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4	
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43 Abstract

44 Many studies indicate that food matrix microstructure and type of dietary oil or fat play a key role in carotenoid absorption. Therefore, this work was designed to highlight the 45 46 relationship between processed food microstructure and carotenoid absorption. The aim of 47 this study was to evaluate the consumption of a carotenoid-rich fruit snack on lipid profile, glycemia and especially on carotenoid absorption/bioconversion in Wistar rats. Animals 48 were fed with mixtures based on vacuum-fried papaya chips with either soy oil (PC-S) or 49 palm oil (PC-P) during 7 days, receiving 0.29 mg lycopene/kg/day and 0.35 mg total 50 51 carotenoids/kg/day. Lycopene and retinoids were analyzed in plasma and liver of rats by 52 HPLC-DAD. Results showed that the consumption of mixtures based on papaya chips did not affect the lipid profile or glycemia in rat plasma, regardless the type of oil. Wide-field 53 and confocal microscopy analyses of food matrix helped to understand why lycopene 54 55 accumulation in liver was higher (p<0.05) in rats fed with PC-P (0.442 µg/g liver) than in those fed with PC-S (0.291 µg/g liver). A better dissolution of crystalloid lycopene was 56 57 found in PC-P. Conversely, a higher bioconversion of provitamin A carotenoids was 58 observed for soy products. The effect of type of oil was underlined by epifluorescence microscopy of papaya mixtures showing homogeneous and small lipid droplets for soy 59 products. These results showed that PC-S could be recommanded as a healthy snack, being 60 61 a source of provitamin A carotenoids and bioavailable lycopene in a diversified diet.

62

63 Keywords

Food microstructure; Lycopene; Carotenoid bioconversion; Lipid profile; Vacuum frying;
Wistar rats.

66 1. Introduction

67 Over the last few decades, a dramatic rise of obesity and metabolic syndrome has been 68 occurred in Latin America and the Caribbean, leading to a higher incidence of non-69 communicable diseases (e.g. cardiovascular diseases and diabetes) which are the major 70 causes of death (Popkin, & Reardon 2018; Mattei et al., 2015). This is attributed to 71 nutrition transition (increase of unhealthy dietary habits), negative changes in lifestyle, and migration from rural to urban areas (Cuevas et al., 2011). For instance, during year 2016 in 72 73 Latin America and the Caribbean 58.8% and 60.1% of adult men and women, respectively, 74 were overweight or obese (OPS/OMS, 2019). Also, during the same year, in this region the 75 prevalence of high blood pressure and high blood glucose/diabetes for men and women was 23.8 and 8.9%, and 18.0 and 9.6%, respectively (OPS/OMS, 2019). Among numerous 76 factors, the high availability of energy-dense and nutrient-poor snack foods in low- and 77 78 middle- income countries represent a problem (Pries, Filteau, & Ferguson, 2019). Consumers in Latin America and the Caribbean are attracted by diets based on this kind of 79 80 foods that are composed mainly or solely of sugars and saturated lipids but slight or no 81 content in vitamins, minerals, protein, fiber or essential fatty acids (Poti, Slining, & Popkin, 82 2014).

The consumer trends towards healthier foods requires alternative strategies to the consumption of deep-fat fried products rich in simple sugars, saturated lipids and salt, e.g. potato chips, French fries, doughnuts, extruded snacks, cheese sticks, among others (Da Silva & Moreira, 2008). Numerous epidemiological studies promote the consumption of fruits and vegetables for the prevention of the non-communicable diseases (Angelino et al.,2019; Gan et al., 2015). Consequently, the intake of fruits and vegetables must be 89 increased in diets because they represent a source of bioactive health promoting compounds90 such as vitamins, polyphenols, carotenoids, and fiber.

91 Among phytochemicals, the carotenoids, such as lycopene, are known to be natural 92 antioxidants with beneficial health effects. These compounds enhance the functions of 93 immune system and lower the development of chronic diseases such as macular 94 degeneration, type 2 diabetes, obesity, cardiovascular diseases, and certain type of cancers (Cheng et al., 2017; Kopec & Failla, 2018). However, the most relevant function, is the 95 96 provitamin A activity. Some carotenoids such as β -carotene and β -cryptoxanthin have 97 provitamin A activity. This means they can be converted into retinol and other related 98 retinoids in organism, playing a key role on growth, visual cycle, and gene regulation (Kulczynski, Gramza-Michalowska, Kobus-Cisowska, & Kmiecik, 2017). 99

The application of technological processes is of utmost importance to obtain novel and 100 101 healthy products from fruits and vegetables. In this context, vacuum frying is an alternative 102 technology to produce fruit and vegetable-based snacks bearing the desired sensory quality 103 and better preserving their nutrients compared to the traditional fried snacks (Da Silva & 104 Moreira, 2008; Dueik & Bouchon, 2011). In addition, vacuum frying allows the use of 105 healthier unsaturated vegetable oils due to the low operation temperatures and the absence of oxygen during process (compared to atmospheric frying), thus minimizing oil 106 107 deterioration (Da Silva & Moreira, 2008). For instance, vacuum-fried papaya chips are a 108 good source of lycopene and provitamin A carotenoids such as β -cryptoxanthin and β -109 carotene (Soto et al., 2020). These chips and may be an alternative in Latin America and 110 the Caribbean to control vitamin A deficiency which remains a public health problem in 111 countries of this region (Cediel et al., 2015). However, the bioaccessibility and bioavailability of carotenoids in fruit and vegetable-based foods are influenced by various 112

factors. Especially, food matrix microstructure as well as the presence and type of oil have 113 114 a great impact (Lemmens, Colle, Van Buggenhout, Palmero, Van Loey, & Hendrickx, 115 2014; Schweiggert & Carle, 2017; Xavier & Mercadante, 2019). Carotenoids are enclosed 116 in cell organelle structures (e.g. chromoplasts in fruits). Thus, disruption of cell food matrix 117 during processing (thermal and mechanical treatments) may increase carotenoid bioavailability (Van Buggenhout et al., 2010). Otherwise, the presence of lipids play a role 118 in the transfer and diffusion of carotenoids during processing as well as during digestion, 119 120 since carotenoids need to be released from food matrix and incorporated into lipid emulsion 121 droplets and finally transferred to mixed micelles (Xavier & Mercadante, 2019). Location 122 and deposition forms of carotenoids in raw fruit/vegetable tissues have been widely studied in relation to their bioavailability (Schweiggert & Carle, 2017). In contrast, few studies 123 have investigated the impact of both processed food microstructure and oil presence on 124 125 carotenoid absorption and bioconversion.

