- 1 Title: Omega–3 encapsulation by PGSS-drying and conventional
- 2 drying methods. Particle characterization and oxidative stability
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15 Abstract

Particles from Gas-Saturated Solutions (PGSS)-drying has been used as a green 16 alternative to encapsulate omega-3 polyunsaturated fatty acids (n-3 PUFAs) at mild, 17 non-oxidative conditions. PGSS-dried particles have been compared to those obtained 18 by conventional drying methods such as spray-drying and freeze-drying, finding 19 encapsulation efficiencies (EE) up to 98 % and spherical morphology for PGSS- and 20 spray-dried particles. Freeze-dried powders showed irregular morphology and EE from 21 22 95.8 to 98.6 %, depending on the freezing method. Differential scanning calorimetry (DSC) analysis revealed glass-transition and melting peaks of OSA-starch and a cold-23 crystallization peak corresponding to the encapsulated n-3 PUFA concentrate. 24 25 Compared to conventionally dried powders, PGSS-dried microparticles showed lower primary and secondary oxidation after 28 days of storage at 4 °C. Ascorbic acid addition 26 27 combined with the mild processing conditions of PGSS-drying yielded particles with a maximum peroxide value of 2.5 meq O₂/kg oil after 28 days of storage at 4 °C. 28

30 1. Introduction

An adequate intake of omega-3 polyunsaturated fatty acids (n-3 PUFAs) is 31 recommended in healthy diet guidelines due to their important benefits (Ruxton, Reed, 32 33 Simpson, & Millington, 2004). Long-chain n-3 PUFAs, mainly eicosapentaenoic (EPA, 20:5 n-3) and docosahexaenoic (DHA, 22:6 n-3) acids are eicosanoid precursors, which 34 are immunomodulatory molecules with a key role in the inflammatory response. EPA 35 and DHA are claimed to contribute to the normal brain, eye and cardiovascular 36 functions in adults and help in the normal development of the eyes, the brain and the 37 nervous system in children (EFSA, 2010). 38

The perceived health benefits of these compounds have created a strong demand for 39 40 EPA and DHA concentrates in the pharmaceutical and food industries. However, n-3PUFAs are unstable and very prone to oxidation, easily generating lipid hydroperoxides 41 42 and free radicals under oxidative conditions. These species negatively affect sensory 43 properties, since they can decompose into low-molecular-weight volatile compounds that are perceived as rancid, and what is more, they present potentially cytotoxic, 44 carcinogenic and mutagenic effects (Niki, 2009; Uluata, McClements, & Decker, 2015) 45 For these reasons, n-3 PUFA concentrates are often encapsulated in order to protect 46 47 them from light and oxygen during shelf life; and natural antioxidants such as tocopherols, phospholipids, ascorbic acid, or their mixtures are usually added (Baik et 48 al., 2004; Löliger & Saucy, 1994). 49

50 Materials of different nature can be used as n-3 PUFA encapsulating agents: proteins 51 such as whey protein isolate, sodium caseinate or gelatin, phospholipids such as 52 lecithin, or polysaccharides such as gum Arabic, carboxymethyl cellulose, maltodextrin, 53 chitosan, or modified starch are some examples of carrier materials for 54 microencapsulation of oils rich in n-3 PUFAs (Encina, Vergara, Giménez, Oyarzún-

Ampuero, & Robert, 2016). Among them, *n*-octenyl-succinic-anhydride modified starch
(OSA-starch) has been chosen in this work because it presents good emulsifying
properties and is suitable to encapsulate oils rich in *n*–3 PUFAs, as well as other
bioactive compounds such as essential oils and hydrophobic compounds (Carneiro,
Tonon, Grosso, & Hubinger, 2013; de Paz, Martín, Bartolomé, Largo, & Cocero, 2014;
Drusch, Serfert, Scampicchio, Schmidt-Hansberg, & Schwarz, 2007; Jafari, Assadpoor,
He, & Bhandari, 2008; Varona, Martín, & Cocero, 2011).

62 Different encapsulation techniques can be used to encapsulate n-3 PUFAs, such as emulsification, spray-drying, freeze-drying, coacervation, in situ polymerization, 63 extrusion, or fluidized-bed coating (Bakry et al., 2016). Among these, the most widely 64 65 used technique in the food and pharmaceutical industries is spray-drying, followed by freeze-drying. Freeze-drying is often applied to thermolabile and easily oxidizable 66 compounds due to the protective low temperatures and vacuum conditions involved in 67 68 the process. Its main drawback is the energy consumption, linked to the low temperature and high vacuum conditions as well as the long residence times required to completely 69 dry the product, which in turn translate into high processing costs. On the contrary, 70 spray-drying is a low-cost microencapsulation technology which operates in a relatively 71 72 simple and continuous way, thus it is commonly used at industrial scale (Bakry et al., 2016). 73

Prior to the drying step, the non-soluble n-3 PUFAs need to be dispersed into the 74 75 encapsulating agent solution, obtaining an oil-in-water (O/W) emulsion. Several methods can be used to prepare O/W emulsions, such as conventional emulsification 76 (colloid milling, high speed blending and high-pressure homogenization), ultrasound 77 membrane 78 (US)assisted emulsification, emulsification, and micro-channel emulsification (Chatterjee & Judeh, 2015). Among them, US-assisted emulsification has 79

grown in importance among the pharmaceutical, cosmetic, and food industries, thanks 80 81 to its versatility and the possibility of obtaining high quality food products with enhanced functional properties (Abbas, Hayat, Karangwa, Bashari, & Zhang, 2013). 82 83 US-assisted emulsification can be applied to improve stability and bioavailability of the dispersed bioactive compounds and, in particular, it can be used to obtain O/W 84 85 emulsions with nanometric droplet size and narrow size distribution. Typically, USassisted emulsification consists on applying low-frequency sound waves of 20-100 kHz 86 through a metallic sonotrode immersed in the liquid medium, in order to generate 87 disruptive forces that break down the macroscopic phases to nanosize droplets. The 88 89 nano-scale emulsions obtained present interesting functional properties and enhanced stability against oxidation (Abbas et al., 2013). 90

Supercritical fluids, and particularly supercritical carbon dioxide (SC-CO₂), are a 91 92 convenient medium to produce particles loaded with bioactive compounds. Carbon 93 dioxide is an inert, non-toxic solvent, and is completely released from the product as a gas once back to atmospheric conditions. Besides, the accessibility of the supercritical 94 state of carbon dioxide ($T_C = 31.1$ °C; $p_C = 73.8$ bar) and its advantageous physical 95 properties (high density and diffusivity, and low viscosity) make SC-CO₂ the solvent of 96 choice in many particle formation processes. (Türk, 2014). Among the several available 97 techniques, the Particles from Gas Saturated Solutions (PGSS) process overcomes the 98 problems of solubility limitations and high gas consumption of other particle formation 99 100 methods using SC-CO₂ (Türk, 2014). This technique can be used for drying aqueous 101 solutions, dispersions or, as in this work, O/W emulsions, in the so-called PGSS-drying 102 process (Türk, 2014).

Basically, the PGSS-drying technique consists on mixing an aqueous solution with
supercritical carbon dioxide upon saturation, and subsequently expanding the gas-

105 saturated solution down to atmospheric pressure through a nozzle. This technique can 106 be used as an alternative to conventional spray-drying, achieving a more efficient atomization due to the sudden vaporization of the dissolved CO₂ and the expansion of 107 108 gas bubbles in the solution during depressurization from supercritical to atmospheric 109 conditions. Both effects improve the atomization of the sprayed solution forming small droplets, thus reducing the particle size of the dried powder and enhancing the drying 110 process (Martín & Weidner, 2010; Weidner, 2009). Besides, and because of the intense 111 112 and deep cooling caused by the Joule-Thomson effect, it is possible to dry the product at low temperature (40-80 °C) (de Paz, Martín, & Cocero, 2012; Weidner, 2009). The 113 114 mild-temperature conditions, combined with the intrinsically inert atmosphere due to oxygen displacement, prevent, or at least delay, oxidative degradation of the 115 encapsulated bioactive compounds (de Paz et al., 2012; Weidner, 2009). Operating 116 117 conditions in the spray tower, particularly temperature and gas-to-product ratio (GPR), must be taken into account in order to operate above the dew line of the carbon dioxide-118 119 water system (Martín & Weidner, 2010), and ensure the complete drying of particles.

