Valorization of okara oil for the encapsulation of *Lactobacillus plantarum*

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**ABSTRACT**

Oil-in-water (O/W) emulsions of okara oil-caseinate (1:2; 1:3 and 1:4 O/W ratios) were used to encapsulate *Lactobacillus plantarum* CIDCA 83114. Once encapsulated, microorganisms were freeze-dried or spray-dried, and observed by scanning electronic and confocal microscopies. A physical characterization of the dehydrated capsules was carried out by determining their moisture content, water activity, particle size, polydispersity index and zeta potential. Determining the induction times and peroxide values provided information about their susceptibility to oxidation. In turn, bacterial stability was analyzed by plate counting before and after freeze-drying and spray-drying, and during storage at 4 °C.

Spray-dried emulsions had lower Z-sizes and polydispersity indexes, higher induction times and lower peroxide values than the freeze-dried ones, thus resulting better systems to protect *L. plantarum* CIDCA 83114. In addition, the culturability of spray-dried bacteria did not decrease neither after spray-drying nor up to 60 days of storage at 4 °C.

The results showed that the better physical-chemical stability of spray-dried capsules determined the greater stability of microorganisms. This demonstrates the importance of defining adequate emulsions’ formulations for an efficient encapsulation of microorganisms, with promising applications in the development of novel functional foods.

1. Introduction

Okara is the by-product remaining from the soy milk production, after filtration of the smashed soybeans seeds (Stanoevic, Barac, Pesic, Jankovic, & Vucelic-Radovic, 2013). It is obtained in large quantities (about 1.1 kg per kilogram of soybean processed for soymilk production), thus posing an important disposal problem (Li et al., 2012). To avoid such problem, great efforts have been employed to use okara for the formulation of different products (Vong & Liu, 2016).

Dehydrated okara contains about 9–10% humidity, 21% proteins, 55% whole fiber, 1.5% ash and 13–14% fats and oils (Quintana, Gerbino, & Gomez-Zavaglia, 2017). Because of the nutritional value of its components (proteins of high nutritional value, dietary fiber, anti-oxidants, unsaturated lipids), okara appears as a valuable source of different ingredients for the formulation of functional foods (Vong & Liu, 2016). Defatted okara can be used as primary ingredient for the formulation of dry breakfast cereals or as meat extender (Shurtleff & Aoyagi, 2000). Okara oil has potential applications in cosmetic, pharmaceutical and food industry. In this regard, using super-critical CO₂ enables the recovery of nutritionally valuable compounds from okara oil, namely polyunsaturated fatty acids, phytosterols and phenolic compounds, such as soy isoflavones, genistein and daidzein (Borhan, Gani, & Shamsuddin, 2014; Quitain, Oro, Shunsaku, & Moriyoshi, 2006). This represents an added value in preventing chronic diseases (FAO, 2010) and opens opportunities for the development of novel applications aiming at using this by-product.

Lactic acid bacteria have an important role in the food and pharmaceutical industries, as they are extensively used as starters in the development of food and probiotic products. However, the decrease of water activity occurring during preservation and technological processes is often detrimental, leading to damages on the cellular structures or death (Santos, Gerbino, Araujo-Andrade, Tymczyszyn, & Gomez-Zavaglia, 2014; Tymczyszyn et al., 2008). To prevent these problems, bacterial microencapsulation into polymer matrices appears as an adequate strategy to ensure mechanical integrity during production processes (exposure to high/low temperatures, oxidation, shear, etc.), desiccation and storage (packaging and environment conditions, including moisture, oxygen, temperature, etc.) (Chávarri, Marañón, & Villarán, 2012). Encapsulation shell agents include a variety of polymers, carbohydrates, fats and waxes, depending on the material to be encapsulated.
protected. The coating of sealed capsules must be semipermeable and mechanically resistant to the adverse conditions above mentioned (Chávarri et al., 2012).

