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1	Bee pollen as a natural antioxidant source to prevent lipid oxidation in black pudding
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19	Abstract: The antioxidant activity of bee pollen (mainly composed by Cistus ladanifer pellets)
20	was explored in the context of black pudding production. For this purpose, three black pudding
21	formulations comprising varying antioxidant compounds (sodium ascorbate, bee pollen and bee
22	pollen extract) were produced.
23	Bee pollen was characterized according to the botanical origin, antioxidant activity, total phenol
24	and flavonoid contents and phenolic profile. Black pudding was characterized by the
25	microbiological safety, lipid oxidation, pH, water activity and humidity for 1, 10, 21, 30 and 37
26	days. Sensory acceptance was evaluated on the four first periods of storage. Salmonella spp.,
27	Escherichia coli and Listeria monocytogenes were absent in all samples. Small variations on
28	humidity and pH were observed during the black pudding's storage. Regarding lipid oxidation,
29	it increased, on average, from 1.36 mg to 2.11 mg malondialdehyde/kg meat. Differences
30	among the three formulations were only significant on the first days of storage. The sensory
31	assessment did not differ between products. This study suggests that bee pollen may be used as
32	a natural antioxidant in meat products, yet a careful labelling is essential to alert allergic
33	consumers.
34	Key words: black pudding, pollen, antioxidant, lipid oxidation

37

38 Introduction

The safety and quality of food products are some of the main concerns of health agencies and consumers worldwide. Also, the consumers are increasingly demanding for a diverse range of food options particularly those containing biologically active ingredients with health promoting capacities and free of food additives. However, for many food products, like those containing animal derivatives, the lipid oxidation is an important source of quality deterioration, reducing their shelf lifetime and impairing its consumption (Jayawardana et al., 2011; Morrissey, Sheehy, Galvin, Kerry, & Buckley, 1998; Shah, Bosco, & Mir, 2014).

46 Black pudding is a meatless sausage containing pork blood as a main ingredient and is a product of excellence in the traditional Portuguese charcuterie called "morcela de assar". Blood 47 48 sausages are produced and consumed throughout Europe and each Region reveal their own 49 specificity and tradition. However, all black pudding are based on pork blood as its main raw 50 material. In this case, the raw material is considered an important source of nutrients because meat derived products contain high amounts of proteins, vitamins (A, B12, and folic acid), 51 52 essential minerals such as iron, zinc and selenium (Fellendorf, O'Sullivan, & Kerry, 2017). In 53 addition, the blood also provides an important source of proteins and lysine (Fellendorf et al., 54 2017).

In the central region of Portugal, the black pudding is manufactured with pork fat, pork blood,
bread, onion, coriander, sugar, olive oil and salt. The shelf-life of this product commonly ranges
from 20 to 30 days, although for some specific formulations it can be increased to 90 days
(Silva et al., 2014).

As far as the authors know the information available in the literature regarding the black pudding produced in Portugal is scarce. However, the physicochemical and sensory characterisation of *Morcilla de Burgos* a traditional Spanish blood sausage were studied by Santos et al (Santos, González-Fernández, Jaime, & Rovira, 2003). Ramos et al. (Ramos et al., 2013) provided an important study concerning the composition and quality of different blood

sausages from different countries with diverse raw materials and composition, while alsoplotting the importance of the mineral content in this kind of food product.

66 All blood sausages include antioxidants in its additive list, which allow to minimize lipid oxidation levels. However, both producers and consumers are looking for products where 67 the synthetic antioxidants are replaced by natural ones derived from plants. In this sense, one 68 must highlight bee pollen as a functional food product, since this is rich in proteins, lipids, free 69 70 sugars, carbohydrates, minerals, phenolic acids, flavonoids, sterols, terpenoids, carotenoids and vitamins (Bogdanov, 2011). In fact bee pollen has gained widespread attention due it's 71 72 purported antioxidant (Estevinho, Dias, & Anjos, 2019), anti-inflammatory (Maruyama, Sakamoto, Araki, & Hara, 2010), antimutagenic (Tohamy, Abdella, Ahmed, & Ahmed, 2014) 73 74 and antimicrobial (Morais, Moreira, Feás, & Estevinho, 2011) properties. Indeed, new applications for bee pollen are currently being developed (Almeida et al., 2017; Krystyjan, 75 Gumul, Ziobro, & Korus, 2015) mainly due to its use as a free radical scavenger and as lipid 76 77 peroxidation inhibitor.

The aim of this study was to evaluate the shelf life of black pudding using pollen as natural antioxidant. Therefore, different formulations of black pudding with bee pollen, bee pollen extract and synthetic antioxidant were prepared to determine oxidative stability and sensory acceptability of the final product.