126 From a nutritional point of view, this study was embedded in the general approach of 127 making a carotenoid-rich snack using papaya in order to offer a healthier product than 128 traditional fried snacks. Papaya fruit was chosen because: 1) it is a fruit widely spread and consumed in America Latina and the Caribbean (FAO, 2019; Saran, Solanki, & Choudhary, 129 130 2016), 2) it presents a varied profile of carotenoids, xanthophyll (β -cryptoxanthin) and carotenes (β-carotene and lycopene) (Soto et al., 2020) carotenoids are more bioavailable 131 132 from papaya than from other plant foods such as tomato and carrot (Schweiggert et al., 133 2014). The objective of the present study was to highlight the relationship between 134 processed food microstructure and carotenoid absorption/bioconversion in rats fed with a 135 mixture based on vacuum-fried papaya chips obtained with saturated or unsaturated oils. To better understand the effect of processing, freeze-dried papaya mixtures with addition of 136

oils were administred to rats comparatively to mixtures from papaya chips. Lipid profile
(triglycerides, total cholesterol, HDL and LDL) as well as glycemia were analyzed in
plasma while retinoids and lycopene were measured in liver and plasma of rats. Wide-field
and confocal microscopy tools were used to explain the results in relation to the processed
food microstructure.

142

143 **2.** Materials and methods

144 *2.1. Materials*

Red-fleshed papaya fruits (*Carica papaya* L. var. Formosa from Brazil) from a single batch
were acquired from TerreAzur (Montpellier, France) at ripening stage 4 (41-55% of skin
yellowing). Commercial frying oils, soy oil (Huileries Cauvin, Nimes, France) and
hydrogenated palm oil Risso® (Vandemoortele, Gent, Belgium) were used as described
previously by Soto et al. (2020b).

150

151 2.2. Obtention of papaya chips and mixtures for animals

Papaya chips were obtained after vacuum frying (120 °C, 13 min, 25 kPa) either with soy oil (~26 % oil in chips) or palm oil (~24 % oil in chips) and then were packaged in metallized PET/PE bags under nitrogen conditions and stored for 90 days at 25 °C as previously described by Soto et al. (2020). After this storage period the chips were ground (18%, w/w) and mixed with 82% (w/w) of distilled water to obtain the diet mixtures with the adequate viscosity to feed the rats via oral administration (Papaya chips - soy oil, PC-S, and Papaya chips - palm oil, PC-P).

On the other hand, papaya fruit slices were freeze-dried (Usifroid SMH 15, Élancourt,
France) during 72 h. Freeze-dried papaya was ground (14%, w/w), then mixed with 81%

161 (w/w) of distilled water and 5% (w/w) of the different oils (soy or palm oils) to obtain the 162 other diet mixtures (Freeze-dried papaya + soy oil, FDP+S, and Freeze-dried papaya + 163 palm oil, FDP+P) with the adequate texture as well. The formulation of different mixtures 164 was made considering the lycopene concentration (major carotenoid) and the lipid content 165 (Table 1) leading to products with a density of ~1.1 g/mL. Contents of lycopene, total carotenoids and lipids in the diet mixtures expressed per body weight (bw) of rats were 0.29 166 mg/kg bw/day, 0.35 mg/kg bw/day and 0.28 g/kg bw/day, respectively. The different 167 168 mixtures were prepared one week before the diet period and stored at -20 °C. Prior to oral 169 administration to the rats, the mixtures were thawed in a water bath at 37 °C.

170

171 *2.3. Animals*

172 Male Wistar rats (n=38), 7 weeks old, were obtained from Janvier Labs (S^t Berthevin, 173 France). Animals were handled in compliance with European Union animal care regulation rules and the guidelines of the National Institute of Health and the Committee for Animal 174 175 Care at the University of Montpellier (France). Animals were housed two by two in 176 plexiglass cages (enriched with wood toys) at 25 ± 1 °C, subjected to 12 h light/dark cycle and fed with standard diet A04 (SAFE, Scientific Animal Food and Engineering; Augy, 177 France) and water ad libitum. The retinol content in the standard diet was 7500 UI/kg. 178 During the adaptation period (4 days), rats were only fed with standard diet (A04) without 179 180 any carotenoid source.

181

182 *2.4. Study design*

The animals were randomly divided into 5 groups for the diet period (7 days) as shown in
Figure 1. One group of animals (n=6) received only the regular A04 diet (Control). The

185 other four groups (n=8) were orally fed with 1 mL of the different diet mixtures for 7 days 186 two times per day (morning, 8-9 am and afternoon, 3-4 pm) in addition to regular A04 diet. 187 Two groups were fed with PC-S or PC-P, and the other two groups treated with FDP+S or 188 FDP+P (Figure 1). At the end of the experimental period, rats were being fasted overnight 189 before anesthetized (20 μ L pentobarbital/100 g bw), and blood samples were collected by 190 cardiac puncture before euthanasia. For each blood sample an aliquot (500 µL) was taken for glycemia determination. Then, tubes with heparin containing the blood samples were 191 192 centrifuged (Jouan BR41 Multifunction, Thermo Electron Corporation, France) at 1500 193 rpm for 20 min at 15 °C to collect the plasma. Liver samples were collected and weighed after being washed with ice-cold saline solution (0.9 % NaCl, w/v), and then immediately 194 snap frozen in liquid nitrogen. All the samples were stored at -80 °C for further analyses. 195

196

197 2.5. Growth performance

Animals from different groups were weighted to determine their weight gain after adaptation and diet periods. Weights were recorded at the initial day of experimentation (day 0), at the end of adaptation period (day 4), and at the end of diet period (day 7) using a top-pan electronic balance.

