile syringe (26G1 needle/2 13x0.45). From 0.1M CaCl2 solution (Dinâmica®, São Paulo) the mixture was dripped using the extrusion technique, stirring occasionally, forming the microcapsules instantly which were left in contact with the solution for 10 minutes. The microcapsules were filtered on filter paper and kept in cryovials with a 1:1 solution of sterile distilled water and YEG broth at 4ºC. Methodology 2. *S. boulardii* lyophilized with chitosan coverage: from 1000 mg of lyophilized yeast powder 3.5mL of glycerine and 3.5 mL of 2% sodium alginate solution were added. The mixture was homogenized and passed to a 20mL sterile disposable syringe with an 18G1 40 x 1.2 hypodermic needle (Injex São Paulo), dripping in 0.1M CaCl2 by extrusion and stirring eventually, keeping the microcapsules in contact with the solution for 10 minutes, following for filtration. In watch glass with chitosan powder (Vetec, Rio de Janeiro), the microcapsules were transferred and slightly agitated to form the protective chitosan barrier, followed by storage in cryovials at 4ºC.

Methodology 3. In 1500 mg of lyophilized *S. boulardii* were added 3.5 mL of glycerine and 3.5 mL of 2% sodium alginate solution, homogenizing and transferring to a sterile 20 mL syringe with a 40X1 18G1 hypodermic needle. By using the extrusion technique, it was dripped into a 0.1 M CaCl2 solution, stirring occasionally for 10 minutes. Then, the capsules were filtered and stored in cryotubes according to methodology 1.

E. Viability of encapsulated yeast after action of gastric and intestinal juice

To simulate the action in gastrointestinal transit and to evaluate the survival of *S. boulardii* in relation to pH changes, the procedure was carried out based on the simulation method proposed by [58]: gastric juice (pH 2): NaCl 9g/L (Vetec, Rio de Janeiro), pepsin 3g/L (Vetec Rio de Janeiro). Intestinal juice (pH 8): NaCl 6.5g / L; 0.835 g/L KCl (Synth, São Paulo); 0.22 g/L CaCl2; NaHCO3 1.386 g/L (Vetec, Rio de Janeiro); pancreatic 1g/L; bile salts 3g / L (Himedia®, Brazil).

In a sterile plastic bag, 30 mL of gastric juice and three microcapsules were added, where it remained for 2 hours in a stomacher-type homogenizer with speed 1. In parallel, for each100 minutes aliquots of 100µL were removed and spread by spreading on plates of Sabouraud Dextrose agar (SD) (Acumedia®, USA). The incubation took place for 96 hours at 25ºC in aerobicosis. The microcapsules were also transferred to 30 mL of intestinal juice for 45 minutes, and at times 20 and 45 minutes, using the inoculum on SD agar plates, according to the previous process.

F. Microbiological evaluation and shelf-life

The methodologies applied for microbiological analysis were carried out according to the provisions of Normative Instruction SDA No. 62, 26 August 2003, of the Ministry of Agriculture, Livestock and Supply [14].

To verify and guarantee the safety and hygiene of the raw material and the final product, a microbiological evaluation was carried out during the shelf life (5 months). Table II shows the analysis carried out according to the parameters considering the legislation for similar products, based on two microbiological criteria: Safety and Hygiene criteria.

<table>
<thead>
<tr>
<th>TABLE II: MICROBIOLOGICAL PARAMETERS DEFINED FOR EVALUATION OF PET SNACK WITH PROBIOTIC Saccharomyces boulardii</th>
</tr>
</thead>
<tbody>
<tr>
<td>Criteria</td>
</tr>
<tr>
<td>Safety:</td>
</tr>
<tr>
<td>Clostridium (reducing sulfite) at 46ºC**</td>
</tr>
<tr>
<td>Staphylococcus aureus*</td>
</tr>
<tr>
<td>Hygiene:</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
</tbody>
</table>

The culture media used in the analysis included: Tetrathionate broth (Acumedia®, USA), Rappaport Vassiliadis broth (Acumedia®, USA) and Hektoen-Enteric agar (HE) (Himedia®, Brazil) for Salmonella sp., TSC agar for Clostridium reducing sulfite, Baird Parker for Staphylococcus aureus, Potato Dextrose (PDA) for molds and yeasts (Kasvi, Brazil), Plate Count agar for aerobic mesophilic or thermophiles (PCA), and YE agar with glucose (YEG) for S. boulardii. (Acumedia®, USA)

G. Physicochemical analysis

The guarantee levels for products intended for animal feed must comply with the technical regulations of identity and quality issued by the Ministry of Agriculture, Livestock and Supply [16]. Pet food must have the following guarantees on their labels or packaging: Moisture (maximum), Crude Protein (minimum), Etheric Extract (minimum), Fiber Matter (maximum), Mineral Matter (maximum), Calcium (maximum) and Phosphorus (minimum) [13]. The analysis included: moisture, crude protein, ether extract, fibrous matter, mineral matter [35], Calcium [11], and Phosphorus [11].