82

83 1. Material and Methods

84 1.1. Chemicals

Folin Ciocalteu phenol reagents, 2,2-diphenyl-1-picryl-hydrazyl (DPPH), gallic acid, 1,1,3,3
tetramethoxypropane (TMP) and trichloroacetic acid were obtained from Sigma Aldrich
(Sternheim, Germany). Aluminum chloride, sodium carbonate, sodium erythorbate (SE),
potassium acetate, ethanol, ethylenediaminetetraacetic acid (EDTA), thiobarbituric acid (TBA)
and chloroform were purchased from Sigma-Aldrich (Germany) and their purities were all over
99%. Absolute alcohol was obtained from Sigma-Aldrich (Germany). All reagents used were of
analytical grade.

92 **1.2. Bee pollen samples**

- 93 The bee pollen samples were collected directly from local beekeepers in the spring of 2017 in
- 94 Castelo Branco, Portugal and stored frozen at -15 $^{\circ}$ C until further analysis.

95 The percentage of pollen grains belonging to each botanical family was determined based on the 96 observation of 500 pollen grains in slides prepared according the acetolise method. The 97 observation of pollens was carried out with a Leitz microscope (Leica, DML, Wetzlar, 98 Germany) at x400 and an image analysis system Qwin 500 (Leica, England).

99 **1.2.1. Preparation of bee pollen extract**

For the bee pollen extraction 11 g of fresh bee pollen was stirred in a digital shaker (VWR 101 15000-1 Advanced Orbital Digital Shaker) with 200 mL of 80% ethanol-water (v/v) at room 102 temperature and at 4 x g during 24 hours in the dark. After this, samples were centrifuged at 103 4080 x g, during 10 minutes and the supernatant was reserved. The extracts were evaporated at 104 40 °C and then were frozen and lyophilized. After that the samples were stored at -20 °C until 105 further analysis.

106 **1.2.2. Total phenolic and flavonoid compounds**

107 The total phenolic content (TPC) of the bee pollen extracts was determined using the Folin–
108 Ciocalteu method as described by Moreira et al. (Moreira, Dias, Pereira, & Estevinho, 2008)
109 and expressed as mg of gallic acid equivalents per g of bee pollen (GAE/g pollen).

110 For total flavonoids contents (TFC) determination in bee pollen the aluminium chloride method

111 was used. Total flavonoids content was expressed as mg of quercetin equivalents per g of bee

112 pollen (QE/g pollen) (Serra Bonvehí, Soliva Torrentó, & Centelles Lorente, 2001).

113 **1.2.3.** Identification of the phenolic compounds in bee pollen

114 The major phenolic compounds of the bee pollen extracts were identified by UHPLC-DAD-

115 ESI-MSn analysis, using a Ultimate 3000 (Dionex Co., San Jose, CA, USA) apparatus with an

- 116 ultimate 3000 Diode Array Detector (Dionex Co., San Jose, CA, USA) coupled to a Thermo
- 117 LTQ XL (Thermo Scientific, San Jose, CA, USA) ion trap mass spectrometer equipped with an
- 118 ESI source. The chromatographic column was an Hypersil Gold (Thermo Scientific, San Jose,
- 119 CA, USA) C18 column (100 mm length; 2.1 mm i.d.; 1.9 µm particle diameter, end-capped) and

120 the general chromatographic conditions corresponded to those previous described (Wasli, Jelali,

121 Silva, Ksouri, & Cardoso, 2018).

122 **1.2.4.** Antioxidant activity of the extracts

123 In order to determine the antioxidant activity of the extract of bee pollen were tested two

124 different methods, namely DPPH and reducing power assays.

125 Free-radical-scavenging (DPPH) assay

126 The capacity to scavenge the 2,2-diphenyl-1-picrylhydrazyl (DPPH) free radical was monitored

according to a method previously described by Morais et al., 2011).

128 The extract concentrations providing 50% scavenging (EC_{50}) were calculated from the graph of

scavenging effect percentage against extract concentration and the results were expressed as

130 mg/mL.

131 Reducing power assay

Reducing power in the extracts were determined by the procedure described by Berker et al (Berker, Güçlü, Tor, & Apak, 2007). In this procedure, the extract concentration that providing 0.5 of absorbance ($C_{0.5}$) was calculated from the graph of absorbance registered at 700 nm against the correspondent extract concentration and the results were expressed as mg GAE/mL.