202

203 2.6. Chemical analyses for papaya mixtures

Moisture and protein contents were determined by standard AOAC methods 920.151 and 920.152,
respectively (AOAC, 2015). Lipid content was determined by the method described by Carpenter et
al. (1993). Sugars (sucrose, glucose, and fructose) were determined using UPLC-1290 System
Infinity II (Agilent, CA, USA) equipped with a refractive index detector. Sugars were separated
using a Shodex SH1011 column (300 x 8 mm i.d., 6 µm) (Showa Denko K.K., Tokyo, Japan) with a

guard column, and the mobile phase was H₂O with H₂SO₄ (0.01%). The operation temperature was
set at 30 °C. The flow rate was set at 0.7 mL/min and the injection volume was 10 μL. Isocratic
condition was programmed with a run time of 20 min. Quantification was performed after obtaining
linear calibration curves of glucose, fructose, and sucrose. The fatty acid profile was determined
using AOAC standard method 996.06 (AOAC, 2015) and AOCS method Ce 1e-91 (AOCS, 2012).

214

215

216 2.7. Carotenoid analyses

217 2.7.1. Carotenoid extraction from papaya mixtures

Procedures and conditions for extraction of different papaya mixtures were described 218 previously by Soto et al. (2020b). Briefly, samples were weighed (700 mg) in 20 mL tubes. 219 220 Then, 2 mL of an ethanol solution containing 1 % pyrogallol was added. The mixture was 221 homogenized using a Vortex mixer and incubated for 2 min in the dark in a water bath at 70 222 °C. Then, after cooling saponification of samples was performed for 30 min in a water bath 223 at 70 °C by adding 1.5 mL of saturated KOH (12 N). After incubation, the tubes were 224 cooled in an ice bath and 2 ml of distilled water and 5 mL of hexane were added. Then, 225 after mixing and decantation, the aqueous phase was extracted twice with 5 mL of hexane. The organic phases were pooled and evaporated under nitrogen at 30 °C until dryness. 226 227 Finally, the residue was dissolved in 500 µL of methyl tert-butyl ether (MTBE)/methanol 228 (80/20) and placed in an amber vial prior to HPLC analysis.

229

230 2.7.2. Carotenoid and retinoids extraction from plasma

The carotenoids were extracted according to the method previously described by Poulaert et

al. (2014) with some modifications. The plasma previously obtained (700 μ L) was put into

a tube of 8 mL, then 500 μ L of ethanol 96 % (v/v) containing canthaxanthin (2 mg/mL) as internal standard and 2 mL of hexane were added. The mixture was homogenized (using a Vortex) for 60 s and then centrifuged (Allegra 21 Centrifuge, Beckman Coulter, Switzerland) at 1400 x g for 5 min at 25 °C. The organic phase was collected, then the aqueous phase was reextracted with 1 mL of ethanol and 2 mL of hexane. The organic phases were pooled and evaporated under nitrogen at 30 °C until dryness. The dried extract was dissolved in 100 μ L of CH₂Cl₂/Methanol (50:50, v/v) prior HPLC analysis.

240

241 2.7.3. Carotenoid and retinoids extraction from liver

242 The frozen liver was cut into small pieces on an ice-cold table. Then, 1 g of liver was mixed 243 with 1 mL phosphate-buffered saline (PSB) and homogenized (using a 15 mL Potter tissue 244 grinder) at 600-1000 rpm for 4 min. This step was done twice. Then the homogenates were 245 pooled and homogenized using an Ultra-turrax homogenizer (IKA T10 Basic, Germany) for 60 s. For carotenoids and retinyl palmitate extraction, 900 µL and 500 µL of liver 246 247 homogenates, respectively, were extracted using 500 µL of ethanol 96 % (v/v) containing 248 canthaxanthin (2 mg/mL) as internal standard and 2.5 mL or 2 mL of hexane for carotenoids and retinyl palmitate, respectively. The mixture was homogenized again using 249 250 a Vortex for 60 s and then centrifuged (Allegra 21 Centrifuge, Beckman Coulter, Switzerland) at 1600 x g for 5 min, at 25 °C. The organic phase was collected, and the 251 252 aqueous phase was reextracted with 1 mL of ethanol and 2 mL of hexane. The organic 253 phases were pooled and evaporated under nitrogen at 30 °C until dryness. The dried extract 254 was dissolved in 100 µL and 850 µL of CH₂Cl₂/Methanol (1 g/20 mL) for carotenoid and 255 retinyl palmitate analysis, respectively.

257 2.7.4. HPLC analysis in papaya mixtures

258 Carotenoid identification was performed by HPLC using a HPLC-DAD Agilent 1100 259 system (Massy, France). Carotenoids were separated using a C30 column (250 x 4.6 mm 260 i.d., 5 µm) (YMC EUROP GmbH, Germany) with a guard column, and the mobile phase 261 was H₂O as eluent A, methanol as eluent B, MTBE as eluent C. Operation temperature was 262 set at 25 °C. The flow rate was set at 1mL/min and the injection volume was 20 µL. The gradient program was described by Soto et al. (2020b). β -cryptoxanthin and β -carotene 263 264 were detected at 450 nm, and lycopene was detected at 470 nm. In papaya mixtures the β -265 carotene and lycopene contents were expressed as the sum of their all-E- and Z-isomers.

266

267 2.7.5. HPLC analysis in plasma and liver samples

268 Carotenoid and retinoids were separated with the same C30 column as previously described in 269 section 2.6.4. The mobile phase was H₂O as eluent A, methanol as eluent B, and MTBE as eluent C. 270 Temperature was set at 25 °C. The flow rate was set at 1mL/min and the injection volume was 60 271 μL. A solvent gradient was programmed as follows: 2% A-96% B-2% C (initial conditions); 0-27 272 min, 2% A-18% B-80% C; 27-35 min, 4% A-11% B- 85 C% and back to the initial conditions for 273 re-equilibration. Chromatograms were generated at 325 nm to identify retinol and retinyl esters, and 274 at 470 nm for lycopene identification. Carotenoid and retinoids were identified by comparing their 275 retention time and spectra with the respective standards. Quantification was achieved by 276 establishing calibration curves with all-E-lycopene, retinol and retinyl palmitate, being the 277 determination coefficients 0.994, 0.997 and 0.998, respectively. For carotenoid HPLC analysis 278 the limit of detection (LOD) was 0.0040 µg, and the limit of quantification (LOQ) was 279 0.0150 µg.