120 In this work, an n-3 PUFA enriched fish oil has been encapsulated by the alternative 121 and green technology Particles from Gas Saturated Solutions (PGSS)-drying. The main hypothesis of the study is to explore whether or not the potential benefits of 122 supercritical carbon dioxide technologies applied to particle formulation and 123 encapsulation may affect particle properties and oxidative stability of heat-sensitive and 124 125 easily oxidizable compounds such as n-3 PUFAs, compared to other conventional 126 drying methods. This way, the PGSS-dried particles have been compared to those obtained by spray-drying and freeze-drying, which are commonly applied in the 127 128 pharmaceutical, cosmetic, and food industries to dry aqueous solutions and dispersions. 129 Characterization of the particles obtained by the different drying methods has been

performed in terms of particle morphology, residual humidity, and particle size distribution of the reconstituted particles. Besides, encapsulation efficiency and oxidative stability (primary and secondary oxidation) of the encapsulated n–3 PUFA concentrate have been monitored over time in the particles formulated with each of the drying methods. Additionally, an antioxidant (ascorbic acid) has been added to some of the formulations as a strategy to potentially enhance the oxidative stability of the encapsulated *n*–3 PUFA concentrate.

137

138 2. Materials and methods

139 **2.1.** Materials

n–3 PUFA concentrate from fish oil, AlgatriumTM Plus, was kindly donated by Brudy
Technology S.L. (Spain). It has been stored at 4 °C in darkness and N₂ atmosphere. HiCapTM 100, an octenyl-succinic-anhydride modified starch (OSA-starch) derived from
waxy maize, was provided by Ingredion Inc. (Germany). Carbon dioxide (99.9%) was
provided by Air Liquide S.A. (Spain). Ascorbic acid (L(+)-Ascorbic acid, AA) was
purchased from Panreac AppliChem (Spain).

146 37% hydrochloric acid (HCl), diethyl ether, 1-butanol, 2-propanol, methanol, 2thiobarbituric acid (TBA), and trichloroacetic acid (TCA) were provided by VWR 147 Chemicals (Germany). Hexane, absolute ethanol, Iron(II) sulphate heptahydrate, and 148 149 ammonium thiocyanate were purchased from Merck KGaA (Germany). 2,2,4trimethylpentane (isooctane) and barium chloride dihydrate were supplied by Macron 150 Fine Chemicals (France) and Panreac AppliChem (Spain), respectively. Cumene 151 hydroperoxide and 1,1,3,3-tetraethoxypropane (TEP) standards were purchased from 152 Sigma Aldrich (USA). 153

155 **2.2.** Characterization of the *n*-3 PUFA concentrate

Neutral lipid profile of the *n*-3 PUFA concentrate has been analyzed by normal-phase
HPLC (NP-HPLC). Separation was carried out at room temperature in a Lichrospher
Diol column (5 mm, 4 mm×250 mm) and detection was performed by evaporative light
scattering (ELSD) (Agilent Technologies 1200 Series, USA) at 35 °C and 3.5 bar.
Solvent gradient and calibration procedure have been reported elsewhere (Solaesa,
Sanz, Falkeborg, Beltrán, & Guo, 2016).

162 Fatty acid profile of the n-3 PUFA concentrate has been determined according to the 163 AOAC Official Method (AOAC International, 2012) in a Hewlett Packard gas chromatograph (6890N Network GC System) equipped with an auto-sampler (7683B 164 165 series) and a flame ionization detector (FID). The separation was carried out in a fused silica capillary column (Omegawax-320, 30 m×0.32 mm i.d.) with helium (1.8 mL/min) 166 as carrier gas. Injection and detection temperatures, as well as ramp conditions have 167 been previously reported (Rebolleda, Rubio, Beltrán, Sanz, & González-San José, 168 2012). Most of the fatty acids were identified by comparison of their retention times 169 170 with those of chromatographic standards (Sigma Aldrich). As indicated by the AOAC Official Method (AOAC International, 2012), an internal standard (methyl-tricosanoate, 171 172 C23:0) was used for quantification purposes.

173 HPLC with diode array detection (HPLC-DAD) of the n-3 PUFA concentrate was 174 carried out in order to detect tocopherol isomeric forms and other vitamin E analogs 175 added to the n-3 PUFA concentrate, as their presence was reported by the provider. The 176 analytical method is based on the IUPAC official method (Pocklington & 177 Dieffenbacher, 1988) with slight modifications, as reported in Rebolleda et al., (2012).

Separation was performed in an ACE 5 silica 250 mm × 4.6 mm column with 1 mL/min of hexane:2- propanol (99:1) as the mobile phase. An isocratic gradient was used, and the total run time was 15 min. α -, β -, γ -, and δ -tocopherols were monitored at λ = 296 nm. For identification and quantification of each tocopherol isomer, a calibration curve with different amounts of the respective standard compound (Sigma Aldrich) was constructed.

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185 2.3. Ultrasound-assisted emulsification

O/W emulsions were formulated in a weigh proportion of 70:24:6 (water:carrier:n-3 186 PUFA concentrate), which in preliminary experiments was found to be the optimal in 187 terms of obtaining the smallest droplet size. First, an aqueous solution of the 188 encapsulating agent was prepared by dissolving 24.0 g of Hi-Cap[™] 100 in 70.0 mL of 189 distilled water. Subsequently, 6.0 g of n-3 PUFA concentrate were added drop by drop 190 to the carrier solution under continuous stirring. Then, the mixture was stirred for 191 192 5 minutes to obtain a pre-emulsion, which was subsequently processed in a 20 kHz 750 W ultrasonic liquid processor Vibra-Cell 75043 (Sonics & Materials Inc.) with a 193 Ø13 mm titanium alloy sonotrode. Based on previous studies, amplitude was set at 100 194 195 % and sound waves were delivered in pulses (5 s On/5 s Off) in order to avoid excessive heating of the sample, for a total processing time of 180 s. O/W emulsions were 196 produced in batches of 100 g. 197

198

199 2.4. PGSS-drying

200 O/W emulsions were processed using the PGSS-drying technique in order to remove 201 water and obtain a solid powder with the encapsulated n-3 PUFA concentrate loaded

202 into the OSA-starch microparticles. Fig. 1 presents the schematic flow diagram of the 203 PGSS-drying apparatus, in which CO₂ was fed by a membrane pump (LEWA) and preheated using a silicone bath before injection into the static mixer, where it was mixed 204 205 with the O/W emulsion at the selected pressure and temperature. The CO₂ mass flow 206 rate was measured with a Coriolis flow meter (Danfoss) with an accuracy of ± 0.1 kg 207 CO_2/h . Temperature before and after the static mixer was measured by means of Pt100 208 thermoresistances (accuracy of ± 0.1 K), being the later under PID control. Pressure in 209 the CO₂ line and after the static mixer was measured with pressure transmitters (DESIN Instruments) with an accuracy of \pm 0.05 MPa. Bourdon manometers (Nuova Firma) 210 211 were installed to provide secondary lectures of the operating pressure.

212 The O/W emulsion was pumped into the static mixer by a GILSON 305 piston pump 213 (max. flow rate: 25 ± 0.1 mL/min). The gas-saturated emulsion was then expanded into 214 the spraying tower through a capillary nozzle with an internal diameter of 400 µm 215 (Spraying Systems Co., Ref.: PF1650-SS). The spraying tower was made of PVC and 216 heated by electrical resistances. Temperature in the spray tower was also measured with a Pt100 probe and controlled using a PID. CO₂ was vented off the spraying tower and 217 passed through a water vapor condenser before final release. As security elements, a 218 rupture disk, check valves, and a relief valve were installed at different points in the 219 220 high-pressure circuit.

Typically, a PGSS-drying experiment began with the preheating of the system up to the desired temperature in the static mixer, fixed at 110 °C, and in the spraying tower, which was set at 55 °C. When temperature was achieved, CO_2 was pumped up to the desired pressure, which was fixed at 10.0 MPa. Pressure in the static mixer and temperatures in the static mixer and the spraying tower were selected based on previous studies (Varona et al., 2011). Once temperature and pressure conditions were stable, the emulsion pump was started at a flow rate such that the desired GPR, which was selected at 30 g/g, was obtained. After all the O/W emulsion was processed, CO₂ was allowed to flow through the system at the same pressure and temperature conditions during 15 minutes in order to completely dry the particles. After that, the system was depressurized and particles were collected from the walls and bottom of the spraying tower and stored in darkness and refrigeration at 4 °C for subsequent analyses.