H. Palatability Analysis

The palatability analysis with dogs was approved by the Ethics Committee on the Use of Animals of the Academic institution where the study was carried out and has received the acceptance number protocol no. 67/2018. The animals included in this study were n=20 dogs, males and females, adults, healthy of medium and small breeds staying in a clinic and inn for dogs. As an inclusion factor in the institution where the study was carried out and has received the acceptance number protocol no. 67/2018. The animals were separated so that there was no interference among them; two meals a day were offered with a snack between them. The offer was repeated on a non-subsequent day for 7 days. The snack acceptance assessment was performed based on: if acceptance was immediate within 10 min; if the animals...
presented voluntary consumption; if there was total consumption in 2 hours; if there were leftovers, (regret); what were the reactions presented. All relevant information was recorded in the period (animal behavior, food dispute, environment, contact with owners).

The palatability of the food was assessed based on the acceptance methods Equation 2 was used to calculate the product’s Acceptability Index (AI):

\[
AI (\%) = \frac{A}{B} \times 100
\]

Where: A = average grade obtained for the product, and B = maximum grade given to the product [49]. AI with good repercussions has been considered ≥ 70% [26], [57].

III. RESULTS AND DISCUSSION

A. Probiotic Yeast Viability Tests Under Different Culture Conditions

The Table III shows that the best growth temperature for S. boulardii was 25°C. However, data from other studies state that the ideal temperature for growth is 37°C, which would be like body temperature [31], [24], [25].

S. boulardii behaved as an optional aerobic, growing normally both in the presence of oxygen and at low oxygen tension. Therefore, it can adapt according to the conditions of the environment [46].

In relation to the different concentrations of salt (NaCl) to which it was submitted, the microorganism developed better at a concentration of 1%, with an average cell density of 1.81 x 10^7 cells/mL (Table III), meaning that a high concentration of salt is able to interfere with its growth.

Considering the best growth conditions tested, it was found that at 25°C in aerobic conditions by cultivation on SD / PDA agar plates the yeast reached 1.52 x 10^7 CFU / 24 hours. It was also estimated that each colony formed on a YEG agar plate can generate an average of 2.03 x 10^8 cells/mL / 24 hours (Fig.1). Gravimetric analysis resulted in a final dry mass of 0.2623 g.

In relation to the pH there was a rapid multiplication of yeasts, reaching 6.78 x 10^7 cells. mL^-1 at pH 3.5 (Table III). A study by [27], presented very similar results when comparing a strain of Saccharomyces cerevisiae with S. boulardii, in which both were inoculated at different pH’s and, while S. cerevisiae showed a great decline in its development according to more acidic pH, S. boulardii remained in equal growth until reaching pH 2.

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### TABLE III: MAXIMUM LOGARITHMIC CYCLES REACHED BY Saccharomyces Boulardii AFTER CHALLENGE UNDER DIFFERENT CONDITIONS OF TEMPERATURE, pH AND SALINITY

<table>
<thead>
<tr>
<th>Condition</th>
<th>Log cycles/mL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature/48hours</td>
<td></td>
</tr>
<tr>
<td>Initial</td>
<td>2.88</td>
</tr>
<tr>
<td>25°C</td>
<td>7.75</td>
</tr>
<tr>
<td>28°C</td>
<td>7.62</td>
</tr>
<tr>
<td>35°C</td>
<td>8.64</td>
</tr>
<tr>
<td>37°C</td>
<td>7.63</td>
</tr>
<tr>
<td>Salinity/24hours</td>
<td>------------</td>
</tr>
<tr>
<td>1%</td>
<td>7.25</td>
</tr>
<tr>
<td>1.5%</td>
<td>7.23</td>
</tr>
<tr>
<td>2%</td>
<td>6.89</td>
</tr>
<tr>
<td>2.5%</td>
<td>6.70</td>
</tr>
<tr>
<td>3%</td>
<td>6.62</td>
</tr>
<tr>
<td>pH/72hours</td>
<td>------------</td>
</tr>
<tr>
<td>1.5</td>
<td>6.68</td>
</tr>
<tr>
<td>2</td>
<td>6.88</td>
</tr>
<tr>
<td>2.5</td>
<td>7.30</td>
</tr>
<tr>
<td>3</td>
<td>7.63</td>
</tr>
<tr>
<td>3.5</td>
<td>7.82</td>
</tr>
<tr>
<td>4</td>
<td>6.77</td>
</tr>
<tr>
<td>4.5</td>
<td>7.74</td>
</tr>
</tbody>
</table>