Microorganisms are usually suspended in a given encapsulating agent, forming emulsions or suspensions. Such emulsions are then dehydrated, generally by freeze-drying or spray-drying (Petrovic, Nedovic, Dimitrijevic-Brankovic, Bugarski, & Lacroix, 2007). Entrapment of probiotic bacteria in emulsion droplets has been suggested to stabilize different species of lactic acid bacteria (Hou, Lin, Wang, & Tzen, 2003; Pimentel-González, Campos-Montiel, Lobato-Calleros, Pedroza-Islas, & Vernon-Carter, 2009; Rodríguez-Huezo et al., 2014; Zhang, Lin, & Zhong, 2015). In this regard, Hou et al. (2003) succeeded at increasing about 10^4 times the intestinal survival rate of Lactobacillus delbrueckii ssp. bulgaricus by entrapping microorganisms in the droplets of reconstituted sesame oil body emulsions. Pimentel-González et al. (2009) successfully encapsulated Lactobacillus rhamnosus using water-in-oil-in-water emulsions prepared with canola oil and sweet whey (Pimentel-González et al., 2009). Milk fat was also used for encapsulation of lactic acid bacteria. In addition, preparing emulsion droplets with multiple lipid-protein-pectin layers provides additional protection for Lactobacillus salivarius strains (Zhang et al., 2015). Canola oil was also used to encapsulate Lactobacillus plantarum in double emulsions containing aguamiel or sweet whey as inner aqueous phases, and the obtained emulsions were successfully incorporated during cheese manufacture (Rodríguez-Huezo et al., 2014). The presence of whey proteins, insulins or fructo-oligosaccharides are reported to play an important role in the protection of the encapsulated microorganisms. Although emulsification technologies have demonstrated a great efficiency for the encapsulation of probiotics, there is still a long way to undergo in the development of more efficient entrapment systems. In this sense, when analyzing the recovery of entrapped bacteria, the focus has been always put on the composition of the aqueous components of emulsions. In fact, for improving bacterial recovery, different aqueous components have been assayed to formulate emulsions (Hou et al., 2003; Pimentel-González et al., 2009; Rodríguez-Huezo et al., 2014; Zhang et al., 2015). However, little attention has been paid to the oil components. This aspect is especially important because the chemical and physical properties of such oils determine their ability to form small droplets, leading to stable emulsions after homogenization (Bai & McClements, 2016). Lipid oxidation is another factor defining the stability of the emulsions, as it usually limits the shelf-life of fat containing products (Jacobsen, 2010). For this reason, an adequate selection of oils is very important to prevent oxidation during storage, thus contributing for the long term stability of encapsulated microorganisms.

The aim of this work was to formulate different okara oil-caseinate emulsions and use them to encapsulate L. plantarum CIDCA 83114, a strain with demonstrated inhibitory properties against E. coli O157:H7, Shigella and Salmonella (Hugo, Kakisu, De Antoni, & Pérez, 2008; Golowczyc, Silva, Teixeira, De Antoni, & Abraham, 2011; Kakisu, Abraham, Tironi Farinati, Ibarra, & De Antoni, 2013; Kakisu, Bolla, Abraham, de Urraza, & De Antoni, 2013). The O/W emulsions were freeze-dried and spray-dried, and the obtained particles were characterized by determining their size, zeta-potential (ζ-potential) and polydispersity index (Đ). The stability of oil to oxidation was assessed by determining the induction time and peroxide values. In turn, the bacterial stability was evaluated by plate counting before and immediately after encapsulation, and during 90 days of storage at 4 °C.
2. Materials and methods

2.1. Materials

Okara, obtained from Soyana S.H. (San Martín, Argentina); sulphuric acid, methanol, chloroform, acetone, butanol, isooctane and isopropanol (Merck, Darmstadt, Germany); sodium thiocyanate, ferrous chloride and cumene hydroperoxide (Sigma-Aldrich, St. Louis, MO, USA); KH₂PO₄ (J. T. Baker, Mexico); NaCl (Sigma-Aldrich, USA); Na₂HPO₄ (Anedra, Argentina); sodium caseinate (Sigma Chemical, St. Louis, MO, USA); non-fat milk (Fisher Scientific, MA, USA); TO-PRO-3 (Sigma-Aldrich, St. Louis, MO, USA); Nile red (Sigma-Aldrich, St. Louis, MO, USA).

2.2. Methods

The experimental sequence of experiments is shown in Scheme 1.

2.2.1. Obtaining and characterization of okara oil

After being received, okara was centrifuged 5 times to remove the excess of water. The sediment was frozen at ~ −80 °C for 48 h and then, freeze-dried on a Heto FD4 equipment (Heto Lab Equipment, Denmark) for 48 h (temperature of condenser: −50 °C; chamber pressure: 0.04 mbar). The okara oil was extracted with diethyl ether solvent in a Soxhlet system (AOAC, 1995).