136 **1.3. Black pudding samples**

The black pudding preparation was conducted under formulation and traditional procedures used in the local factory named "Salsicharia Rebolosa". Regarding producer confidentiality issues the quantity of each ingredient is not publicised in this work, as well the quantity of pollen added.

The basic mixture of ingredients, without the commercial antioxidant, was divided into three lots. Several black puddings were performed containing three different antioxidant sources, namely: 1- fresh bee pollen; 2- lyophilized ethanolic extract of bee pollen; 3- sodium ascorbate (E301, a commercial antioxidant commonly used in the industrial process that in this study was used as control). The bee pollen (fresh and lyophilized ethanolic extract) was added dissolved in the volume of olive oil necessary for the black pudding preparation.

The black pudding for three treatments (different antioxidant sources) at five times (0, 10, 21, 30 and 37 days) and in three replicates was made, totalizing 45 samples. All samples were divided in sealed polyethylene bags under vacuum and stored at 4 °C in a refrigerator. Because the legal shelf life of the black pudding is 30 days, and because is impossible to have the results of microbiological analysis at the same data of the sensory analysis, the evaluators do not made the sensory evaluation for the 37th days to avoid possible health risk.

153 **1.3.1. Microbiological analysis**

For the microbiological analysis of L. monocytogenes, 25 g of sample was homogenised for 2 154 155 min in 225 mL of Half Fraser Base CM0895 (Oxoid, Hampshire, UK), using a Stomacher 400 homogenizer (Seward, Basingstoke, England). The enumeration was performed according to the 156 157 ISO 11290-2:1998/Amd. 1:2004(E) procedure (ISO, 1998). After incubation of the initial suspension for 1 h at 20 °C, a 0.1 mL volume was surface-inoculated on Oxoid Chromogenic 158 159 Listeria Agar Base CM1084 (OCLA, Oxoid) and incubated at 37 °C for 48 h. The detection of L. monocytogenes was according to the ISO 11290-1:1996/Amd. 1:2004(E) procedure (ISO, 160 161 1996). The initial suspension was supplemented with SR0166G selective supplement (Oxoid), 162 incubated at 30 °C for 24 h. To the primary-enriched sample, 0.1 mL was streaked on OCLA and incubated at 37 °C for 48 h, for the secondary-enriched sample, 0.1 mL of the same initial 163 164 supplemented suspension was transferred into 10 mL Fraser Broth supplemented with SR0156E 165 (Oxoid), incubated at 37 °C for 48 h. If no growth was detected in primary-enriched sample, 0.1 166 mL of the secondary-enriched sample was streaked on OCLA and incubated at 37 °C for 48 h. 167 The colonies L. monocytogenes that grew on OCLA was green-blue surrounded by an opaque halo. The determinations per sample were carried out in duplicate and the results were 168 169 expressed in CFU/g.

170 **1.3.2.** Physicochemical analysis

171 The samples were analysed for physicochemical composition (moisture, pH and water activity 172 (a_w)) using standard procedures, along the storage time. Moisture content of samples, along the 173 storage time, was quantified directly, according to the loss of mass after drying at 105 °C in an

174 oven (Thermo Scientific, Heratherm IMH 180) until constant weight, using AOAC procedures

175 (AOAC, 1995). The results were expressed in percentage.

- 176 The pH of samples was determined weighing 10 g of black pudding and mixed with 100 mL of
- 177 ultrapure water until a homogeny solution. The measurements were performed at room
- temperature (around 24 °C).
- 179 Water activity was determined by means of a Rotronic (HygroskopDT, Swiss) coupled with a
- 180 Julabo (F35) thermostatized Baths.

181 **1.3.3.** Oxidative stability - thiobarbituric acid reactive substances content (TBARS)

In order to determine the oxidative stability of the black pudding the method of thiobarbituric acid reactive substances (TBARS) was performed according Almeida et al. (Almeida et al., 2017). Measurements were made on the day of their production and over the storage time (1, 10, 21, 30 and 37 days). Concentrations of 0.6 and 3.0 mmol/L of 1,1,3,3 tetramethoxypropane (TMP) were used as the standards. The results were expressed as mg of MDA/kg of sample (MDA: malondialdehyde). All measurement was carried out in triplicate.

188 **1.3.4.** Sensory analysis

189 The sensory acceptance test was performed using 32 untrained assessor's usual consumers of 190 black pudding (14 women and 8 men with ages ranging between 23 and 54 years) performed the 191 sensory evaluation.

192 The sensory analyses were performed at a room temperature and the samples were presented to 193 the panel cut as 1 cm thick slices of roasted black pudding, under white natural lighting 194 (according to the International Standards (ISO, 1988). Water and apple was provided for mouth 195 rinsing between samples.