233

234 2.5. Spray-drying

235 Spray-drying is a conventional, well-known drying technique which is widely used in 236 the pharmaceutical, cosmetic and food industries; thus, it was chosen to compare the characteristics of the powder that may be obtained conventionally to those of the 237 238 powder obtained by the alternative PGSS-drying process. The spray-drying process was carried out in a commercial Buchi B-290 mini Spray-dryer. The O/W emulsion, 239 obtained as described in section 2.3, was fed into the spray-drying apparatus at an inlet 240 241 temperature of 155 °C, and %pump of 8 %, which was equivalent to a mass flow of emulsion of 3.0 g/min. Outlet temperature was 100 °C. The emulsion was sprayed 242 through a nozzle with 1.5 mm diameter and dried under a N_2 flow of 360 L/h. 243

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245 **2.6.** Freeze-drying

O/W emulsions obtained by the US-assisted method described in section 2.3 were submitted to two different freezing methods: (1) conventional at -20 °C overnight, and (2) freezing with liquid nitrogen (-196 °C). Samples were then equilibrated at -80 °C for 2 h and submitted to freeze-drying in a Labconco Freeze Dry System at $1.5 \cdot 10^{-4}$ mbar during 48 h. These two different freezing methods were chosen in order to evaluate the

effect of the freezing step, since the slower conventional freezing process is more likely to form large crystals of water, which could adversely affect the emulsion stability and structure, whereas the rapid freezing achieved with liquid nitrogen could better preserve the physical structure of the emulsion.

255

256 2.7. Characterization of the O/W emulsion

257 2.7.1. Droplet size analysis of the O/W emulsions

The droplet size distribution of the O/W emulsions (original and reconstituted) was measured by a Laser Diffraction (LD) equipment (Malvern Mastersizer 2000). A small amount of sample was suspended in the suspension container filled with distilled water under gentle agitation. In the case of the reconstituted O/W emulsions, the dried powders were firstly dissolved in distilled water, maintaining the original ratio of 70:24:6 wt. (water:carrier:n-3 PUFA concentrate).

Droplet size measurements are reported as relative volume distribution and defined by the mean diameter over volume (DeBroukere mean, D[4,3]) and the volume/surface mean diameter (Sauter mean, D[3,2]), calculated as in Eqs. 1 and 2, respectively.

267
$$D[4,3] = \frac{\sum_{i=1}^{n} D_{iv_{i}}^{4}}{\sum_{i=1}^{n} D_{iv_{i}}^{3}}$$
 (1)

268
$$D[3,2] = \frac{\sum_{i=1}^{n} D_{iv_{i}}^{3}}{\sum_{i=1}^{n} D_{iv_{i}}^{2}}$$
 (2)

269 where D_i is the diameter of the *i*th particle.

The median particle size $(d_{0.5})$, defined as the maximum particle diameter below which 50 % of the sample volume exists, is also reported. The span value, defined in Eq. 3, was also calculated.

273 span =
$$\frac{d_{0.9} - d_{0.1}}{d_{0.5}}$$
 (3)

where d_x is the maximum particle diameter below which x% of the sample volume exists. Span values near to 1 indicate a narrow particle size distribution (PSD).

276

277 2.7.2. Emulsion stability

278 Physical stability of the O/W emulsion was analyzed by static multiple scattering in a vertical scan analyzer Turbiscan Lab Expert (Formulaction Inc.) with ageing station 279 280 AGS. By means of two optical sensors, the instrument measures the light transmitted through the emulsion (180° from the incident light, transmission, T) and the light 281 282 backscattered by the emulsion droplets (45° from the incident light, backscattering, BS). 283 The scanning process is made vertically along the glass cell from bottom to top, and the T/BS are each plotted as a function of the emulsion height in the glass cell. By 284 285 monitoring the T/BS profiles at different time intervals, physical changes in the 286 emulsion can be followed over time, which gives a detailed overview of dispersion stability or instability. In the current work, the stability of the original emulsion was 287 monitored at 4 h intervals during 24 days. Emulsion samples were kept in the ageing 288 station at a constant temperature of 25 °C. As variations in T profiles were lower than 289 290 2%, only BS profiles at different storage times were analyzed in this study.

291

292 2.7.3. Density of the O/W emulsions

293 Density of the O/W emulsions was measured in an Anton Paar DMA 5000 instrument at
294 25 °C. Measurements were carried out in triplicate.

295

296 **2.8.** Characterization of the dried powders

297 2.8.1. Yield, moisture, encapsulation efficiency and bioactive loading

Yield of particles was calculated as the ratio between the mass of collected particles ($m_{collected particles}$) and the theoretical mass fed to the PGSS-drying, spray-drying, or freeze-drying apparatus $m_{initial feed}$, expressed as weight percentage (Eq. 4).

301 Yield
$$(\%) = \frac{m_{\text{collected particles}}}{m_{\text{initial feed}}} \cdot 100$$
 (4)

Moisture content of the dried particles was determined gravimetrically. Samples (*ca*. 0.5 g) of particles obtained by the different methods used in this work were weighed before and after drying in an oven at 120 °C until constant weight.

305 Encapsulation efficiency (EE) was determined according to the method described by Wang et al. (Y. Wang, Liu, Dong, & Selomulya, 2016) with some modifications. For 306 307 the non-encapsulated oil determination, samples (ca. 1.0 g) of particles obtained by the 308 different methods used in this work were suspended with 25 mL of hexane in a Falcon 309 centrifuge tube, which was vortexed for 15 s at room temperature and centrifuged at 3000 rpm during 20 min. Immediately afterwards, the supernatant was taken and 310 311 filtered, and its oil content was measured spectrophotometrically at $\lambda = 286$ nm. The 312 same procedure was repeated two additional times to extract the potentially remaining non-encapsulated oil. A calibration curve was previously constructed with known 313 314 quantities of n-3 PUFA concentrate dissolved in hexane.

Total oil in the dried particles obtained by the different methods used in this work was 315 determined by acid digestion of approximately 1.0 g of powder with 37% HCl, and 316 subsequent extraction with diethyl ether and petroleum ether, following the AOAC 317 318 Official Method (AOAC International, 2005). After centrifugation at 3000 rpm during 20 min, the solvent phase with the extracted oil was taken and transferred to tared 319 round-bottom flasks in order to evaporate the solvent under vacuum (Heidolph rotary 320 evaporator). Total oil in the samples was determined by mass difference of the initial 321 322 clean round-bottom flask and that containing the extracted oil residue. As a blank, the same procedure was also followed with known quantities (ca. 1.0 g) of the carrier alone 323 (Hi-CapTM 100). The fat traces found in the carrier were subtracted from the total oil 324 content of the powders. 325

Encapsulation efficiency (EE) was calculated from Eq. 5.

327
$$\operatorname{EE}\left(\%\right) = \frac{\operatorname{TO} - \operatorname{nEO}}{\operatorname{TO}} \cdot 100$$
 (5)

328 where TO is the total oil content and nEO is the non-encapsulated oil.

The bioactive loading, which is also an important parameter of microencapsulated bioactive compounds (Encina et al., 2016), has been also calculated. It can be referred as to the total oil content (TO), expressed as mg oil/g sample.

332

333 2.8.2. Particle size analysis of the dried powders

Particle size analysis of the dried powders was carried out in a Malvern Mastersizer 2000 equipment, following the same procedure as in the original O/W emulsion (see section 2.7.1), yet dispersing the particles in absolute ethanol to avoid dissolution of the encapsulating agent.

339 2.8.3. Scanning electron microscopy (SEM)

Morphology of the dried particles was observed in a Scanning Electron Detector microscope JEOL JSM-6460LV with Energy Dispersive X-ray (JEOL Ltd. Japan) operating at 20 kV. Samples were gold-sputtered and observed with magnifications of 1500, 5000 and 10000x for PGSS- and spray-dried particles, and 50, 400 and 2000 or 3000x for the freeze-dried powders.