The fatty acid composition of the oil was analyzed according to IUPAC 2.302 standard method (IUPAC, 1992). The fatty acids methyl esters were prepared by adding 2 mL sulphuric acid (20 g/L in methanol) to 3 mg oils, heating at 60 °C for 2 h, extracting the esters with 1 mL chloroform-water (2:0.7 v/v) and washing twice with 0.7 mL ethanol) to 3 mg oils, heating at 60 °C for 2 h, extracting the esters with 1 mL chloroform-water (2:0.7 v/v) and washing twice with 0.7 mL water. The obtained fatty acids methyl esters were analyzed on a gas chromatograph interfaced with a mass spectrometer detector (Shimadzu QP 5050A, Tokyo, Japan) using capillary column ZB-5 (30 m × 0.25 mm). The conditions of analysis were: injection temperature 250 °C, detector temperature 280 °C and column temperature (30 m × 0.25 mm). The conditions of analysis were: injection temperature 250 °C, detector temperature 280 °C and column temperature 250 °C, detector temperature 280 °C and column temperature (30 m × 0.25 mm). The conditions of analysis were: injection temperature 250 °C, detector temperature 280 °C and column temperature (30 m × 0.25 mm). The conditions of analysis were: injection temperature 250 °C, detector temperature 280 °C and column temperature 250 °C, detector temperature 280 °C and column temperature (30 m × 0.25 mm). The conditions of analysis were: injection temperature 250 °C, detector temperature 280 °C and column temperature (30 m × 0.25 mm). The conditions of analysis were: injection temperature 250 °C, detector temperature 280 °C and column temperature (30 m × 0.25 mm). The conditions of analysis were: injection temperature 250 °C, detector temperature 280 °C and column temperature (30 m × 0.25 mm). The conditions of analysis were: injection temperature 250 °C, detector temperature 280 °C and column temperature (30 m × 0.25 mm). The conditions of analysis were: injection temperature 250 °C, detector temperature 280 °C and column temperature (30 m × 0.25 mm). The conditions of analysis were: injection temperature 250 °C, detector temperature 280 °C and column temperature (30 m × 0.25 mm). The conditions of analysis were: injection temperature 250 °C, detector temperature 280 °C and column temperature (30 m × 0.25 mm). The conditions of analysis were: injection temperature 250 °C, detector temperature 280 °C and column temperature (30 m × 0.25 mm).

2.2.2. Bacterial strain and growth conditions

L. plantarum CIDCA 83114 was isolated from kefir grains (Garrote et al., 2001) and maintained frozen at ~ −80 °C in 120 g/L non-fat milk solids. The strain belongs to the CIDCA culture collection and was identified by molecular methods, as previously published (Garrote et al., 2001; Golowczyc et al., 2008).

Microorganisms were cultured twice in MRS broth (de Man, Rogosa, & Sharpe, 1960) at 37 °C in aerobic conditions. Cultures in the stationary phase (~1 × 10¹³ CFU/mL) were harvested by centrifugation at 8000 rpm for 10 min, washed twice with phosphate saline buffer (PBS, KH₂PO₄ 0.144 g/L, NaCl 9 g/L, Na₂HPO₄ 0.795 g/L, pH 7.2) and suspended in the same volume of 6% w/v sodium caseinate.

2.2.3. Preparation of oil-in-water (O/W) emulsions

Microencapsulation was performed using an oil-in-water (O/W) emulsion system containing different concentrations of okara oil (1.5, 2.0, and 3.0% w/v), and a constant concentration of sodium caseinate (6% w/v) (Dianawati, Mishra, & Shah, 2016). This way the O/W ratios remained 1:4, 1:3 and 1:2, respectively. Okara oil and caseinate mixtures were then emulsified using an Ultra Turrax T25 high-shear probe mixer (IKA, Staufen im Breisgau, Germany) for 5 min at 13500 rpm and 20 °C. O/W emulsions (1:4 ratio) in which no microorganisms were added to the aqueous phase were used as controls.