196 It was made a ranking descriptive analysis (RDA) (Richter, de Almeida, Prudencio, & de 197 Toledo Benassi, 2010), in which the samples were presented at the same time to the panelists 198 who had to rank the samples for the attribute aroma quality and the flavor, according to a proof 199 sheet prepared for this specific purpose.

200 **1.4. Statistical analysis**

All tests were performed in triplicate and the results were presented as mean \pm standard deviation. A factorial variance analysis was performed to assess the effects of the different antioxidant used as well the shelf life period.

For each significant factor or interaction, the variance percentage was calculated and a Schefee post-hoc test with 95% confidence was applied to the corresponding variables. For the statistical analysis of the sensory data resulting from the ranking test, the Friedman's test was performed based on the sum of the ordinations assigned by the tasters. All the calculations were performed using Statistica from Statsoft (vs 7.09) (Tulsa, OK, USA).

209

210 2. Results and discussion

211 **2.1. Bee pollen characterization**

It is well know that the chemical composition of bee pollen varies depending on the plant
sources, growth conditions and storage conditions (Anjos, Paula, Delgado, & Estevinho, 2019;
Atrouse, Oran, & Al-Abbadi, 2004; Bogdanov, 2011; Elamine et al., 2019; Letícia M.
Estevinho, Dias, & Anjos, 2018; Leticia M. Estevinho, Rodrigues, Pereira, & Feás, 2012;
Komosinska-Vassev, Olczyk, Kaźmierczak, Mencner, & Olczyk, 2015; Serra Bonvehí et al.,
2001).

Palynological analysis found as predominant pollen *Cistus ladanifer* (42.6 %) followed by *Echium* spp. (13.6%) and *Apiaceae* (13.2%). 8.6 % of pollen of *Cistaceae* family were also
founded. *Cistus ladanifer* pollen is very usual in Mediterranean regions, and in particularly in
the region of the study which was well characterized previously by Raimundo et al. (Raimundo
et al., 2018).

The others pollen founded in the mixtures were: *Brassicaceae* spp. (10.1%); *Cichorieae* spp.
(8.0%); *Asteraceae* spp. (1.9%); *Lavandula* spp. (1%); *Plantago* spp. (0.5%); *Silene* spp (0.5%).
The values of TPC and TFC of bee pollen were 35.05±0.5 mg GAE/g of pollen and 6.81 ± 0.08
mg QE/g of pollen, respectively (Table 2). Our results showed a TPC higher than observed by
Morais et al (Morais et al., 2011) that studied the honeybee-collected pollen from five
Portuguese Natural Parks. They are also superior to those described by Campos et al. (Campos,

Webby, Markham, Mitchell, & da Cunha, 2003) who studied pollens from New Zealand and Portugal. Furthermore, the present values are comparable of TPC and TFC of bee pollen collected in Portugal with similar amount of *Cistus ladanifer* pollen (Anjos et al., 2019). The TPC of this pollen mixture were higher that the results founded for the pollen mixtures used by Almeida et al. (Almeida et al., 2017) that studied the use of lyophilized bee pollen extract as a natural antioxidant source in refrigerated sausages.

Because different antioxidant agents present different mechanism for their antioxidant capacities in this work was evaluated the antioxidant activity by two methods (DPPH and reducing power assay). On the other hand, and as say before, the antioxidant activity of bee pollen is well knowing as well the properties of *Cystus ladanifer* pollen. In this work the evaluation of this property is only to calculate the quantity of bee pollen that must be added in the black pudding formulation, in order to allow a similar antioxidant power of that of the commercial one.

The results of antioxidant activity of bee pollen assessed by free-radical-scavenging (DPPH) assay, expressed in terms of EC_{50} value, and reducing power assay are summarized in Table 2. The EC_{50} values of bee pollen sample is 2.62 ± 0.09 mg/mL. This value indicates a good antioxidant activity and higher that the values reported by some authors (Negri, Barreto, Sper,

245 Carvalho, & Campos, 2018; Suriyatem R., Auras R. A., Intipunya P., 2017)

Concerning the values obtained for reducing power assay they are also higher than thosereported for Rape Bee Pollen (Sun, Guo, Zhang, & Zhuang, 2017).

The chromatographic profile at 280 nm of bee pollen extract is represented in Figure 1, while Table 1 summarizes the retention time, UV–vis and MSn spectral data of the identified compounds. Globally, the bee pollen extract was mainly rich in myricetin and quercetin *O*derivatives (Table 1). Please note that the presence of flavonoids such as quercetin derivatives in the bee pollen has been previously related to the biological quality of the pollen, including its high antioxidant function (Lv, Wang, He, Wang, & Suo, 2015; Serra Bonvehí et al., 2001), which is one of the main claimed advantages to the use of bee pollen as an healthy product.