![Fig. 1. Growth curve of Saccharomyces boulardii at 25°C/ 24 hours of in Yeast Extract brotgh and Glucose under agitation at 200 rpm. Reading of optical density at 600nm and colonies forming units (CFU)/mL count by spread plate in YE agar plates after 72 hours/25°C.](http://dx.doi.org/10.24018/ejfood.2020.2.4.69)

B. Microencapsulation

In relation to microencapsulation methods, the methodology I by using culture in YEG broth was efficient, but dehydration occurred during the storage process. The microcapsules were reduced in size even when stored in distilled water and YEG broth (1: 1). A similar result of microcapsule deterioration was observed by [18] when using calcium alginate, where the microspheres deteriorated in about 30 days. Besides that, the process of obtaining microspheres with fresh culture is time-consuming and is not economically viable for industrial application.
For methodology 2, it was used lyophilized yeast culture surrounded by a layer of chitosan, being the most efficient technique in protection. Likewise, [23] performed microencapsulation of probiotic and prebiotic cultures in alginate and chitosan capsules and found that the combination of protected even more against some exogenous factors. Although according to [39], chitosan is widely used in the food industry due to its antifungal action, here in our study, contamination by fungi, despite the imposed aseptic conditions, made this methodology unfeasible. Thus, the quality and homogeneity of production inputs is an objective to be achieved.

In relation to the methodology 3, microencapsulation was performed with lyophilized culture of *S. boulardii* with a higher concentration, obtaining the best result in microcapsules for application in pet snack, by using only 2% sodium alginate and 0.1 M CaCl₂. The capsules, measuring 1.5 cm in diameter, proved to be inert during the storage period, showing no deterioration, besides they were free from contamination. Likewise, [37] performed the encapsulation of *Lactobacillus casei* with sodium alginate coating, claiming that alginate protection was highly efficient in acid pH tests, bile salts and heat treatment.

C. Evaluation Of Simulated Gastric And Intestinal Juice Action On Microencapsulated Saccharomyces Boulardii

The results indicated that although there was a release (loss) of cells during the process of passing through gastric juice (pH 2.0), corresponding to 2.54 Log cycles of colonies grown on SD agar plates (Fig. 2), with the intestinal juice the result was 5.92 Log cycles (Fig. 3).

![Fig. 2. Encapsulated Saccharomyces boulardii subjected to in vitro action of gastric juice (pH 2.0). Results based on yeast release and growth on Sabouraud Dextrose agar plates at 25°C/24 hours.](image1)

![Fig. 3. Encapsulated Saccharomyces boulardii subjected to in vitro action of intestinal juice (pH 8.0) right after the action of gastric juice at pH 2.0. Results based on release and growth on Sabouraud Dextrose agar plates at 25°C/24 hours.](image2)

Although the amount of microencapsulated yeasts resulting after passing through gastric juice was estimated to be just over 3 Log cycles, the final release of yeasts in the intestinal juice corresponded to 6 Log cycles. Therefore, the action of the intestinal juice allowed the total release yeasts.

Alginate when used as an encapsulating agent, especially sodium alginate, is insoluble in some organic solvents and in acidic media, with a pH below 3 [9]. However, in the present study, microencapsulated yeast cells were partially released during the action of gastric juice (pH 2.0), indicating that some change in the structure of the microcapsule occurred.

The survival of probiotic microorganisms during passage in the gastrointestinal system, resisting low pH, digestive enzymes, and bile salts, was maintained by the microencapsulation process, allowing the probiotic action of the microorganism, as recommended by the [28].

Therefore, it was estimated at about 4.18 x 10⁶ CFU the total viable colonies in the animal's intestine, taking into account the 15 microcapsules into snack pet with meat pork, however already considering a loss of 2.54 Log cycles during the process.