345

346 2.8.4. Differential scanning calorimetry (DSC)

A TA Instruments Q200 differential scanning calorimeter with refrigerated cooling 347 system (RCS90) and nitrogen purge gas was used. Melting point and enthalpies of 348 indium were used for temperature and heat capacity calibration. Samples (ca. 10 mg) 349 350 were placed in TA Tzero 40-µL aluminum pans and closed with hermetic aluminum lids 351 with a pinhole. An empty pan closed with pinholed lid was used as a reference. Starting temperature of the DSC analysis was set at 40 °C, and held for 30 min. Then, the system 352 was cooled down to -80° C at 10° C·min⁻¹. After an isothermal period of 30 min, samples 353 were heated from -80 °C to 350 °C at a constant heating rate of 10°C·min⁻¹. DSC 354 thermograms were recorded and analyzed with the Advantage v. 5.5.20 software (TA 355 Instruments). 356

357

358 2.9. Measurement of lipid oxidation

Oxidative status of the dried powders was determined in terms of primary oxidation (peroxide value, PV) and secondary oxidation (Thiobarbituric Acid Reactive Substances, TBARS). To observe the effect of each drying method, PV and TBARS were determined in the n-3 PUFA concentrate, as well as in the O/W emulsions before drying.

For the dried powders, PV and TBARS were measured right after each drying method (PGSS-drying, spray-drying, and freeze-drying) and monitored over a 28-day storage period. Dried powders were placed in closed containers and stored at 4 °C in darkness. Samples were withdrawn at 7-day intervals and dissolved in distilled water to obtain reconstituted emulsions with the original water:carrier:n-3 PUFA concentrate proportion (70:24:6 wt.). PV and TBARS analyses were carried out as described below.

370

371 **2.9.1.** Peroxide Value

PV was measured spectrophotometrically with a Hitachi U-2000 apparatus and 372 following the method described by Shanta et al. (Shantha & Decker, 1994) with slight 373 374 modifications. In brief, 10-50 mg of oil or 0.025-1.0 mL of emulsion, depending on the 375 expected PV, were taken in a centrifuge tube and mixed with 1.5 mL of isooctane:2propanol (3:1 v/v). The tube was vortexed for 15 s and centrifuged at 5000 rpm during 376 377 10 min. Immediately afterwards, 0.2 mL of the supernatant were transferred to a new centrifuge tube and 2.8 mL of methanol:1-butanol (2:1 v/v) were added. After vortexing 378 for 15 s, 15 μ L of 3.94 M ammonium thiocyanate and 15 μ L of a Fe²⁺ solution were 379 added. The Fe²⁺ solution was obtained by mixing 0.132 M barium chloride in 0.4 M 380 381 HCl and 0.144 M Iron(II) sulphate heptahydrate (1:1 v/v), centrifuging at 5000 rpm for 10 min, and taking the supernatant. Samples were vortexed again for 15 s and kept in 382 darkness for 20 min. Blanks were prepared the same as above with 0.3 mL of distilled 383 384 water instead of the oil or emulsion sample. Hydroperoxyde concentration was 385 determined spectrophotometrically at $\lambda = 510$ nm. A calibration curve was constructed using known concentrations of cumene hydroperoxide, ranging from 0.13 to 3.28 mM. Results were expressed in milliequivalents of oxygen per kg of n-3 PUFA concentrate (meq O₂/kg oil).

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390 2.9.2. TBARS analysis

TBARS present in the n-3 PUFA concentrate were determined following the method 391 392 described by Ke and Woyewoda (Ke & Woyewoda, 1979). Briefly, 10 mg of n-3 PUFA 393 concentrate were weighed in a screw-capped glass test tube. 5 mL of TBA work 394 solution, which was prepared by mixing 0.04 M 2-thiobarbituric acid in glacial acetic acid, chloroform, and 0.3M sodium sulphite (12:8:1 v/v), were also added to the screw-395 396 capped glass test tube. The mixture was vortexed for 15 s and incubated in a water bath 397 at 95 °C during 45 min. After cooling down the test tubes under running cold water, 2.5 398 mL of 0.28 M trichloroacetic acid were added to the samples, which were then mixed by inversion. Samples were then centrifuged at 2500 rpm for 10 min in order to separate 399 the pink aqueous phase from the chloroform phase. Absorbance of the aqueous phase 400 was measured at $\lambda = 538$ nm in a Hitachi U-2000 spectrophotometer. Blanks were 401 402 prepared the same as above, yet without the oil, and subtracted from the absorbance 403 measurement.

TBARS analysis of the original and reconstituted O/W emulsions was carried out following the method described by Mei *et al.* (Mei, McClements, Wu, & Decker, 1998) with slight modifications. Briefly, 0.025-1.0 mL of emulsion, depending on the expected oxidative status, were taken in screw-capped glass test tubes. Distilled water was used to complete to 1.0 mL if necessary. Subsequently, 2 mL of a TCA/TBA mixture – which was prepared by dissolving 7.5 g of TCA into 10 mL of 0.25M HCl,

adding this solution to 0.1875 g of TBA and completing to volume with 0.25M HCl in a 410 411 50 mL volumetric flask - were added and the glass test tube was tightly closed, 412 vortexed for 15 s and immersed in a water bath at 95 °C during 15 min. Then the vials 413 were cooled down under running cold water and centrifuged at 5000 rpm during 10 min. 414 Immediately afterwards, the supernatant was collected and its absorbance measured in a Hitachi U-2000 spectrophotometer at $\lambda = 538$ nm. Blank runs were also performed the 415 same as above, but without adding the emulsion, and its absorbance subtracted from the 416 417 measurements.

TBARS concentration in the emulsion and the *n*–3 PUFA concentrate samples was determined using a TEP standard curve with concentrations ranging from 2.5 to 20 nM. Results were expressed in mg malondialdehyde equivalents (MW = 72.06 g/mol) per kg of *n*–3 PUFA concentrate (mg MDA/kg oil).

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423 **2.10.** Statistical analysis

All results reported in this work represent the average of at least three independent measurements. Drying experiments performed in this work have been duplicated. Statistical analyses were performed using Statgraphics Centurion XVII software. Statistical significance was determined by analysis of variance (ANOVA) using the Fisher's least significant difference test. Results were deemed as statistically significant when p < 0.05.

431 **3. Results and discussion**

432 **3.1.** Characterization of the *n*-3 PUFA concentrate

Results obtained in the characterization analysis are summarized in Table S-1 of the 433 provided supplementary material. As it can be seen from Table S-1a, the fatty acid 434 profile of the n-3 PUFA concentrate is constituted by more than 90 % n-3 PUFAs, 435 436 being 73.49 % identified as DHA. Neutral lipid profile of the n-3 PUFA concentrate (Table S-1b) showed that more than the 75 % of the neutral lipids in the n-3 PUFA 437 concentrate are in the form of triacylglycerides, with 21.6 % being in the form of fatty 438 acid ethyl-esters. Traces of diacylglycerides and monoacylglycerides (1.2% and 0.7%, 439 respectively) were also found. The high content of triacylglycerides is an important 440 441 feature of the n-3 PUFA concentrate, since these compounds are the natural form of food lipids and they may present better bioavailability and stability against oxidation 442 443 (Rubio-Rodríguez et al., 2010). Tocopherol analysis by HPLC-DAD revealed a racemic 444 mixture of tocopherol added as antioxidant (again, this is in consonance with consumer's preference for natural sources). α -, β -, γ -, and δ -tocopherol isomers were 445 identified and quantified. Results are showed in Table S-1c. 446

447

448 **3.2.** Characterization of the O/W emulsion

449 3.2.1. Droplet size of the O/W emulsions

Results from the analysis of droplet size distribution are reported in Table 1 for the original and reconstituted O/W emulsions. In general, similar values for D[4,3] and D[3,2] were found in all samples, with the exception of the conventionally freeze-dried powder that showed significantly higher values for both D[4,3] and D[3,2], which

means that the reconstituted emulsion from the conventionally freeze-dried powderpresented larger mean diameters both in volumetric and surface basis, respectively.