2.2.4. Freeze-drying, spray-drying and storage

100 mL of O/W emulsions containing bacteria in the stationary phase were frozen at ~ −80 °C for 12 h, and freeze-dried in a Rificor L-A-B4-C (Buenos Aires, Argentina), operating with the condenser at ~ −45 °C in a chamber pressure of 0.04 mbar. The freeze-drying process lasted for 48 h. Non-encapsulated bacteria suspended in PBS, and 1:4 O/W freeze-dried emulsions not containing microorganisms were used as controls.

In parallel assays, 500 mL O/W emulsions containing microorganisms in the stationary phase were spray-dried in a laboratory-scale spray-dryer (Büchi B290 mini spray-dryer, Flawil, Switzerland) at a constant air inlet temperature of 145 °C and an outlet temperature of 60 °C. Controls: microorganisms grown in MRS broth harvested, neutralized and resuspended in PBS (non-encapsulated microorganisms) and spray-dried O/W emulsions not containing microorganisms.

Before and after freeze-drying or spray-drying, microorganisms were serially diluted in PBS, plated on MRS agar and incubated at 37 °C in aerobic conditions for 48 h. Plate counts were expressed as log CFU/g.

Dehydrated bacteria (both freeze-dried and spray-dried) were stored for 90 days at 4 °C in silica gel containing recipients, and plate counted every 15 days. Plate counts were expressed as log N₀Nₚ where N₀ was the plate count at a time 0 and Nₚ the plate count at each time of storage.

2.2.5. Microscopic observations

2.2.5.1. Scanning electronic microscopy. Dehydrated samples were frozen in liquid nitrogen and fractured using a cold scalpel blade, and examined with a FEI model Quanta 200 electron microscope (The Netherlands). Samples were mounted onto bronze stubs by using a double-sided tape and examined without any metal or carbon coating at low pressure and an acceleration voltage of 12.5 kV.

2.2.5.2. Confocal laser scanning microscopy analysis. Two dyes were used for confocal laser scanning microscopy: TO-PRO-3 and Nile red. Microorganisms were visualized using TO-PRO-3, which is an excellent far red-fluorescent nuclear and chromosome counterstain. The dye exhibits far-red fluorescence with excitation at 642 nm and emission at 661 nm. In turn, Nile red was used to stain the O/W emulsions. A 0.01% w/v Nile red staining solution was prepared by dissolving 0.001 g of the dye in 10 mL of acetone. The solution was stored in the dark at 20 °C. Nile red dye exhibits green fluorescence with excitation at 510–560 nm and emission at 590 nm.

Freeze-dried and spray-dried emulsions were dispersed in 10 mL distilled deionized water under constant magnetic stirring to obtain a concentration of ~10¹⁰ cells/mL. Then, 1% v/v of Nile red was added and samples were incubated for 1 h at 37 °C. Afterwards, an equal volume of 50% v/v methanol was incorporated and samples were incubated at 37 °C for 5 min. Finally 1 μL of TO-PRO-3 (final concentration: 0.8 μM) was added and samples were incubated at 37 °C for 5 min. All samples were observed in a confocal laser-scanning microscope (Leica TCS SP5 Leica Microsystems, Wetzlar, Germany), using an excitation wavelength of 642 nm. Emulsions were observed with no observation wavelength.

2.2.6. Characterization of dehydrated emulsions

2.2.6.1. Moisture content. Moisture content of the freeze-dried and spray-dried samples were determined by measuring their weight loss upon drying in a vacuum oven at 70 °C until constant weight (AOAC, 1980). Dehydrated emulsions without microorganisms were used as controls. Moisture results were expressed as grams of water per 100 g of dried sample (ds).

2.2.6.2. Water activity (aw). The aw of freeze-dried and spray-dried samples was determined using an Aqualab water activity instrument (Aqualab, Model Series 3TE, USA). Capsules not containing bacteria were used as controls. The equipment was calibrated using standard salt.
solutions provided by the manufacturer.

2.2.6.3. Particle size, polydispersity index (D) and zeta potential (ζ). All measurements were recorded at 37 °C on freshly prepared emulsions as well as after freeze-drying and spray-drying, using a Malvern Instrument Zetasizer Nano-Z (Malvern Instruments, Malvern, UK). For all measurements, samples were dispersed in 0.15 M NaCl (pH 7; 37 °C, 30 min) under constant magnetic stirring. They were then appropriately diluted to obtain about 10^5 particles/mL and vortexed for 30 s before determinations.