255 **2.2. Black pudding characterization**

In all samples, *Salmonella* spp., *Escherichia* coli and *Listeria monocytogenes* ATCC 19117.
were analysed, according the Portuguese legislation and were absent in all of them.

258 The results obtained for the pH, moisture content, water activity and lipid oxidation by TBARS 259 analysis during the storage period of black pudding are presented in Table 3. Overall, the pH of 260 black pudding samples were similar to those studied by Santos et al. (Santos et al., 2003) and higher to those founded by Diez et al. (Diez, Santos, Jaime, & Rovira, 2008) that studied blood 261 262 sausages produced with rice. The pH of the different formulations of black puddings was influenced by the antioxidant added and the storage period (Table 3). The higher values are 263 264 founded for the black puddings produced with bee pollen as antioxidant, and the variations during the time is different for the different formulation ($TxD = 44.0^{***}$, Table 4). These 265 266 variations could be explained by the fact that the bee pollen have a lower pH than black 267 pudding. According to Anjos et al (Anjos et al., 2019) the pH of pollen ranging between 3.4 and 268 5.9. The pH of the pollen used in the present study is 4.70 ± 0.47 .

269 Moisture content of black pudding depends on the fat content and the final preservation process: 270 cooked, drying or smoking (Ramos et al., 2013). The moisture content of studied products 271 ranging, on average, between 46.33% and 49.73%. The different treatments and storage days 272 were significant factors explaining 48% and 7% of the total variance, respectively, regardless 273 variability between samples also had a high impact (explaining 11.6% of the total variance). The 274 lower moisture content is observed for the black pudding made with pollen and the higher 275 values for the black pudding made with pollen extract. Our values are lower than observed by Fellendorf et al. (Fellendorf et al., 2017) that studied black puddings usually consumed in 276 Ireland and the United Kingdom. Differences can however be due to the distinct list of 277 278 ingredients among the formulations.

a_w is a feature of great importance in food products preservation and particularly in black pudding that was produced with meat and blood. During the manufacturing process of these products they were subjected to high temperatures and, as expected, vegetative cells do not survive, but after the high temperatures process, post-contamination of the product may occur. The higher values of a_w for these kind of products are always higher (Santos et al., 2003) and

because of that it is very important to performed a restrict quality control. Our values for a_w ranging between 0.90 and 0.92 was lower than observed in other studies (Santos et al., 2003). For this parameter, no significant differences were found among the sausages formulations neither along the storage period (Table 4).

TBARS is generally used as an indicator of the degree of lipid oxidation for pork meat and pork
meat sausages, that reflects the content of MDA formed during the oxidation of polyunsaturated
fatty acids (Tang, Sheehan, Buckley, Morrissey, & Kerry, 2001).

291 Concerning the TBARS, Selani et al. (Selani et al., 2011) refer that values lower than 3 mg of 292 MDA/kg sample can be considered in good condition. All the black pudding samples analysed 293 could be considered in good condition during all storage periods (values lower 2.56 mg 294 MDA/kg sample) (Table 3). In the first day, the sample prepared with pollen extract had a value 295 of 3.04 mg MDA/kg of black pudding. The black pudding prepared with pollen as natural 296 antioxidant had values similar to those observed for the black pudding prepared with the 297 commercial antioxidants, except for the first 15 days after production. Further studies may be 298 performed in order to evaluate the optimum quantity of pollen.

For TBARS all factors are highly significant, but the stored days are the most important and explain 44.7 % of the total variance. The variation between days is also different for the different antioxidants used. The bee pollen presents a similar antioxidant effect than the commercial product.

303 **2.3. Sensory evaluation**

The sensory evaluation of the 12 black pudding products is plot in Figure 2.

The black pudding was sensory analysed only until 30 days, because is the legal self-life in the factory. According Silva et al (Silva et al., 2014) after 30 days of storage, the over-wrap packed blood sausages present mould and yeast. Nevertheless, for the vacuum-packed blood sausages the mould and yeast appears only after 45 days (Silva et al., 2014). In our study we use the vacuum-packed system but because no studies were performed in this product to extend the self-life we consider only the legal limit (30 days) stablished for this product for sensory analysis, excluding for this propose the samples with 37 day of self-life. In fact, the