The shelf-life of the trachea snack with probiotic followed for 150 days. The pathogens *Salmonella* sp./25g and *Clostridium* sulphite reducing were not detected, so the product did not present a risk to the health of the animals (Table IV). In relation to the probiotic in the trachea snack, it resulted in 3.08 Log cycles in the last month, a reduction of approximately 3 Log cycles in the period of the final month. But, even with this loss it can still be considered viable and with probiotic potential. Given this result, the shelf life was defined at 120 days. *S. boulardii* presented an average of 5.62 Log cycles, so there is a safety margin in the product and a guarantee of benefits provided by the indicated amount of probiotic to be ingested. Ideal for probiotic products is an initial density of 10⁸ to 10⁹ CFU/day which must be ingest for functional effects [53]. However, lower values at the product are accepted if their effectiveness is proven [15].

Comparing the count of molds and yeasts obtained from the trachea with probiotic and that obtained from the trachea without probiotic, the result was 2 more Log cycles of difference in the first due to the presence of *S. boulardii*. Molds and yeasts are widely distributed in the environment and can be found in the air and in food, which are an important source of contamination.

The trachea snack used here is an edible by-product containing MSM-based filling. But, in handling this type of by-product, contamination by mesophilic bacteria, molds and yeasts is a risk to be considered. The presence of probiotic yeast in trachea snack can contribute to reduce that risk. In view of the possible susceptibilities of the immune system of dogs and cats to inflammatory bowel disease [22] the continuous use of this yeast helps in the better quality of life of the animal, which is directly linked to its health.
TABLE IV: MICROBIOPHICAL ANALYSIS OF TRACHEA SNACK ADDED OF Saccharomyces boulardii

<table>
<thead>
<tr>
<th>Time (days)</th>
<th>10</th>
<th>10</th>
<th>30</th>
<th>30</th>
<th>60</th>
<th>60</th>
<th>90</th>
<th>90</th>
<th>120</th>
<th>120</th>
<th>150</th>
</tr>
</thead>
<tbody>
<tr>
<td>Safety level</td>
<td>TP</td>
<td>TN</td>
<td>TP</td>
<td>TN</td>
<td>TP</td>
<td>TN</td>
<td>TP</td>
<td>TN</td>
<td>TP</td>
<td>TN</td>
<td>TP</td>
</tr>
<tr>
<td>Salmonella sp./25g</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Clostridium reducing sulfite at 46°C</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>

Hygiene criterion
Staphylococcus aureus (CFU/g) | (Log_{10}) 3.88 | 2.56 | 6.28 | 3.01 | 5.80 | 3.46 | 5.87 | 3.58 | 5.60 | 3.08 | 2.72 |
Molds and yeasts (CFU/g) | (Log_{10}) 5.62 | 4.60 | 4.70 | 4.65 | 4.68 | 4.18 | 4.70 | 5.05 | 4.71 | 0 | 0 |
Aerobic mesophiles (Log_{10} CFU/g) | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
Aerobic thermophiles (Log_{10} CFU/g) | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
S. boulardii (Log_{10} CFU) | 6.79 | No | 6.18 | No | 5.89 | No | 5.89 | No | 5.62 | No | 3.08 |

TP- Trachea with Probiotic; TN- Trachea without probiotic; No-Absence

D. Physical And Chemical Analysis

The trachea snack is a specific food, considered not complete in nutritional terms. In Table V the reference is made to complete foods according to the identity standards of the food products of the Ministry of Agriculture, livestock and supply in Brazil [13]. Therefore, in relation to pet food, they must present on their labels or packaging, at least, the following guarantees: (maximum); crude protein (minimum); ethereal extract (minimum); fibrous matter (maximum); mineral matter (maximum), Calcium (maximum) and Calcium (minimum), and Phosphorus (minimum). Thus, Table V compares the physical-chemical composition of the stuffed trachea without probiotic with the stuffed trachea with probiotic with reference to the guarantee levels for a complete food for adult dogs.

TABLE V: COMPARISON OF THE RESULTS OF THE GUARANTEE LEVELS FOR STUFFED TRACHEA SNACK WITHOUT PROBIOTIC Saccharomyces Boulardii AND SNACK WITH PROBIOTIC WITH THE IDENTITY STANDARDS OF FOOD PRODUCTS