282 *2.8. Glycemia*

Determination of glucose was performed in blood samples using the Accu-Chek Performa
blood glucose meter (Roche, Basel, Switzerland).

285

286 2.9. Lipid profile and free fatty acids in plasma

Lipid profile analyses were performed in plasma: triglycerides (TRIG), total cholesterol (CHO); low-density lipoprotein (LDL) and high-density lipoprotein (HDL) using the Biolabo enzyme kits (Biolabo SAS, Maizy, France). Content of free fatty acids (FFA) was determined using the Abcam enzyme kit (Allscience, Miami, USA). These analyses work under colorimetric reactions, for which a Spark 96-plate micro spectrophotometryfluorescence device was used (TECAN, Männedorf, Switzerland).

293

294 2.10. Microscopy analyses

295 Mixtures from vacuum-fried papaya chips and freeze-dried papaya were directly observed 296 with a wide-field microscope Eclipse Ni-E (Nikon Instruments Inc., NY, USA). The pictures were obtained with the 20X Plan-APO 0.75 NA objective under transmitted light 297 or differential interference contrast (DIC). Size distribution of lipid droplets, expressed as 298 299 D₉₀, was calculated by using ImageJ v1.8.0 software. D₉₀ means that 90% of spherical 300 particles have a diameter less than the specified value (µm). The lipids were visualized in 301 fluorescence with the lipophilic stain Nile Red $(3.14 \,\mu\text{M})$ with a blue excitation filter (B2A: 302 450-490 nm, long-pass emission 505 nm).

The same mixtures and the papaya chips were observed with a confocal microscope Zeiss
880 (Zeiss, Jena, Germany) with a 488 Argon laser with an objective 20X Plan APO 1.0

305 NA to visualize the autofluorescence of carotenoids (lycopene and β -carotene/ β -306 cryptoxanthin).

307 Spectral analysis was carried out using the advanced Linear Unmixing function (LSM 880 308 software, Zeiss) which separates mixed signals pixel by pixel using the entire emission 309 spectrum of each defined autofluorescent compound in the sample. This function was 310 applied with the advanced iterative option and one residual channel. After spectral imaging acquisitions on samples, this Linear Unmixing function allowed visualization with coded 311 312 colors of the fluorescence of each standard (lycopene, β -carotene, and β -cryptoxanthin) 313 based on their reference spectra (Talamond, Verdeil, & Conéjéro 2015). The spectral 314 detector of this microscope was used to obtain the emission spectra of lycopene and β-315 carotene/ β -cryptoxanthin and visualize specifically these molecules in the samples between 316 500 and 690 nm.

317

318 2.11. Statistical analyses

Results obtained from plasma and liver samples were analyzed by one-way analysis of variance (ANOVA) after examining for homogeneity of variances by Bartlett's test. Pvalue <0.05 was considered significant. If significantly different, means were further compared using Fisher's test (p<0.05). Composition of the different papaya mixtures was also compared by one-way ANOVA and post hoc Tukey-HSD (p<0.05). Statistical analyses were performed using the XLSTAT Software version 2020 (Addinsoft Inc, USA).

325

326 **3. Results and discussion**

327 3.1 Carotenoid consumption and growth performance

328 Table 1 shows the composition of the different papaya mixtures (PC-S, PC-P, FDP+S, 329 FDP+P) that were made to feed the rats by oral administration. There were no significant 330 differences (p>0.05) in lycopene, total carotenoid, and lipid contents among the different 331 papaya mixtures. During the diet period rats were orally fed with 1 mL of the different 332 papaya mixtures (PC-S, PC-P, FDP+S, FDP+P) two times per day, which means 2 mL 333 papaya mixture/day. Therefore, the animals fed with mixtures consumed per day ~96 µg of lycopene, ~113 µg of total carotenoids, and ~90 mg of lipids (either from soy or palm oils). 334 Mixtures from papaya chips (PC-S and PC-P) presented around 2-fold lower (p<0.05) 335 336 contents of β -cryptoxanthin and β -carotene and thus of vitamin A (expressed as Retinol 337 Activity Equivalent, RAE) than those in mixtures from freeze-dried papaya (FDP+S and FDP+P). This was explained by thermal degradation of these carotenoids during vacuum 338 frying process (Soto et al., 2020). 339

340 Daily lycopene intake in this study was similar to that applied in previously studies realized

in rats by Jain et al., (1999) (142 μ g/day), and in gerbils by Mills et al., (2007) (60 μ g/day).

In most studies with rats, in which are analyzed the absorption, bioconversion or the health effects of carotenoids, the authors usually apply much higher doses of carotenoids, in diets, to feed animals. For instance, some studies reported a daily lycopene intake that ranged from 720 to 10.000 μ g (Takayama et al., 2013; Shalaby, & Fouad, 2019; Yilmaz et al., 2018; Xu et al., 2019).

Rats gained weight during the experimental period and the final body weight ranged from
326.6 g to 346.6 g (Table 2). However, after the diet period there were no significant
differences (p>0.05) in weight gain between rats consuming papaya mixtures (PC-S, PC-P,
FDP+S, FDP+P) and Control group (Table 2).