312 microbiological results confirm that no mould or yeast have been developed in the samples, so it 313 was need future research in other to establish better the shelf-life, is not an aim of this work. 314 The ANOVA made for all samples and for the appearance and flavor revel that no significant 315 difference exists for the storage period (appearance: p=1.000; flavor: p=0.999) and for the 316 different formulation period (appearance: p=0.328; flavor: p=0.235). These results confirm that the new additives, pollen or pollen extract do not affect the preference of the consumers. 317 318 Many of the tasters referred that the forced choice required by the triangular test was very difficult because the tree samples are very similar (24 % of the tasters). Other comments given 319 320 by the tasters that help they to identify some differences are: homogeneity of the product (18%); visible pieces of onion (5%) and more fat quantity in a specific sample (3%). However, this 321 322 kinds of comments are all related to manufacture process of this product. The different raw 323 materials are cut in small pouches and mixed but not crushed.

324

325 **3.** Conclusion

The inclusion of bee pollen as an antioxidant could be a natural alternative to prevent the lipid oxidation in black pudding. These products could be added dissolved in the olive oil that will be used in the preparation of the sausage and have the advantage to be a recognized healthy food product.

Additionally, the use of bee pollen as antioxidant improve product quality and consumeracceptance and do not affect their traditional flavor.

Furthermore, it is important to note that the use of bee pollen must be very well mentioned in the label, to prevent allergic risks. More studies will be need in order to identify the more appropriate concentration of bee pollen to use as well the influence of botanical origin of bee pollen.

336

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491 492	Table 2. TPC, TFC and antioxidant activity	ty of bee pollen extract.
	Parameters	Bee pollen
	TPC (mg GAE/g of pollen)	35.05±0.5
	TFC (mg QE/g pollen)	6.99±0.33

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EC₅₀ (mg/mL)

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495 **Table 3.** Chemical analysis of different formulation of black pudding.

Reducing power assay (mg GAE/mL)

			Treatment	
	Days	E301	Pollen	Pollen extract
	1	6.75 ± 0.02^{bB}	6.70 ± 0.01^{aA}	6.68 ± 0.01^{aA}
	10	6.72 ± 0.01^{bB}	6.75±0.01 ^{bC}	6.72 ± 0.00^{aA}
pН	21	6.60±0.02 ^{cA}	6.80±0.01 ^{cB}	6.59 ± 0.04^{bA}
	30	6.55 ± 0.02^{aA}	6.77 ± 0.02^{bC}	6.61 ± 0.02^{bB}
	37	6.51±0.01 ^{aA}	6.71 ± 0.02^{aC}	6.66±0.01 ^{aB}
	1	48.53±0.22 ^{aA}	48.87 ± 0.14^{bA}	48.63±0.19 ^{abA}
Maintenna ann tant	10	49.07 ± 0.28^{aB}	46.54±0.43 ^{aA}	48.66±0.28 ^{abB}
Moisture content	21	47.93±0.73 ^{aB}	46.33±0.59 ^{aA}	49.73±0.08 ^{bC}
(%)	30	48.39±0.48 ^{aB}	46.47 ± 0.44^{aA}	48.10±0.89 ^{aAB}
	37	48.50±0.34 ^{aA}	47.16±0.30 ^{aB}	48.56±0.31 ^{abA}
	1	0.92 ± 0.01^{dA}	0.92 ± 0.01^{bA}	0.92 ± 0.03^{bA}
	10	$0.90{\pm}0.01^{bA}$	0.92 ± 0.01^{bC}	0.91 ± 0.01^{aB}
Water activity	21	0.91±0.01 ^{cA}	0.91 ± 0.01^{aA}	0.91 ± 0.01^{aA}
	30	0.89 ± 0.01^{aA}	0.91 ± 0.02^{aA}	0.92 ± 0.05^{bA}
	37	0.92 ± 0.02^{cA}	0.92 ± 0.01^{bA}	0.92 ± 0.01^{bA}
V	1	1.30±0.08 ^{aA}	2.56±0.30 ^{aB}	3.02±0.13 ^{aC}
TBARS	10	1.28 ± 0.02^{aA}	2.21 ± 0.23^{aB}	2.46 ± 0.24^{bB}
(mg of MDA/kg of	21	1.24 ± 0.15^{aA}	1.33 ± 0.16^{bA}	1.32 ± 0.06^{cA}
black pudding)	30	1.15 ± 0.07^{aA}	1.34 ± 0.10^{bA}	1.27 ± 0.07^{cA}
-	37	1.13 ± 0.06^{aA}	1.30 ± 0.15^{bA}	1.50 ± 0.06^{cB}

 2.62 ± 0.09

6.51±0.30

E301- sodium ascorbate. Different lower-case letter in the same column indicate significant difference (P < 0.05) by Scheffe test. Different capital letters in the same row indicate significant difference (P < 0.05) by Tukey's test.