The average hydrodynamic particle size (Z-average) and D were determined by using dynamic light scattering at backward scattering (173°) with the Zetasizer 6.20 software. ζ-potential was determined using a combination of measurement techniques: electrophoresis and laser Doppler velocimetry (Laser Doppler Electrophoresis). The ζ-potential values were provided directly by the instrument. An average value of ζ-potential was obtained from at least 20 determinations for each sample.

2.2.6.4. Induction time. The induction time, lag period during which oils show stability to oxidation, for both freeze-dried and spray-dried emulsions was determined according to AOAC Cd12-57, using a Rancimat 743 equipment (Metrohm AG, Herisau, Switzerland). Emulsions were determined according to AOAC Cd12-57, using a Rancimat 743 equipment (Metrohm AG, Herisau, Switzerland). Each sample was prepared with the oil at an amount of 0.2% (w/w) of the total emulsion. The induction time was determined using a combination of measurement techniques: electrophoresis and laser Doppler velocimetry (Laser Doppler Electrophoresis). The ζ-potential values were provided directly by the instrument. An average value of ζ-potential was obtained from at least 20 determinations for each sample.

2.2.6.5. Peroxide determination. Lipid oxidation was evaluated by determining the peroxide value. The oil was extracted according to Partanen, Hakala, Sjövall, Kallio, and Forssell (2005). Briefly, 0.5 g of dehydrated samples containing microorganisms were suspended in 5 mL of distilled water and shaken until complete dissolution. 300 μL of suspensions were taken and vortexed 3 times for 10 s with 1.5 mL of an isooctane/isopropanol (3:1 v/v) mixture. The phases were separated. Peroxide values were determined on the upper phase according to McClements (1999). About 10–100 μL of the extraction medium were added to 2.8 mL of methanol/butanol (2:1 v/v), 15 μL of 3.94 M thiocyanate solution and 15 μL of 0.072 M acidic solution of ferrous iron. The samples were briefly vortexed, left 20 min in the dark, and the absorbance at 510 nm was read in a spectrophotometer (Shimadzu, Kyoto, Japan). Standard curves were carried out using cumene hydroperoxide. Capsules not containing microorganisms (O/W: 1:4) were used as controls.

2.2.7. Statistical analysis

All the assays were conducted in triplicate and in three independent assays. Average values were used for data analysis. Analysis of variance (ANOVA) was carried out for all the assays, using the program GraphPad Prism 5 (GraphPad Software Inc., San Diego, CA, 2007). Analysis of variance (ANOVA) was carried out using the statistical program STATISTIX 8 Software (Analytical Software, Tallahassee, FL, USA). Means were compared by two-sided Dunnett’s multiple comparison test and the difference was considered significant when \( p < 0.05 \).

3. Results and discussion

3.1. Characterization of okara oil

Okara constituents are valuable ingredients for the food and/or pharmaceutical industries. In this work, okara oil was used for the formulation of stable O/W emulsions aiming at protecting L. plantarum CIDCA 83114 during both dehydration and storage. The efficiency of such encapsulation strategy requires an integrative analysis considering not only the bacterial culturability but also the oil stability.

Table 1 shows the fatty acid composition of okara oil. C18:2 (n – 6) and C18:1 (n – 9) were the main fatty acids, accounting 74.65% of the total. The contribution of polyunsaturated acids (C18:2 and C18:3) was 62.58%.

3.2. Microscopic observations

Fig. 1 displays the SEM images of freeze-dried and spray-dried emulsions. Freeze-dried samples appeared as isolated particles not uniformly distributed (Fig. 1AI). On the contrary, spray-dried ones occurred as greater amounts of smaller and regularly dispersed particles (Fig. 1BI). Confocal laser microscopy enabled the observation of microorganisms stained with TO-PRO-3 (Fig. 1AII and BII, left panels). Nile red stain enabled the observation of the capsules (Fig. 1AII and BII, middle panels). The overlap of both images allowed the observation of both capsules (green) and microorganisms (red) (Fig. 1AII and BII, right panels). The dimensions of the spray-dried capsules containing microorganisms were comparable to those of other microorganisms encapsulated using other emulsions (1000–2000 nm for microorganisms and 5000 nm for emulsions), such as skim milk or acacia gum (Zhang et al., 2015), which may be important at a technological level for their incorporation in food products.