352 3.2 Glycemia

353 Figure 2 shows the results of glycemia measured in rats from Control group and the rats fed 354 with the different papaya mixtures (PC-S, PC-P, FDP+S, FDP+P). The glycemia of rats fed 355 with mixtures from papaya chips (185.75 \pm 34.56 mg/dL for PC-S, and 178.13 \pm 30.71 356 mg/dL for PC-P) was not significantly different from Control group (160.67 \pm 20.70 357 mg/dL). Unexpectedly, the glycemia increased in rats consuming freeze-dried papaya mixtures (223.00 \pm 48.09 mg/dL for FDP+S, and 198.50 \pm 36.98 mg/dL for FDP+P). These 358 359 values were significantly higher (p < 0.05) than those in Control group. These differences in 360 glycemia could be attributable to the higher contents of glucose and fructose present in freeze-dried papaya mixtures (FDP+S, FDP+P) (~98% of total sugars) compared to the 361 mixtures made with papaya chips (PC-S, PC-P) (~30% of total sugars) (Table 1). During 362 vacuum frying of papaya, the degradation of glucose and fructose was observed whereas 363 364 the formation of sucrose occurred (Soto et al., 2020c). According to these authors, factors related to frying process (high temperature and fast rate of water loss) and intrinsic 365 characteristics of papaya (proximal composition, sugar concentration, organic acids, 366 367 acidity) are involved in sucrose formation.

368

369 **3.3 Lipid profile and free fatty acids in plasma**

Table 3 shows the lipid profile and content of free fatty acids (FFA) in plasma of rats fed with the papaya mixtures (PC-S, PC-P, FDP+S, FDP+P) and the Control group. No significant differences (p>0.05) were found in cholesterol determinations (total, CHO; high-density lipoprotein, HDL; and low-density lipoprotein, LDL) among rats fed with papaya chip mixtures (PC-S, PC-P) and rats from Control group. Similarly, there were no significant differences (p>0.05) for CHO, HDL and LDL among Control group and rats fed
with freeze-dried papaya mixtures (FDP+S, FDP+P) (Table 3).

Moreover, the level of triglycerides (TRIG) in plasma was similar among rats of Control group and rats fed with papaya chips mixtures (PC-S, PC-P) and those fed with FDP+S mixture. Rats consuming FDP+P mixture presented the highest TRIG level (p<0.05). These results demonstrated that consumption of mixtures based on papaya chips (PC-S, PC-P) did not increase CHO, HDL, LDL or TRIG in rat's plasma, regardless the type of oil.

Only FFA increased significantly (p<0.05) in plasma of rats consuming papaya chips mixtures (PC-S, PC-P) compared to the Control group. The heat treatment (120 °C) applied during vacuum frying of papaya could induce some hydrolysis of triglycerides from frying medium (soy and palm oils) (Chung, Lee, & Choe, 2004). Thus, the oil absorbed in papaya chips (PC-S, PC-P) during process could alter the levels of FFA in plasma of rats. On the other hand, the mixtures based on freeze-dried papaya (FDP+S, FDP+P) were made with fresh oils without any thermal degradation.

389

390 3.4 Lycopene absorption

Lycopene was detected in liver of rats fed with the different mixtures based on freeze-dried 391 392 papaya (FDP+S, FDP+P) and papaya chips (PC-S, PC-P) (Figure 3). Significant differences 393 (p<0.05) were observed between groups of rats fed with FDP+S and PC-S. Lycopene 394 accumulation in the liver was higher in rats fed with FDP+S (0.522 µg/g liver) than in those 395 fed with PC-S (0.291 µg/g liver). To support these results, wide-field microcopy of the 396 mixtures was performed and revealed that FDP+S mixture exhibited a fine emulsion with 397 numerous droplets (the D₉₀ statistical diameter was 18 µm) probably due to the soy oilbased formulation (Figure 4.a). In opposite, the oil droplet size of PC-S mixture was higher 398

399 (the D_{90} statistical diameter was 46 μ m) and crystalloid remnants of lycopene were 400 identified (Figure 4.b, see arrows). A better dissolution of crystalline lycopene in the soy oil 401 fine emulsion was hypothesized.

402 To confirm the identification of carotenoids, the spectral imaging of confocal microscopy 403 allowed to identify these compounds with fluorescence spectra as well as to locate the 404 carotenoids in papaya mixtures. In the FDP+S mixture, the two type of carotenoids β -405 carotene/ β -cryptoxanthin (provitamin A carotenoids) and lycopene seemed to dissolve in 406 the core of lipid droplets of emulsion (Figure 4.c). Inversely, only fluorescence emission 407 spectra of lycopene were found in PC-S mixture within solid aggregates (Figure 4.d, see 408 arrowheads).

409 According to Schweiggert et al. (2012) and Schweiggert & Carle (2017), lycopene from 410 papaya occurs in a solid crystalline deposition form whereas β -carotene/ β -cryptoxanthin are 411 liquid-crystalline or lipid-dissolved in globular-tubular substructure of chromoplast. The morphology of chromoplast and deposition form strongly influence carotenoid 412 413 bioavailability. Also, the storage of lycopene crystalloid form was associated with a lower 414 bioaccessibility and absorption (Schweiggert et al., 2014). The addition of soy oil and water to freeze-dried papaya (FDP+S) generated a fine emulsion favorizing the dissolution of 415 416 small crystalline lycopene, thereby increasing its absorption during digestion (Salvia-417 Trujillo et al., 2017). Indeed, before carotenoid absorption the lipid digestion rate was 418 higher for a fine emulsion with small droplets contributing to micelle formation, which 419 represents the absorbable form for enterocytes (Salvia-Trujillo et al., 2019). In the case of 420 PC-S, the formed emulsion was coarse with larger oil droplets and crystalloid remnants of 421 lycopene (Figure 4.b).

422 On the other hand, significant differences (p<0.05) in lycopene absorption were observed 423 between rats fed with PC-S mixture (0.291 μ g/g liver) and those with PC-P mixture (0.422 424 μ g/g liver). As shown in Figure 5, crystalloid lycopene form remained in PC-S (Figure 5.a). 425 This figure shows specific red color aggregates (lycopene) within the food matrix. Whereas 426 lycopene seemed to be better dissolved in the matrix of PC-P. In fact, the red color 427 represents a diffuse veil within the food matrix (Figure 5.b). Lycopene transfer to oil during vacuum-frying of papaya fruit could have been better with palm oil than with soy oil. The 428 429 fatty acid composition of oils (chain length and degree of unsaturation) influences the 430 incorporation of lycopene into micelles (Lemmens et al., 2014). Lycopene bioaccessibility or absorption was reported higher with olive oil and could be explained by the presence of 431 432 C18:1 (Clarke et al., 2000; Colle et al., 2012; Nagao et al., 2013). In addition, fatty acids with medium-chain length such as C16:0 increased lycopene bioaccessibility compared to 433 434 long-chain length and this was particularly significant when up to 5% of lipids was added. Medium-chain fatty acids are hydrolyzed to a higher extent than those with long-chain 435 436 length. A complete digestion of medium-chain fatty acids led to better lycopene 437 solubilization capacity (Colle et al., 2012). This can also explain why upon addition of sunflower oil up to 5%, lycopene bioaccessibility from papaya remained unaffected 438 (Schweiggert et al., 2012). 439