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505	Table 4 . Component variance analysis for the measured parameter of black pudding considered
506	the threes treatment and the 5 storage period
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the threes treatment and the 5 storage period

	Variance origin	DF	F	р	Variance percentage
	Treatment (T)	2	234.1	0.0000***	37.8
II	Days (D)	4	59.4	0.0000***	15.8
pH	TxD	8	55.2	0.0000***	44.0
	Residual	30			2.4
	Treatment (T)	2	63.2	0.0000***	48.0
Moisture	Days (D)	4	6.4	0.0007***	7.0
Moisture	TxD	8	9.7	0.0000***	33.4
	Residual	30	7		11.6
	Treatment (T)	2	276.3	$0.157^{\text{n.s.}}$	
Watan activity	Days (D)	4	196.7	$0.064^{\text{n.s.}}$	
Water activity	TxD	8	116.8	$0.281^{\text{n.s.}}$	
	Residual	30			
	Treatment (T)	2	149.8	0.0000***	25.7
	Days (D)	4	156.1	0.0000***	44.7
TBARS	TxD	8	32.3	0.0000***	27.0
	Residual	30			2.6

significant, 0.001 ; *** highly significant, <math>p < 0.001

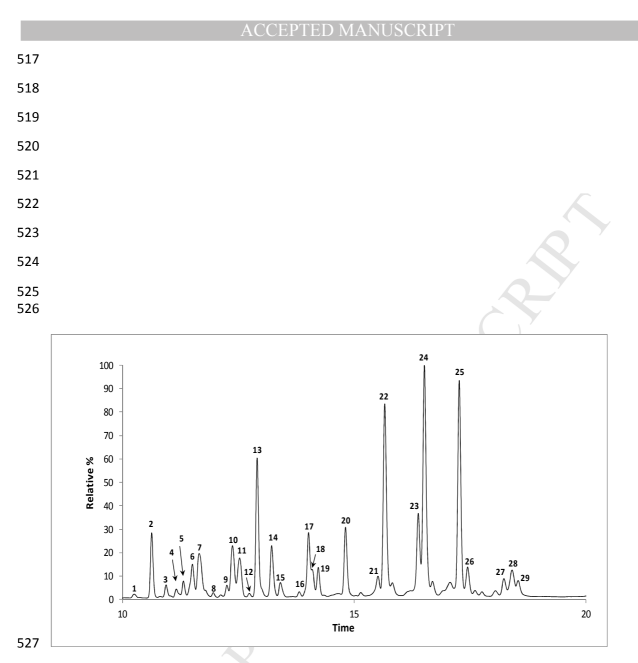
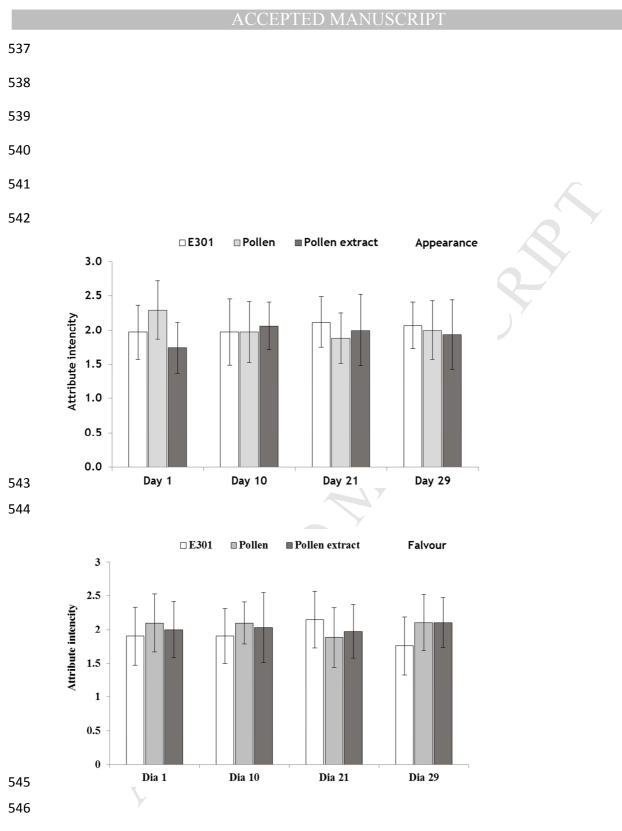


Figure 1. Chromatographic profile at 280 nm of bee pollen extract. Numbers in the figure correspond
to the eluted UHPLC peaks for which UV and MS data is summarized in Table 1.