<table>
<thead>
<tr>
<th>Dry food Warranty levels (%)</th>
<th>For adult dogs** Official standards</th>
<th>Stuffed trachea snack- no probiotic</th>
<th>Stuffed trachea snack with probiotic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Humidity (max.)</td>
<td>12</td>
<td>25.40</td>
<td>17.90</td>
</tr>
<tr>
<td>22/5000</td>
<td>16</td>
<td>36.10</td>
<td>57.50</td>
</tr>
<tr>
<td>Crude protein (min.)</td>
<td>4.5</td>
<td>6.55</td>
<td>11.44</td>
</tr>
<tr>
<td>Ethereal extract (min.)</td>
<td>6.5</td>
<td>0.11</td>
<td>0.78</td>
</tr>
<tr>
<td>Fibrous matter (max.)</td>
<td>12</td>
<td>3.13</td>
<td>2.56</td>
</tr>
<tr>
<td>Mineral material (max.)</td>
<td>2.4</td>
<td>1.56</td>
<td>1.83</td>
</tr>
<tr>
<td>Calcium (max.)</td>
<td>0.6</td>
<td>0.52</td>
<td>1.04</td>
</tr>
</tbody>
</table>

**[13]

The values presented are well accepted, including representing a final product with excellent protein value. The addition of salt concentration interferes with a small decrease in the tolerance of microorganisms, but still with good results. Therefore, in the comparison between stuffed trachea without probiotic and stuffed trachea with probiotic on the minimum level of phosphorus. The first one presented 0.08% below the limit for complete foods. For humidity both types of trachea snacks were above the limit for dry product as mentioned in the ordinance above. The Fig. 4 summarizes final data obtained for stuffed trachea snack with probiotic. Except for mineral material, the trachea snack with probiotic showed the highest levels for protein, ether extract, fibers, calcium, and phosphorus. This is since yeast is constituted on average by 48.2% total dietary fiber (43.7% insoluble fiber and 4.5% soluble fiber); 7.9% crude protein and 19.8% ethereal extract [43].

![Fig. 4. Percentual levels of mineral material, fibers, calcium, phosphorus and other in stuffed trachea snack with probiotic Saccharomyces boulardii.](image)

The yeasts are sources of unicellular protein and B vitamins, enzymes, volatile fatty acids, chelated minerals, among others, which provide better performance, greater resistance, and less stress to the animal [17]. The chemical composition depends on a series of factors, such as the nature of the substrate used, degree of aeration of the medium, yeast species, treatments imposed on the culture medium and concentration of salts, among others. However, for a product to play a role on health, it is not enough that it has advantageous nutritional characteristics. It must reach the consumer, be a product with a favorable taste with attractive properties [7], [34].
E. Palatability

When developing a new product, one of the fundamental points is the evaluation of its acceptability to predict the behavior towards the consumer market [45].

The palatability of the food is determined by the association of chemical and physical aspects, based on odor, texture, size, temperature, and flavor. Being able to relate the canine preferences with that of the owners, as well as the physical and social environment in which the dogs find themselves [51].

In the present study, acceptability was tested with 20 animals. Average scores were obtained by race (Fig. 5). Among the animals, the Poodle breed was the only one who disliked the product. Details about this breed provided by the veterinarian who accompanied the test indicated that it has a higher requirement in the selection of its foods compared to other breeds, which may be the reason for the low average.

![Fig. 5. Average degrees of reaction of dogs by the palatability test for trachea snack stuffed with meat and added of probiotic.](image)

According to [51] canine preferences can relate to that of the owners, as well as the physical and social environment in which the dogs find themselves. In addition, they consider other factors that can influence the perception of odours such as: race, sex, environment, and medication administration are the most important.

In the present study the average age of dogs was 5.4 years. Comparing between genders, the female had greater acceptance, with a difference of 2.2 points. Only one dog was on medication, but this factor had no influence on the animal response.

According to [57] and [26], for the product to be considered as accepted, in terms of its sensory properties, it is necessary that it obtains an Acceptability Index (AI) of at least 70%. Based on the grades for acceptability and AI calculation (equation 2), the product presented AI in 77.8% considered to be of good palatability.

Comparing with work involving the incorporation of encapsulated probiotic microorganisms for human food, [32] observed, in yogurt, that the calcium alginate capsules were visible and perceptible in the mouth when ingested, causing the product to be rejected by the tasters. However, for use in cereal bars, the incorporation of microorganisms occurred without them being able to be sensorially detected [5].

The palatability of the snack for dogs was found to be positive, as there was no leftover food offered to the taster dogs, characterizing that the presence of the microcapsules did not influence the feeding.

Based on the data obtained, it can be considered that the yeast with probiotic potential Saccharomyces boulardii developed its greatest growth at 25°C. The microencapsulation methodology adapted for lyophilized yeast (methodology 3) was more efficient, avoiding the greatest loss of cells. In addition, the microcapsules remained viable during the simulation of the action in the simulated gastrointestinal transit, with greater release of cells in the intestinal juice, this will allow their effectiveness (survival and viability) in real conditions.

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