3.3. Characterization of the emulsions

The moisture content of freeze-dried and spray-dried samples was within 1.50 and 2.60 g/100 g ds, and emulsions not containing microorganisms (controls) showed significantly lower values (0.50–1.00 g/100 g ds) (Table 2). In turn, \( n_m \) was significantly lower for freeze-dried samples than for spray-dried ones and no differences for the different O/W ratios were observed (Table 2).

Table 2 shows the Z-size and ζ-potential of O/W emulsions before and after freeze-drying and spray-drying. Fresh emulsions showed the lowest Z-sizes and no significant differences related with the O/W ratios were observed (Fig. 2A). The Z-size of spray-dried samples increased up to 1.7 times with regard to that of fresh emulsions, with no significant differences related with the O/W ratios (Fig. 2A). For spray-dried samples the obtained D was lower than 0.300, indicating the homogeneity of the population (Table S1). On the contrary, freeze-dried samples were those showing the largest Z-sizes (Fig. 2A) and the greatest D (above 0.550, Table S1), indicating a greater heterogeneity. In this case, the 1:2 O/W freeze-dried emulsions were the largest ones and most heterogeneous, followed by 1:3 O/W and 1:4 O/W, thus demonstrating that the higher the oil content, the larger the Z-size and the greater the heterogeneity (Fig. 2A and Table S1).

When intended to encapsulate bioactive compounds for food applications, the size of the beads is critical and may determine their applications. For example, large particles can provide sandy textures, affecting the sensory properties of the products and thus, limiting their application in food products (Walstra, Wouters, & Geurts, 2005). Therefore, Z-sizes must be as low as possible. In this work, the Z-sizes of dehydrated emulsions were higher than those of the fresh ones (Fig. 2).

This can be explained considering that some aggregation can occur

| Table 1 |
| --- |
| Fatty acid composition of okara oil (% of total fatty acids). |
| Fatty acid | Okara oil |
| C16:0 | 11.57 |
| C16:1 | 0.08 |
| C18:0 | 4.43 |
| C18:1 (n – 9) | 20.04 |
| C18:1 (n – 7) | 1.30 |
| C18:2 (n – 6) | 54.61 |
| C18:3 (n – 3) | 7.97 |

* Standard deviation below 1% in all cases.
upon dehydration. In addition, the type of dehydration process strongly determined the stability of the obtained beads in terms of Z-size and Đ (Fig. 2 and Table S1). Hence, spray-dried emulsions had lower Z-sizes and Đ values than freeze-dried ones. The SEM and confocal images were consistent with these results, showing a homogeneous distribution of spray-dried particles, which was not observed among freeze-dried beads (Fig. 1). The nature of the spray-drying process explains the lower Z-sizes and Đ of the spray-dried emulsions in comparison with the freeze-dried ones. Drying in contact with a hot drying medium enables the obtaining of small droplets and size distributions during the atomization step (Sander & Penovic, 2014). Moreover, the smaller Z-sizes of spray-dried emulsions lead to greater surface contact, which in turn, can better interact with water. This explains the higher $a_w$ of spray-dried emulsions in comparison with the freeze-dried ones (Table 2).

Regarding the ζ-potential, no significant differences arising from the processes (freeze-drying or spray-drying) were observed (Fig. 2B). However, some differences arising from the O/W ratios were detected (Fig. 2). 1:2 O/W were those with the most negative values, followed by 1:3 and 1:4 O/W. In this latter case, ζ-potentials of the freeze-dried emulsions were within $-34$ and $-36$ mV. These results indicated that the higher the oil content, the more negative the superficial charge (Fig. 2B). In addition, controls without microorganisms, prepared with 1:4 O/W were significantly more negative than their counterparts containing bacteria.
3.4. Stability of the emulsions

Fig. 3 shows the stability of freeze-dried and spray-dried emulsions as determined by the induction time and the peroxide values. Freeze-dried samples had induction times within 6 and 13 h, with no significant differences related with the O/W ratios (Fig. 3A). On the contrary, spray-dried ones showed much larger induction times. They were about 30 h for 1:2 and 1:3 O/W, and about 80, for 1:4 O/W and for the control (also prepared with the same O/W ratio). In regard to peroxidation, spray-dried samples showed significantly lower peroxide values than freeze-dried samples (p < 0.05) (Fig. 3B), which reinforce their higher stability.