Palm oil contained 39% of C18:1and 41% of C16:0 whereas soy oil only contained 26% and 11%, respectively. Therefore, the fact that palm oil has a different fatty acid composition, added to the effect of processing, could explain the difference for lycopene absorption in rats consuming the papaya chips but processed with different oil. Furthermore, saturated coconut oil (rich in medium-chain triglycerides) enhanced lycopene tissue accumulation to a greater degree than safflower oil (rich in long-chain triglycerides)

in Mongolian gerbils fed with tomato powder (Conlon et al., 2012). Borel et al. (1996)
pointed out that the solubility of carotenes in bulk triglycerides increased with decreasing
of fatty acid chain length.

449

450 3.5 Provitamin A carotenoid bioconversion in plasma and liver

451 The bioconversion of pro-vitamin A carotenoids was demonstrated by an increase of retinol 452 in plasma and retinyl palmitate content in liver of rats consuming papaya mixtures (PC-S, 453 PC-P, FDP+S, FDP+P) comparatively with those of Control group fed with a regular diet 454 (non-supplemented with carotenoids). Retinol content in plasma was significantly higher (p<0.05) in rats fed with the papaya mixtures (PC-S, $3.05 \pm 0.20 \mu$ mol/L; PC-P, 2.80 ± 0.31 455 μ mol/L; FDP+S, 3.11 ± 0.38 μ mol/L; FDP+P, 2.87 ± 0.14 μ mol/L) than rats of Control 456 457 group $(2.38 \pm 0.25 \mu \text{mol/L})$. However, no significant differences (p>0.05) were found in 458 retinol content among the group of rats fed with the different papaya mixtures (PC-S, PC-P, FDP+S, FDP+P). 459

460 Figure 6 shows that contents of retinyl palmitate following administration of mixtures with 461 palm oil (PC-P, FDP+P) were lower than those in soy oil mixtures (PC-S, FDP+S). The better bioefficacy observed after ingestion of soy products could be explained by the 462 different behavior of emulsions formed in mixtures administrated to the rats. Wide-field 463 464 epifluorescence microscopy with help of lipid coloration showed emulsions with small and 465 round-shape oil droplets for the soy products (Figure 7.a, and Figure 7.c) in opposition to 466 emulsions with larger and undefined-shape oil droplets for the mixtures with palm oil 467 (Figure 7.b, and Figure 7.d).

468 Absorption of carotenoids requires their liberation from food matrix and their solubilization469 into oil droplets to be then transferred to bile salts mixed micelles. According to Salvia-

470 Trujillo (2013), the droplet size of oil in water emulsions generated during the first step of 471 digestion was one of the most important factors in carotenoid absorption because of the 472 hydrolysis of triglycerides. Digestion of triglycerides and carotenoid bioaccessibility were 473 higher for small lipid droplets than for bigger ones (Salvia-Trujillo, 2013). In addition, the 474 lipid digestion of long-chain fatty acids such as acid oleic and linoleic acids induced the 475 assembly and basolateral secretion of chylomicrons for their transfer to the lymph (McClements, Li, & Xiao, 2015). Failla et al. (2014) showed that unsaturated fatty acids 476 477 from soy oil promoted the uptake and secretion of β -carotene by Caco-2 cells.

478 It was also observed a processing effect between vacuum-fried papaya chips mixtures (PC-479 S, PC-P) and freeze-dried papaya mixtures (FDP+S, FDP+P). In fact, the bioconversion of provitamin A carotenoids was generally better when vacuum-fried chips products were 480 ingested (average of retinyl palmitate was 161 μ g/g liver for papaya chips mixtures versus 481 482 114µg/g live for freeze-dried papaya mixtures). This is relevant because the vitamin A content (RAE) in mixtures based on papaya chips was around 2-fold lower than in freeze-483 484 dried papaya mixtures (Table 1). This result underlined that vacuum frying process 485 favorized the carotenoid bioconversion. The incorporation of oil into the papaya matrix during vacuum frying (120 °C, 13 min, 25 kPa) could play a major role on the absorption of 486 provitamin A carotenoids compared to freeze drying with subsequent oil addition. It was 487 488 reported that heat treatments increased carotenoid bioavailability especially when lipids 489 were absorbed into the food matrix during deep-fat frying or vacuum frying (Lemmens et 490 al., 2011; Berni et al., 2015; Tumuhimbise, Namutebi, & Muyonga, 2009). Mutsokoti et al. 491 (2016) showed that high temperatures (>100°C) and short time process (10 min) were 492 sufficient to observe a maximal carotenoid transfer to oil. These conditions were similar to those used during vacuum frying of papaya. The increase of carotenoid solubility due to 493

494 vacuum frying of papaya fruit resulted in a higher bioconversion of provitamin A495 carotenoids and accumulation of lycopene in liver of rats.