Median droplet size by volume $(d_{0.5})$ of the emulsion is sub-micrometric, with $d_{0.5} =$ 456 457 0.114 μ m and a D[4,3] and D[3,2] of 0.144 μ m and 0.114 μ m, respectively. On the other hand, the drying methods proposed in this work significantly increased $d_{0.5}$ after 458 459 reconstitution, with the exception being the freeze-dried particles with liquid N₂, in 460 which no statistically significant differences were found with the original emulsion (p < p461 0.05). Still, most droplet populations were around 0.130 µm for particles obtained by 462 PGSS-drying, freeze-drying and spray-drying methods, which demonstrates that the proposed drying methods do not produce aggregation of oil droplets. The span values 463 464 followed the same trend as $d_{0.5}$, with original emulsion < freeze-drying (liq N₂) < spraydrying \approx freeze-drying (-20 °C) < PGSS-drying. The higher span values in the 465 466 reconstituted emulsions may be due to higher polydispersity.

467

468 3.2.2. Emulsion stability

Physical stability of the US-assisted O/W emulsion was analyzed by static multiple 469 470 scattering. Changes in the backscattering profile (ΔBS) of the O/W emulsion sample 471 were recorded every 4 h during 24 days of storage at 25 °C and plotted vs. time. Results 472 are provided as supplementary material in Fig. S-1. As shown in this figure, ΔBS in the top-section reached 5% increment on day 2 and started to decrease in the lower section 473 $(|\Delta BS| > 2\%)$ on day 5, indicating creaming destabilization due to phase separation and 474 475 migration of the lighter oil droplets to the top zone. Moreover, a slight BS increase over time in the middle section of the glass cell can be seen (Fig. S-1), indicating emulsion 476 droplet size slightly increased over the 24-day storage period. 477

479 3.2.3. Density of the O/W emulsions

480 Density measurements were carried out for the original emulsion as well as for the481 reconstituted dried powders. Results obtained are shown in Table 1.

No statistical difference (p < 0.05) was found between the densities of the original and reconstituted emulsions, being those the means of three independent measurements. Thus, the average value of 1.091281 g·cm⁻³ was used for the volume-to-mass transformations necessary in the PV and TBARS calculations.

486

487 **3.3.** Characterization of the dried powders

488 3.3.1. Yield and bioactive loading

489 Calculated yield of particles and loading of fish oil concentrate of each of the proposed drying methods is showed in Table 1. The spray-drying method exhibits the lowest 490 yield, which is because the particles deposited on the wall of the spraying tower were 491 492 collected separately and finally not considered due to its low oxidative quality (results not shown). In the PGSS-drying method, some of the finer particles were blown away 493 by the vented CO₂ and deposited in the condenser. This wet powder was not collected, 494 slightly reducing the final yield. In the case of the freeze-dried particles, the observed 495 496 yield is very close to unity. This trend was also observed by other authors (Lévai et al., 2017) and may be attributed to the one-pot processing and the preservation of the 497 498 emulsion structure during freezing.

Regarding the bioactive loading, it is close to the maximum theoretical loading of 200 mg/g sample in all cases, and no statistical differences (p < 0.05) are observed no matter the drying method used to obtain the particles. Nevertheless, the spray-dried 502 particles present a slightly lower average value, which may be attributed to the higher 503 moisture content that will be discussed next. On the other hand, the freeze-dried 504 particles present the highest fish oil concentrate loading, which is possibly linked to the 505 aforementioned preservation of the emulsion integrity.

506

507 3.3.2. Moisture content

508 The moisture content of the particles prepared with different drying methods is showed in Table 1. The spray-dried particles showed the highest residual humidity, whereas the 509 510 PGSS-drying technique gave the lowest moisture value. Humidity values for the spray-511 dried particles found in this work are higher than those reported in the literature, which 512 are usually around 1-3 % (Carneiro et al., 2013; Hogan, McNamee, O'Riordan, & 513 O'Sullivan, 2001). In the case of the freeze-dried particles, no significant difference in 514 the final humidity was found (p < 0.05), no matter the freezing method used 515 (conventional at -20 °C or with liquid nitrogen).

516

517 3.3.3. Encapsulation efficiency

Encapsulation efficiency is one of the most important quality parameters in encapsulated fish oil and n-3 PUFA concentrates. The presence of free oil may adversely affect the physical properties of the final product, such as flowability and bulk density, and would also enhance lipid oxidation (Y. Wang et al., 2016). Table 1 shows the initial encapsulation efficiency of the drying methods proposed in this work.

In general, high initial encapsulation efficiencies, no matter the drying method used, were obtained. It can be noticed that the powder obtained by freeze-drying with conventional -20°C freezing presents a significantly lower (p < 0.05) initial

526 encapsulation efficiency, with $EE = 95.8 \pm 0.2$ % (Table 1). As it has been previously 527 mentioned, it is likely that partial destabilization of the emulsion and release of small amounts of n-3 PUFA concentrate may have happened, probably due to the mechanical 528 529 and hygroscopic forces caused by the growing of large water crystals during the slow 530 freezing process. By comparison, freeze-dried particles obtained with liquid nitrogen present the highest encapsulation efficiency with 98.6 \pm 0.1 % (Table 1), which reflects 531 that the emulsion casting is preserved with a rapid and deep freezing step. Similar 532 533 results have been also obtained by Lévai et al. (Lévai et al., 2017) dealing with freezedried quercetin encapsulated in soybean lecithin. Still, more than 95 % of the total n-3534 535 PUFA concentrate loading was encapsulated by conventional freeze-drying, and almost 98 % encapsulation efficiency was obtained by PGSS-drying (97.9 \pm 0.3 %), which is 536 537 similar to the EE value of the spray-dried microparticles (97.5 \pm 0.1 %). Carneiro *et al.* 538 (Carneiro et al., 2013) compared combinations of maltodextrin and Hi-Cap and other 539 wall materials to encapsulate flaxseed oil by spray-drying, finding Hi-Cap as the best in 540 terms of EE, with 95.7 %. Results obtained in this work are slightly higher in all cases 541 except for conventionally freeze-dried particles, which may be attributed to the optimized US-assisted emulsification process. 542

Surface oil of the dried particles has been analyzed over time during 28 days of storage 543 544 at 4°C in darkness and ambient oxygen concentration to check if some of the n-3 PUFA 545 concentrate could have been released. Results obtained are summarized in Fig. S-2 of 546 the supplementary material. As Fig. S-2 shows, spray-dried particles released around 547 2% of the total encapsulated n-3 PUFA concentrate during the first 7 days and then the 548 release continued at a lower rate, down to 94 % encapsulated oil after 28 days. In the case of the conventionally freeze-dried particles, a slight decrease in the encapsulated 549 550 oil can be seen after the second week of storage; whereas for the PGSS- and freezedried particles frozen with liquid N_2 , no significant changes in the encapsulation efficiency were noted during the first 21 days and only a slight decrease started to occur after the fourth week of storage.

554

555 3.3.4. Particle Size Analysis

The particle size distribution plot of PGSS-dried and spray dried particles is provided in Fig. 2. Particle mean diameters ($d_{0.5}$) varied from 28.605 µm for PGSS-dried particles to 35.375 µm for the spray-dried particles. The span value of the PGSS-dried particles (1.663) was also lower than that of the spray-dried particles (6.082).

The microparticles produced by spray drying showed a bimodal distribution with a group of particles centered around 30 μ m and a second population around 250 μ m. This justifies the high span value and may be linked to particle swelling during drying as well as to agglomeration due to the higher moisture content. This agglomerated clusters are also visible in the SEM images showed in Fig. 3b and discussed in the next section.

565 On the other hand, the PGSS-dried particles show a monomodal particle size 566 distribution with smaller mean diameter. As it has been reported in previous works (de 567 Paz et al., 2012), the effective atomization caused by CO_2 vaporization may have led to 568 the production of smaller and monodisperse particles.