The fatty acid composition of okara oil could explain its stability in the desiccated emulsions (Fig. 3, Table 1). In fact, canola and sesame oils, as well as milk fats have been used to encapsulate different species of lactic acid bacteria (Hou et al., 2003; Pimentel-González et al., 2009; Rodríguez-Huezo et al., 2014; Zhang et al., 2015), but they often require double emulsification strategies and/or deposition of additional aqueous phases to correctly shell the obtained beads, and thus protect

| Samples            | Moisture (g/100 g ds) | a w     |
|--------------------|-----------------------|---------|
| Freeze-drying      |                       |         |
| O/W 1:2            | 1.51 ± 0.10          | 0.13 ± 0.04a |
| O/W 1:3            | 1.60 ± 0.13          | 0.14 ± 0.03a |
| O/W 1:4            | 2.26 ± 0.09          | 0.17 ± 0.05b |
| O/W 1:4 (control)  | 0.50 ± 0.02          | 0.24 ± 0.07ab |
| Spray-drying       |                       |         |
| O/W 1:2            | 1.90 ± 0.10          | 0.30 ± 0.05A |
| O/W 1:3            | 2.59 ± 0.19          | 0.32 ± 0.05A |
| O/W 1:4            | 2.14 ± 0.15          | 0.32 ± 0.05A |
| O/W 1:4 (control)  | 0.96 ± 0.09          | 0.27 ± 0.01A |

Different letters indicate significant differences (p < 0.05).
microorganisms. Canola oil has 57% of monounsaturated and 21% of polyunsaturated fatty acids (Matthaus, Özcan, & Al Juhaimi, 2016). In turn, sesame oil includes 39% of monounsaturated, 46% of polyunsaturated, and 13% of saturated fatty acids (Nzikou et al., 2009). It must also be considered that milk fat includes higher amounts of shorter fatty acids and lower amounts of mono and polyunsaturated ones (Månsson, 2008). In contrast, okara oil had a high contribution of polyunsaturated fatty acids, in particular, C18:2 (Table 1). In addition, the total content of polyunsaturated fatty acids (18:2 + 18:3) reached 62.58%, and that of C18:1, 21.34% (Table 1). The presence of higher amounts of polyunsaturated fatty acids would explain the stability of okara oil emulsions (Fig. 3). As the droplets in an O/W emulsion are usually liquid because both the oil and water phases must be in a fluid state when emulsions are prepared (temperature ~ 20 °C), polyunsaturated fatty acids (with lower melting temperatures), appear as more adequate ones to form more stable emulsions, avoiding crystallization processes (Bai & McClements, 2016).

In spite of these advantages, the presence of high concentrations of polyunsaturated fatty acids could make emulsions more susceptible to oxidation. The induction times and peroxide values of dehydrated emulsions (in particular those obtained by spray-drying 1:4 O/W emulsions) were much higher and lower, respectively (Fig. 3), than those of other edible oils (Bae & Lee, 2008; Binski et al., 2017; Ixtaina, Julio, Wagner, Nolasco, & Tomás, 2015; Pereyra Boué, Costamagna, Rodríguez, Speltini, & Coppo, 2009). This behavior can be explained considering the presence of antioxidant polyphenolic compounds (genistein, daidzein) in okara oil (Quittain et al., 2006).

Two comparisons could explain the high induction times for spray-dried 1:4 O/W emulsions: a-comparing the dehydration process (freeze-drying vs spray-drying), and b-comparing the O/W ratios.

If consider that freeze-dried samples were larger and less homogeneous than spray-dried ones (Table S1 and Fig. 1), they expose a larger surface to the air, thus being more prompt to oxidation. In this regard, Miyagawa and Adachi (2017) showed demonstrated that a reduction of the oil droplet size suppresses or retards autoxidation. Microencapsulated lipids or oils exhibit distinct features in the oxidation process. In addition, a reduction in the oil droplet size is also effective for suppressing or retarding the oxidation of microencapsulated oils.