496

497 4. Conclusions

498 Our study showed that a carotenoid-rich healthy snack obtained by vacuum frying did not 499 increase glycemia, cholesterol or triglycerides in rat plasma regardless the type of oil. Moreover, the consumption of mixtures based on vacuum-fried papaya chips favorized the 500 501 absorption of lycopene and provitamin A carotenoids in rats. It was shown that carotenoid 502 absorption depends on the type of oil and processing as well as carotenoid deposition in food microstructure. The crystalloid form of lycopene in papaya was better dissolved in 503 504 palm oil during vacuum frying and was better absorbed in rats. This could be explained by 505 the presence of C18:1 and medium-chain fatty acids such as C16:0 in palm oil. Inversely, 506 consumption of mixtures based on vacuum-fried chips with soy oil increased the 507 bioconversion of provitamin A carotenoids in liver of rats. The lipid-dissolved form of 508 provitamin A carotenoids associated with the presence of unsaturated soy oil (rich in 509 linoleic and α -linolenic acids) in vacuum-fried papaya chips favorized the formation of a fine emulsion in diet mixture. This facilitated the formation of micelles and consequently a 510 511 better carotenoid absorption in rats. Finally, these results showed that vacuum-fried papaya 512 chips obtained with soy oil represent an interesting source of provitamin A carotenoids and 513 bioavailable lycopene in a diversified diet. . In other words for the contributions of pro-514 vitamin A carotenoids, a portion of 25g of vacuum-fried papaya chips presented a 515 nutritional value of 75 µg Retinol Activity Equivalent (RAE) which corresponds to 10 % of 516 the recommended daily allowance (RDA) (800 µg for adults). It could be claimed that food products with 10% or more of the RDA for vitamin A (RAE) as "good source of vitamin 517

518	A" (FDA, 2020). Vacuum-fried papaya chips could be an alternative to offer the consumers
519	a healthier product than traditional snacks. For intance, 25-30 g could be a portion for
520	papaya chips (this correspond to classic commercial potatoes chips for individual portion).
521	According to Soto et al. (2020) 25 g of papaya chips contains 6.4 g of lipids (which
522	corresponds to 9.8% of daily intake recommended for lipids) and 3.9 mg of carotenoids.
523	Daily intake recommended for carotenoids does not exist; however, 4.8 mg/day is needed
524	to meet the requirement of 800 μ g of vitamin A for an adult (Toti et al ., 2018).
525	

527 Declaration of competing interest

528 No conflict of interest involving any of the authors.

529

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711

712 Figure captions

Figure 1. Study design for evaluating the effect of the consumption of different papaya
mixtures on carotenoid absorption and lipid profile in plasma and liver of Wistar rats.

715

Figure 2. Glycemia of Control group and rats fed with papaya mixtures: vacuum-fried papaya chips with either soy oil (PC-S) or palm oil (PC-P), and freeze-dried papaya mixed with either soy oil (FDP+S) or palm oil (FDP+P). Bars are expressed as the means \pm standard error of the mean, SEM (n=6 for Control, n=8 for the other groups). Means with the same letter are not significantly different (Fisher's t-test, p<0.05).

721

Figure 3. Lycopene content in liver of Wistar rats fed with papaya mixtures: vacuum-fried papaya chips obtained with either soy oil (PC-S) or palm oil (PC-P), and freeze-dried papaya mixed with either soy oil (FDP+S) or palm oil (FDP+P). Bars are expressed as the means \pm standard error of the mean, SEM (n=8). Means with the same letter are not significantly different (Fisher's t-test, p<0.05).

727

Figure 4. Wide-field and confocal microscopy of papaya soy mixtures for feeding rats. (A)
Wide-field microscopy (DIC 20 X objective) of freeze-dried papaya soy mixture (FDP+S).
(B) Wide-field microscopy (DIC 20 X objective) of vacuum-fried papaya chips soy mixture
(PC-S). (C) Spectral imaging by confocal microscopy of FDP+S mixture (red line,
lycopene; green line, β-carotene/β-cryptoxanthin). (D) Spectral imaging by confocal

microscopy of PC-S mixture (red line, lycopene). *Arrows and arrowheads* mark crystalloid
remnants of lycopene. The scale bars represent a length of 100 µm.

735

Figure 5. Spectral imaging by confocal microscopy (20X objective) of (A) Vacuum-fried papaya chips with soy oil, (B) Vacuum-fried papaya chips with palm oil. Lycopene is represented by red color and β -carotene/ β -cryptoxanthin by green color. The scale bars represent a length of 50 µm.

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Figure 6. Retinyl palmitate content in liver of Control group and rats fed with papaya mixtures: vacuum-fried papaya chips obtained with either soy oil (PC-S) or palm oil (PC-P), and freeze-dried papaya mixed with either soy oil (FDP+S) or palm oil (FDP+P). Bars are expressed as the means \pm standard error of the mean, SEM (n=6 for Control, n=8 for the other groups). Means with the same letter are not significantly different (Fisher's t-test, p<0.05).

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748 Figure 7. Wide-field epifluorescence microscopy (20X objective-Nile Red coloration 749 fluorescent stain for lipids) of papaya mixtures for feeding rats. (A) Vacuum-fried papaya 750 chips soy mixture (PC-S). (B) Vacuum-fried papaya chips palm mixture (PC-P). (C) Freeze-dried papaya soy mixture (FDP+S). (D) Freeze-dried papaya palm mixture 751 752 (FDP+P). The 100 scale bars represent a length of μm.

754 Figure 1



Figure 2



Figure 3



- , , ,

798 Figure 4



810 Figure 5



50 µm

813 Figure 6



830 Figure 7



Table 1. Composition of the different papaya mixtures used for feeding the Wistar rats.