569

570 3.3.5. Particle morphology (SEM)

571 Visual morphology of the dried powders can be observed in the SEM micrographs
572 (Fig. 3). Both PGSS- and spray-dried particles present spherical morphology. For the
573 PGSS-dried particles, small spheres with diameters ranging from 2 µm to 5 µm can be

observed together with some larger agglomerates around 10-20 µm diameter (Fig. 3a). 574 575 Fractured particles are also seen in some micrographs, showing a porous internal structure in which the n-3 PUFA concentrate is probably encapsulated. As it has been 576 577 also observed in the particle size analysis, spray-dried particles show more variance in size. A small population of microparticles around 2 µm was detected together with 578 some specimens larger than 20 µm and particle clusters around 150 µm (Fig. 3b), which 579 is also in accordance with the results obtained in the particle size analysis (section 580 581 3.3.4). This variety in size has been also reported in the literature (Carneiro et al., 2013), and seems to be a typical characteristic of particles produced by spray drying. Spray-582 583 dried particles also showed a rougher surface than PGSS-dried samples, with more imperfections or 'teeth'. These surface depressions are associated to the collapse of the 584 585 particle hollow core once the crust is formed during the initial stages of drying. Similar 586 morphological characteristics have been also found in the literature, either with OSAstarch as encapsulating agent (Carneiro et al., 2013), or with other materialas such as β-587 588 glucans (Salgado, Rodríguez-Rojo, Alves-Santos & Cocero, 2015).

In the case of the freeze-dried particles, larger and more irregular particles have been 589 590 produced. Conventionally freeze-dried powder presents a flakey or scaly appearance, forming planar structures with some dimensions being larger than 100 µm (Fig. 3c). 591 592 Some dents can be seen in the surface of several particles, probably corresponding to the voids left by water crystals after sublimation. In larger magnifications (3000x) a 593 594 porous internal structure can be also appreciated, being the n-3 PUFA concentrate 595 likely encapsulated inside these vesicles. In the case of the freeze-dried powder frozen with liquid N_2 (Fig. 3d), a powder finer than the conventionally frozen (Fig. 3c) has 596 been obtained. Some particles show an alveolar structure, which may have been formed 597

by liquid nitrogen boiling during freezing of the O/W emulsion. These alveolar holes
present diameters around 5-7.5 μm.

600

601 3.3.6. Differential Scanning Calorimetry (DSC)

602 DSC runs of PGSS-dried particles, modified OSA-starch (Hi-Cap 100) used as a carrier, 603 and *n*-3 PUFA concentrate revealed cold-crystallization, glass-transition (gelatinization) 604 and melting peaks. The peak temperatures of these thermal events are summarized in Table 2. Endothermic peaks near 80 °C were observed in the PGSS-dried and Hi-Cap 605 606 100 samples, which probably correspond to the glass transition (gelatinization) of OSAstarch. A second endothermic peak was found around 220 °C in both PGSS-dried 607 608 particles and Hi-Cap 100, which may be linked to the melting of OSA-starch. Similar 609 glass-transition and melting temperatures have been reported in the literature for this polymer (Yu & Christie, 2001). 610

611 In the lower temperature range, an exothermic cold-crystallization peak was noticeable for the n-3 PUFA concentrate and for the PGSS-dried particles, which may correspond 612 613 to some lipid compound of the n-3 PUFA concentrate transitioning from liquid to solid 614 state. This assumption can be corroborated by the studies of Tolstorebrov et al. 615 (Tolstorebrov, Eikevik, & Bantle, 2014), in which cold-crystallization peaks in the range -75 to -55 °C have been reported for some olein-, linolenin-, and linolein-616 617 containing tryacylglycerides, which are minoritary constituents of the n-3 PUFA 618 concentrate (Table S-1a). The slightly lower crystallization temperature observed in the 619 PGSS-dried particles compared to the n-3 PUFA concentrate alone (AlgatriumTM Plus) 620 is probably linked to the particle shell offering heat transfer resistance to the 621 encapsulated oil, and thus delaying the cold crystallization event.

623 **3.4.** Oxidative stability of the dried powders

624 Peroxide value (PV) and thiobarbituric acid reactive substances (TBARS) have been 625 systematically determined in the PGSS-dried powders with and without ascorbic acid (AA) during 28 days of storage at 4 °C and dark conditions. In order to determine the 626 627 initial oxidative status, PV and TBARS were measured in the n-3 PUFA concentrate 628 and in the original emulsion right after US-assisted emulsification. With the purpose of 629 comparing the different drying methods used in this work, PV and TBARS of the spraydried and the freeze-dried particles were measured after formulation of the powders 630 (day 0) and after 28 days of storage under the same conditions as the PGSS-dried 631 632 particles (4°C, darkness). Results obtained are summarized in Fig. 4.

633 Fig. 4a shows that PV increases from 1.64 \pm 0.05 meq O₂/kg oil in the *n*-3 PUFA concentrate up to 5.6 ± 0.3 meq O₂/kg oil during the US-assisted emulsification process, 634 which slightly surpasses the maximum limit of 5 meq O₂/kg oil for fish oil concentrates 635 intended for direct human consumption (Codex Alimentarius Comission, 2017). It is 636 likely that the high energy input involved in the ultrasonication process promoted a 637 638 temperature increase that may negatively affect the oxidative status of the n-3 PUFA concentrate (Abbas et al., 2013). As a strategy to prevent primary oxidation during US-639 assisted emulsification, 20 mM ascorbic acid (AA) was added to the emulsion 640 formulation. AA concentration was selected based on Uluata et al. (Uluata et al., 2015) 641 642 studies on lipid oxidation in O/W emulsions.

As it can be seen in Fig. 4a inset (O/W emulsion), the antioxidant successfully protected the n-3 PUFA concentrate and even reduced the PV of the emulsion down to 0.19 ± 0.03 meq O₂/kg oil. This behaviour has been also observed by Uluata *et al.* in O/W emulsions with AA (Uluata et al., 2015) and it is likely related to AA's ability to

647 inactivate free radicals such as lipid hydroperoxides. Other mechanisms can be also 648 involved in the observed antioxidant activity, since AA can act as an oxygen scavenger thanks to the enediol group in carbons 2 and 3 (Johnson, 1995; Liao & Seib, 1988), or 649 650 even play a synergistic role by means of regenerating other antioxidants such as the tocopherol originally present in the n-3 PUFA concentrate (Reische, Lillard, & 651 Eitenmiller, 2008). However, it is not easy to determine which of these pathways is 652 taking place in any given food system (Uluata et al., 2015) and it is likely that all of 653 654 them occur simultaneously.

If we focus on the PV results obtained after formulation of the dried particles (Fig. 4a 655 656 day 0), it can be seen that PGSS-drying promoted a slight PV increase up to 5.9 ± 1.5 657 meq O₂/kg oil in the emulsion without AA, although this value is not significantly different (p < 0.05) from the PV of the original emulsion. Furthermore, AA addition had 658 659 a significant (p < 0.05) effect on the PV of the PGSS-dried particles, since only a slight 660 increase from 0.19 \pm 0.03 to 0.5 \pm 0.1 meg O₂/kg oil was observed in the PGSS-dried 661 particles with antioxidant (Fig. 4a day 0). On the other hand, the spray-drying process yielded particles with much lower oxidative quality (PV = $28.0 \pm 1.6 \text{ meq } O_2/\text{kg oil}$). As 662 some authors have pointed out for the spray-drying process (Drusch & Berg, 2008; H. 663 Wang et al., 2011), it is likely that the rapid formation of the particle shell increased the 664 665 resistance to evaporation of water trapped inside the particle core, promoting a rapid 666 temperature increase in the particles and prolonging the n-3 PUFA exposure to high 667 temperatures, thus promoting oxidation and increasing the PV after spray-drying 668 formulation. The freeze-drying process with liquid nitrogen achieved good results, with 669 $PV = 4.6 \pm 1.8 \text{ meq } O_2/kg \text{ oil, which is not statistically different } (p < 0.05) \text{ from that of}$ the original emulsion (Fig. 4a day 0). This result can be related to the freeze-drying 670 671 process being a degradation-free technology, since the samples are not submitted to

high processing temperatures and processed in absence of light and in an almost inert atmosphere due to vacuum conditions. Unexpectedly, the conventionally frozen emulsion did overcome oxidation despite the favourable processing conditions, showing a PV = 12.4 ± 1.5 meq O₂/kg oil (Fig. 4a day 0). This is likely due to oxygen contact during the conventional freezing step, in which the samples were held overnight at - 20° C under ambient oxygen concentration.