Regarding the O/W ratios, spray-dried emulsions containing 1:4 O/W ratios (samples containing bacteria and controls) had much higher induction times. This observation can be explained considering that higher amounts of proteins better stabilize O/W emulsions. This observation has already been reported for other emulsions (Bai & McClements, 2016; Evans, Ratcliffe, & Williams, 2013; Miyagawa & Adachi, 2017).

3.5. Stability of the encapsulated microorganisms

Fig. 4 shows the culturability of L. plantarum CIDCA 83114 after freeze-drying and spray-drying. No significant decrease of culturability was observed after spray-drying, regardless the O/W ratio. On the contrary, freeze-drying was less efficient, leading to significant decreases of culturability (p < 0.05). In addition, the culturability within this group showed to be dependent on the O/W ratio. In fact, using 1:2 O/W was the worst condition, as culturability decreased from 13.91 ± 0.73 log CFU/g to 9.40 ± 0.31 log CFU/g. 1:4 O/W resulted a better condition, as a drop of 1.53 ± 0.08 log CFU/g was observed (Fig. 4). This indicates that the higher the oil content in the emulsion, the lower the bacteria culturability. Although bacterial culturability was above those required for probiotic products (Aquilina et al., 2013; Hill et al., 2014; Tripathi & Giri, 2014), the physical properties of the freeze-dried capsules (especially high D indexes and Z-sizes) make them unappropriated to encapsulate microorganisms. On the contrary, spray-dried emulsions looked as more homogenous (lower D values) and non-sticky powders with lower Z-sizes, larger induction times and lower peroxide values (Figs. 2, 3; Tables 2, S1). Thanks to these properties no
significant decrease of culturability neither during the process nor during storage (Figs. 4, 5), was observed, thus making this system more appropriate to encapsulate *L. plantarum* CIDCA 83114.

Storage also showed a differential behavior for freeze-dried and spray-dried microorganisms (Fig. 5). 1:3 and 1:4 O/W showed a similar behavior for freeze-dried samples, with no significant decrease of culturability up to day 45 (Fig. 5A). The drop of culturability started at day 60, and after 90 days of storage, it decreased about 4 log for both O/W emulsions. The culturability of bacteria freeze-dried in 1:2 O/W emulsions started decreasing already at day 15, and dropped 5.00 ± 0.09 after 90 days of storage (Fig. 5A). On the contrary, spray-dried emulsions were much better encapsulation systems also during storage, as no significant decrease of culturability was observed up to day 60 in none of the conditions assayed (Fig. 5B). Furthermore, the drop of culturability observed after 90 days of storage was not > 1.89 ± 0.87 in none of the cases. If considering that spray-drying is a cost effective technique, easily scalable and with low energetic consumption, the higher culturability of spray-dried microorganisms [consistent with the higher physical-chemical stability of spray-dried emulsions (Fig. 3)] appears as a very promising result.

As a whole, the results obtained in this work clearly showed that a greater physical-chemical stability leads to a greater stability for the encapsulated microorganisms, thus demonstrating the importance of defining adequate formulations for the emulsions aiming at efficiently encapsulate microorganisms.

4. Conclusion

Okara components are nutritionally valuable ingredients for food and/or pharmaceutical industries. Therefore, attempts to incorporate them in the formulation of functional foods is matter of increasing research work. One of the main achievements of this work was to use okara oil in the formulation of stable emulsions. The obtained results demonstrated that the adequate formulation of emulsions results in adequate materials to safely entrap lactic acid bacteria.

Taking into account that okara oil is a by-product and spray-drying is a cost-effective and easily scalable process, with low energetic consumption, the production of spray-dried emulsions entrapping *L. plantarum* CIDCA 83114 appears as an adequate alternative for the production of starters at a large scale. Moreover, the composition of okara oil, including polysaturated fatty acids and antioxidants, provides an additional source of high quality nutrients as vehicles of potentially probiotic microorganisms, whose importance in human health is out of question. Taking this into account, the results support the use of okara oils as suitable ingredients to formulate stable emulsions, with promising applications in the development of novel functional foods.

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Competing interests

The authors declare that they have no competing interests.

Author's contributions

G.Q. did the experimental work. E.G. conceived the experimental design and participated in the experimental work. A.G.-Z. analyzed the results, coordinated discussions and wrote the manuscript. All authors approved the final version of the manuscript.

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