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841

Component	Papaya mixtures					
Component	PC-S	PC-P	FDP+S	FDP+P		
Moisture (mg/mL)	906.38 ± 0.73^{b}	910.71 ± 0.94^{a}	$903.97 \pm 0.74^{\circ}$	909.66 ± 0.95^{a}		
Lipids (mg/mL)	44.67 ± 3.65^{a}	43.93 ± 1.03^{a}	45.01 ± 2.42^{a}	46.00 ± 1.81^{a}		
Protein (mg/mL)	8.10 ± 0.96^{a}	8.19 ± 1.44^{a}	4.90 ± 0.12^{b}	4.66 ± 0.10^{b}		
Carotenoids (µg/mL):						
β -cryptoxanthin	3.01 ± 0.33^{b}	2.50 ± 0.13^{b}	5.67 ± 0.53^{a}	5.38 ± 0.50^{a}		
β-carotene	2.73 ± 0.27^{b}	2.38 ± 0.11^{b}	6.28 ± 0.42^{a}	5.89 ± 0.43^{a}		
Lycopene	49.50 ± 5.28^{a}	46.44 ± 5.12^{a}	48.50 ± 5.45^{a}	47.30 ± 4.47^{a}		
Total carotenoids	55.24 ± 5.41^{a}	51.31 ± 5.15^{a}	60.45 ± 5.77^{a}	58.57 ± 5.31^{a}		
Total carotenoids (µg/mg FM)	1.24 ± 0.12^{a}	1.17 ± 0.12^{a}	1.34 ± 0.13^{a}	1.27 ± 0.12^{a}		
Vitamin A content (µg RAE/mL)	0.32 ± 0.03^{b}	0.27 ± 0.02^{b}	0.70 ± 0.02^{a}	0.65 ± 0.05^{a}		
Sugars (mg/mL):						
Sucrose	71.24 ± 4.25^{a}	69.72 ± 2.95^{a}	1.79 ± 0.06^{b}	1.89 ± 0.12^{b}		
Glucose	14.55 ± 0.89^{b}	15.55 ± 1.09^{b}	49.07 ± 0.42^{a}	50.98 ± 0.83^{a}		
Fructose	13.85 ± 0.74^{b}	14.52 ± 0.90^{b}	47.84 ± 0.56^{a}	49.79 ± 0.83^{a}		
Fatty acid profile (mg/mL):						
Saturated	$7.45 \pm 0.29^{\circ}$	21.04 ± 0.30^{b}	$7.06 \pm 0.10^{\circ}$	22.80 ± 0.10^{a}		
Monounsaturated	12.95 ± 0.22^{b}	17.20 ± 0.36^{a}	12.39 ± 0.55^{b}	18.14 ± 0.32^{a}		
Polyunsaturated	23.44 ± 0.43^{b}	$4.86 \pm 0.21^{\circ}$	25.01 ± 0.45^{a}	$5.04 \pm 0.26^{\circ}$		

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845 PC-S, papaya chips with soy oil; PC-S and PC-P, vacuum-fried papaya chips obtained with 846 soy and palm oil, respectively; FDP+S and FDP-P, freeze-dried papaya mixed with soy and 847 palm oil, respectively. FM, fat matter. Values are expressed as the means ± standard 848 deviation, SD (n=3). Means in the same row with the same letter are not significantly different (Tukey's test, p<0.05).Vitamin A content is expressed as Retinol Activity 849 850 Equivalent (RAE). RAE estimate was calculated for a bioconversion ratio 851 (carotenoid:retinol) of 12:1 for all-E-β-carotene, and 24:1 for all-E-β-cryptoxanthin and Zβ-carotene (US IOM, 2000). 852

Table 2. Body weight and weight gain of Wistar rats fed with papaya mixtures and Control

group at different periods of the study.

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W 7. * .14	Control –	Papaya mixtures			
weight		PC-S	PC-P	FDP+S	FDP+P
Body weight (g):					
Initial	254.8 ± 4.5	252.6 ± 2.5	267.8 ± 2.9	259.1 ± 4.1	253.0 ± 3.8
After adaptation period (4 days)	300.9 ± 4.0	290.7 ± 3.9	306.1 ± 2.0	297.1 ± 5.1	288.5 ± 4.3
After diet period (7 days)	346.6 ± 5.0	326.6 ± 6.2	343.9 ± 5.6	337.3 ± 7.7	331.5 ± 5.3
*Weight gain (g):					
During adaptation period (4 days)	46.2 ± 2.2^{a}	38.1 ± 3.9^{b}	38.3 ± 2.2^{ab}	$38.0 \pm 2.8^{\text{b}}$	35.5 ± 0.8^{b}
During diet period (7 days)	45.7 ± 1.4^{a}	$35.9 \pm 3.4^{\mathrm{a}}$	37.9 ± 4.7^{a}	40.1 ± 3.7^{a}	42.9 ± 3.4^{a}

857 Control, carotenoid deficient diet; PC-S and PC-P, vacuum-fried papaya chips obtained 858 with soy and palm oil, respectively; FDP+S and FDP-P, freeze-dried papaya mixed with 859 soy and palm oil, respectively. Values are expressed as the means \pm standard error of the 860 mean, SEM (n=6 for Control, n=8 for the other groups). *For weight gain, means in the 861 same row with the same letter are not significantly different (Fisher's t-test, p<0.05).

Table 3. Lipid profile and free fatty acids in plasma from Wistar rats fed with papayamixtures and Control group.

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Papaya mixtures	CHO (mg/dL)	HDL (mg/dL)	LDL (mg/dL)	TRIG (mg/dL)	FFA (µmol/uL)
Control	90.05 ± 1.7^{ab}	20.80 ± 3.10^{a}	14.35 ± 1.32^{a}	70.69 ± 6.06^{b}	27.70 ± 9.41^{b}
PC-S	92.05 ± 4.9^{a}	27.06 ± 2.90^{a}	15.77 ± 1.05^{a}	59.00 ± 3.25^{b}	64.93 ± 14.7^{a}
PC-P	78.82 ± 2.78^{b}	19.91 ± 3.17^{a}	13.80 ± 1.48^{a}	65.37 ± 4.15^{b}	58.61 ± 7.34^{a}
FDP+S	85.52 ± 2.99^{ab}	22.69 ± 1.88^{a}	14.93 ± 1.21^{a}	67.59 ± 5.12^{b}	43.14 ± 2.52^{ab}
FDP+P	93.43 ± 3.91^{a}	21.47 ± 1.26^{a}	16.06 ± 1.64^{a}	92.95 ± 5^{a}	40.37 ± 5.26^{ab}

865 Control, carotenoid deficient diet; PC-S and PC-P, vacuum-fried papaya chips obtained 866 with soy and palm oil, respectively; FDP+S and FDP-P, freeze-dried papaya mixed with 867 soy and palm oil, respectively. CHO, total cholesterol; LDL, low density lipoprotein; HDL, 868 high density lipoprotein; FFA, free fatty acids; TRIG, triglycerides. Values are expressed as 869 the means \pm standard error of the mean, SEM (n=6 for Control, n=8 for the other groups). 870 Means in the same column with the same letter are not significantly different (Fisher's t-871 test, p<0.01).

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