In view of the results (Fig. 4a day 0), we can infer that PGSS-drying is a suitable method to formulate dried particles loaded with n-3 PUFAs, more so if we combine the mild processing conditions with the addition of an antioxidant such as AA. As it has been previously stated, the short residence time of the O/W emulsion in the PGSSdrying system as well as the inert CO₂ atmosphere prevent the loaded bioactive compounds from degradation (Weidner, 2009) and as such, the n-3 PUFA concentrate can be successfully protected from oxidation.

Oxidative stability of the PGSS-dried particles was monitored during 28 days of storage 685 686 in darkness at 4 °C (Fig. 4a days 1-28). Results obtained showed a sustained increase of primary oxidation, reaching values of PV = 25.2 ± 0.7 meq O₂/kg oil after 28 days of 687 storage (Fig. 4a). On the other hand, AA successfully protected the PGSS-dried 688 particles from primary oxidation during storage, being the values found significantly 689 lower (p < 0.05) than those of the PGSS-dried particles without antioxidant. The highest 690 691 PV was found after 14 days of storage and was $2.5 \pm 0.5 \text{ meq } O_2/\text{kg}$ oil, still below the 692 maximum allowable limit according to legislation, and remained with no significant 693 changes (p < 0.05) during the rest of the 28-day storage period, reaching a final value of 694 $2.2 \pm 0.3 \text{ meq } O_2/\text{kg oil (Fig. 4a)}.$

Comparing the primary oxidation of the particles obtained by the different drying 695 methods after 28 days of storage, we can see the same trend as in the PV analysis after 696 formulation, although PV increased in all samples (Fig. 4a days 1-28). Freeze-dried 697 particles frozen with liquid nitrogen maintained a relatively low PV of 16.9 ± 0.8 meg 698 O₂/kg oil, which is likely linked to the good encapsulation efficiency and the 699 preservation of the physical structure of the emulsion thanks to the fast and deep-700 701 cooling effect of liquid nitrogen. The same was not true for the conventionally freeze-702 dried particles, with PV = 37.7 ± 3.7 meq O₂/kg oil after 28 days of storage. Spray-dried particles showed the highest PV with 66.0 ± 0.4 meq O₂/kg oil after 28 days of storage. 703 704 The higher oxidation rates of these two samples (spray-drying and conventional freeze-705 drying) are probably due to the high starting PV (day 0) as well as their lower 706 encapsulation efficiency, which implies more oil in the particle surface susceptible to 707 oxidation. A similar encapsulation efficiency vs. oxidation rate inverse relationship has been observed by other authors (Yang & Ciftci, 2017). However, PGSS-dried and 708 709 freeze-dried particles with liquid nitrogen exhibited high encapsulation efficiencies (up 710 to 98%), and still encapsulated n-3 PUFA concentrate was not fully protected against 711 primary oxidation (PV after 28 days = 25.2 ± 2.2 and 16.9 ± 0.8 meq O₂/kg oil, respectively). This trend can be explained by taking into account not only the oxidation 712 713 of the oil present in the particle surface, but also oxygen diffusion through the 714 encapsulating material. It must be also pointed out that the fish oil concentrate used in this work is extremely rich in n-3 PUFAs, which are highly prone to oxidation. This 715 716 highly sensitive-to-oxidation fatty acid profile may also offer an explanation to the higher oxidation rates obtained in this work compared to other studies, even with no 717 718 accelerated storage (Carneiro et al., 2013; Yang & Ciftci, 2017).

TBARS analysis results are summarized in Fig. 4b. Although there is no legal 719 maximum limit for this parameter in food products, we can take the values of 10 µmol 720 MDA equiv/kg fish and 1-2 µmol MDA equiv/g fat given in the FAO guidelines (Huss, 721 722 1995) as an orientative basis to evaluate rancidity of the n-3 PUFA concentrate (1 µmol MDA equiv/g fat corresponds to 72.06 mg MDA/kg oil). From Fig. 4b we can see that 723 initial TBARS of the n-3 PUFA concentrate lay below this rancidity limit (TBARS = 724 41.1 ± 2.7 mg MDA/kg oil). US-assisted emulsification slightly increased the TBARS 725 726 value up to 54.8 ± 0.6 mg MDA/kg oil in the formulation without AA, whereas the addition of AA yielded particles with TBARS = $42.8 \pm 1 \text{ mg MDA/kg oil}$ (Fig. 4b day 727 728 0). In view of the results, AA addition slowed down secondary oxidation during the ultrasonication step since no significant difference (p < 0.05) between the AA-added 729 emulsion and the n-3 PUFA concentrate was found (Fig. 3b inset). 730

Among the dried powders (Fig. 4b, day 0), spray-dried particles showed the highest 731 secondary oxidative status with a TBARS value of 88.5 ± 6.0 mg MDA/kg oil, which is 732 733 above the FAO rancidity limit (Huss, 1995). PGSS-drying process slightly increased TBARS up to 59.4 ± 4.4 mg MDA/kg oil, whereas the addition of AA did not make any 734 735 statistically significant difference (p < 0.05). Both PGSS-drying with and without AA, 736 and freeze-dried powder with liquid N₂ showed no statistically significant differences with the original emulsion, which gives an idea of the protective effect of these drying 737 techniques against secondary oxidation. On the contrary, the conventionally frozen 738 particles were not successfully protected, and TBARS increased up to 74.5 ± 3.5 mg 739 740 MDA/kg oil after the conventional freeze-drying process.

Secondary oxidation products were also monitored in the PGSS-dried particles during the 28-day storage period. In Fig. 4b (days 1-28), we can see that TBARS in the PGSSdried particles without AA did not significantly increase (p < 0.05) up to the second

week of storage, when TBARS value raised from 69.2 ± 1.4 up to 110.9 ± 1.8 mg 744 MDA/kg oil, reaching a final value of 141.0 ± 1.9 mg MDA/kg oil after 28 days of 745 storage. On the other hand, AA addition delayed secondary oxidation for the first 14 746 747 days of storage, obtaining significantly lower (p < 0.05) TBARS values than those of the control sample without antioxidant, yet increasing thereafter and even exceeding the 748 control after 28 days of storage (TBARS = 141.0 ± 1.9 mg MDA/kg oil). As previously 749 mentioned, this behavior has been observed by other authors when studying the effect 750 751 of ascorbic acid on lipid oxidation in O/W emulsions, especially in presence of transition metals such as iron and copper (Mei et al., 1998; Uluata et al., 2015). Uluata 752 753 et al. (Uluata et al., 2015) provide an explanation related to the ability of AA to reduce metal ions, making them more reactive towards peroxides and hydroperoxides. 754 According to this proposed mechanism, reduced metallic species would decompose 755 peroxides and hydroperoxides into secondary oxidation products, increasing the 756 757 observed TBARS and preventing the accumulation of primary oxidation intermediaries 758 (Uluata et al., 2015). This behavior has been also observed in this work, although no 759 metals were added to the O/W emulsion. However, and according to inductively coupled plasma mass spectrometry (ICP-MS) analysis (Table S-2), metal traces are 760 761 present in the encapsulating material, enabling this hypothesis.

Additionally, it has been found that spray-dried and conventionally freeze-dried particles underwent secondary oxidation during the 28-day storage period, with final TBARS values of 137.2 ± 4.7 mg MDA/kg oil and 166.6 ± 0.3 mg MDA/kg oil, respectively (Fig. 4b days 1-28). Again, this high secondary oxidation status might be linked to the poorer encapsulation efficiency of those methods. On the other hand, freeze-dried particles frozen with liquid N₂ showed good stability against secondary

oxidation during storage, maintaining a TBARS value of 79.6 ± 2.4 during 28 days of storage at 4°C.

770

771 **4.** Conclusion

Particles from Gas-Saturated Solutions (PGSS)-drying has been used to encapsulate
omega-3 polyunsaturated fatty acids (*n*-3 PUFAs) into octenyl-succinic-anhydride
(OSA) starch, obtaining a solid powder with high bioactive load.

Similar encapsulation efficiencies (EE) and spherical morphologies have been obtainedby PGSS and spray-drying.

Freeze-dried particles showed irregular morphology. Slow conventional freezing destabilizes the O/W emulsion and negatively affects EE. DSC analysis of the PGSSdried particles successfully identified cold crystallization of the n-3 PUFA concentrate as well as gelatinization and melting peaks of OSA-starch.

PGSS-drying method offers low drying temperature and an intrinsically inert 781 782 atmosphere, which avoid oxidative degradation of n-3 PUFAs during processing, as demonstrated by the oxidative stability analyses. Conventional freeze-drying method 783 yielded particles with low oxidative stability, whereas freezing with liquid N₂ resulted 784 in a powder with oxidative stability comparable to PGSS-dried particles. Combined 785 with the addition of natural antioxidants such as ascorbic acid, the PGSS-drying 786 technique rises as a suitable method to formulate n-3 PUFAs in solid form and protect 787 them against oxidation during shelf life. 788

789

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Table 1. Summary of experimental results.

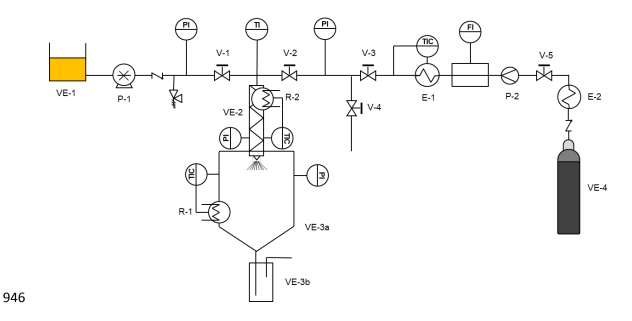
	Density (g·cm ^{·3})	Yield (%)	EE (day 0) (%)	Bioactive loading (mg/g)	Moisture (%)	Droplet size analysis			
Emulsion /drying method						D[4,3]	D[3,2]	d _{0.5} (μm)	span
Original						0.144 ^a	0.114 ^a	0.114 ^a	1.150 ^a
PGSS-drying	1.091281*	61 ± 1	$97.9^{b}\pm0.3$	191 ± 8	3.3 ± 0.3^{a}	0.227 ^b	0.116 ^a	0.134 ^b	2.197 ^d
Spray-drying		30 ± 1	$97.5^{\text{b}}\pm0.1$	187 ± 3	$5.6\pm0.2^{\circ}$	0.197 ^b	0.112 ^a	0.129 ^b	1.636 ^c
Freeze-drying (-20°C)		99 ± 1	$95.8^{c}\pm0.2$	192 ± 2	4.66 ± 0.05^{b}	0.567 ^c	0.121 ^b	0.131 ^b	1.772 ^c
Freeze-drying) (liq N ₂)		99 ± 1	$98.6^{a} \pm 0.1$	192 ± 2	$4.7\pm0.1^{\text{b}}$	0.146 ^a	0.107 ^a	0.118 ^a	1.291 ^b

* Standard uncertainty is $u(\rho) = \pm 0.000002 \text{ g} \cdot \text{cm}^{-3}$ a,b,c,d Different upper-scripts in the same column denote statistically significant differences at p < 0.05

- 943 944
- **Table 2.** Peak temperatures of the thermal events observed in the PGSS-dried powder loaded with n-3 PUFA concentrate (PGSS-drying), the carrier alone (Hi-CapTM 100), and the n-3 PUFA concentrate alone (AlgatriumTM Plus).

	Peak temperature (°C)						
Sample	Cold crystallization	Glass transition	Melting				
PGSS-dried particles	-72.99	76.57	223.55				
Ні-Сар™ 100	n.d.	78.83	217.05				
Algatrium [™] Plus	-71.42	n.d.	n.d.				

n.d.: not detected



948 Figure 1. Schematic diagram of the PGSS-drying apparatus. VE-1: O/W emulsion vessel, VE-2: static mixer, VE-3: a) spraying tower, b) condenser, VE-4: CO₂ vessel, P-1: O/W emulsion pump, P-2: CO₂
950 pump, V-: process valve, E-: heat exchanger, R-: electrical resistance, PI: pressure indicator, TI(C): temperature indicator (and controller).

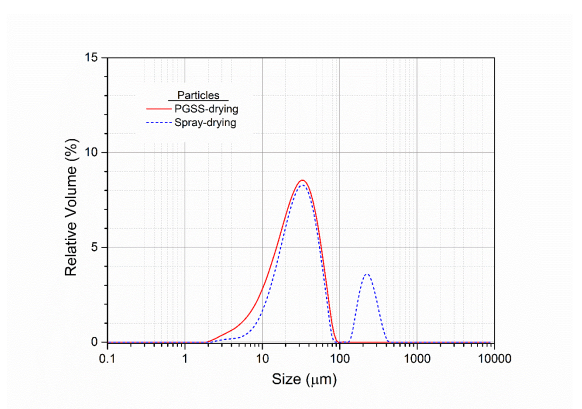
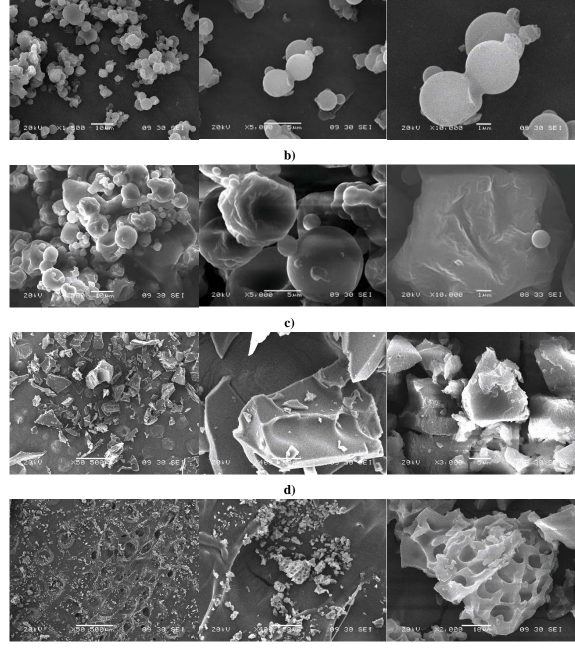


Figure 2. Particle size distribution plot of the particles obtained by PGSS-drying and by spray-drying.



a)

Figure 3. SEM micrographs of the dried powders. a) powder obtained by PGSS-drying, b) powder obtained by spray-drying; from left to right, 1500, 5000 and 10000x magnifications. c) powder obtained by conventional freeze-drying (50, 400 and 3000x). d) powder obtained by freeze-drying with liquid N₂; (50, 400 and 2000x).

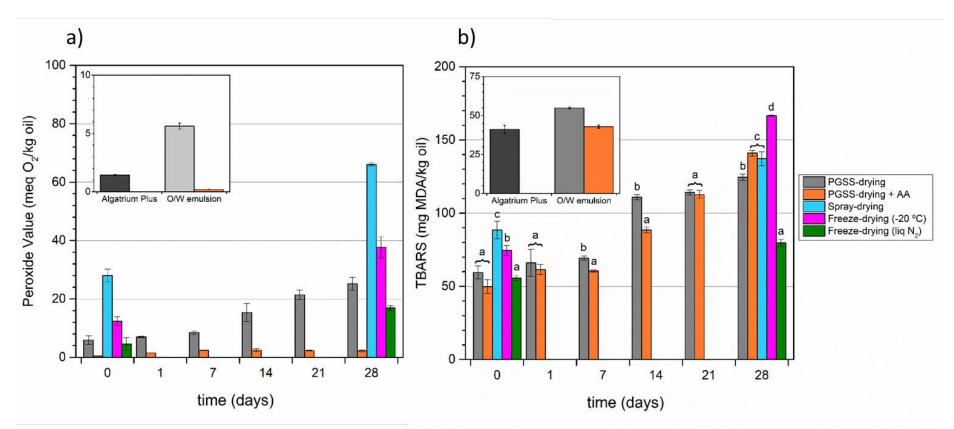


Figure 4. a) Peroxide Value (PV) and **b**) Thiobarbituric acid reactive substances (TBARS) content of the powders obtained by the different drying methods right after drying (day 0) and during storage at 4°C in darkness and ambient oxygen conditions (days 1-28). Samples were reconstituted the day of analysis keeping the water:carrier:n-3 PUFA concentrate proportion the same as the original (70:24:6 wt.). Different letters denote statistically significant differences at p < 0.05. **Insets:** PV and TBARS of the n-3 PUFA concentrate (AlgatriumTM Plus), and the original US-assisted O/W emulsions without and with ascorbic acid (AA).