547 Figure 2. Appearance and flavor evaluation by sensory analysis of black pudding self-life

PN	t _{R (min)}	Amax (nm)	(<i>m/z</i>)	MS^n ions (m/z)	Probable compound
1	10	312	337	MS ² [337]: 173, 162, 191	coumaroyl quinic acid
2	10.3	258, 356	625	MS ² [625]: 316, 317, 271, 461, 479, 609	myricetin-O-rutinoside
3	10.7	264, 351	609	MS ² [609]: 447, 285	luteolin-O-dihexoside
			625	MS ² [625]: 301, 463, 445	quercetin-O-dihexoside
4	10.9	262, 353	479	MS ² [479]: 316, 317	myricetin-O-hexoside
5	11	261, 357	711	MS ² [711]: 667, 316, 317	myricetin-O-(malonyl)rutinoside
6	11.2	270, 355	639	MS ² [639]: 459, 315	isorhamnetin-O-dihexoside
			595	MS ² [595]: 301, 463	quercetin-O-hexosyl-pentoside
7	11.4	256, 308, 354	609	MS ² [609]: 301, 463	quercetin-O-rutinoside
			565	MS ² [565]: 521, 316, 317	myricetin-O-(malonyl)hexoside
8	11.7	266, 353	609	MS ² [609]: 301	quercetin-O-rutinoside
9	12	266, 351	755	MS ² [755]: 609, 593, 573, 285, 255	luteolin-di-O-hexosyl-rhamosíde
10	12.1	257, 353	695	MS ² [695]: 661, 609, 301	Quercetin-O-(malonyl)rutinoside
11	12.2	255, 354	623	MS ² [623]: 315, 459	Isorhamnetin-O-rutinoside
12	12.5	250sh, 297, 308	437	MS ² [437]: 317	hydroxybenzoyl myricetin
13	12.6	256, 354	549	MS ² [549]: 505, 301, 463	quercetin-O-(malonyl)hexoside
14	13.0	265, 350	679	MS ² [679]: 635, 301, 575, 255	quercetin derivative
15	13.2	257, 351	447	MS ² [447]: 301	quercetin-3-O-rhamnoside
16	13.6	271, 351	563	MS ² [563]: 315, 519, 545	isorhamnetin-O-(malonyl)hexoside
17	13.8	256, 354	533	MS ² [533]: 489, 285	luteolin-O-(malonyl)hexoside
18	13.8	mix	317	MS ² [317]: 179, 151	myricetin
19	14.0	255, 353	563	MS ² [563]: 519, 315, 359	isorhamnetin-O-(malonyl)hexoside
20	14.5	245, 296sh, 319	631	MS ² [631]: 495, 317	myricetin-O-dihydroferuloyl protocatechuic acid
21	15.2	245, 296, 310	615	$MS^{2}[615]: 479; MS^{3}[479]: 359; MS^{4}[359]: 317$	myricetin-O-acetyl hydroxybenzoyl protocatechuic acid

Table 1. UHPLC-DAD-ESI-MSⁿ data for bee pollen.

22	15.4	245, 296, 310	615	MS ² [615]: 479; MS ³ [479]: 359; MS ⁴ [359]: 317	myricetin-O-acetyl hydroxybenzoyl protocatechuic acid
23	16.1	240, 295, 308	599	$MS^{2}[599]: 463; MS^{3}[463]: 343; MS^{4}[343]: 301$	quercetin-O-acetyl hydroxybenzoyl protocatechuic acid
24	16.3	240, 295, 309	599	MS ² [599]: 479; MS ³ [479]: 359; MS ⁴ [359]: 317	myricetin-O-acetyl hydroxybenzoyl hydrobenzoic acid
25	17.0	240, 295, 312	583	MS ² [583]: 463; MS ³ [463]: 343; MS ⁴ [343]: 301	quercetin-O-acetyl hydroxybenzoyl hydrobenzoic acid
26	17.2	240, 295, 308	583	MS ² [583]: 463; MS ³ [463]: 343; MS ⁴ [343]: 301	quercetin-O-acetyl hydroxybenzoyl hydrobenzoic acid
27-	17.8-18.4	242, 270-294	785	MS ² [785]: 665; MS ³ [665]: 545; MS ⁴ [545]: 503,	<i>O</i> -dihydroxybenzoyl acetyl malonyl coumaric acid
29				459, 399	flavonoid derivative
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Highlights

- Bee pollen is a healthy product.
- Chemical and sensory characterization of Black pudding with pollen.
- The work suggest that bee pollen could be a natural alternative to prevent the lipid oxidation in black pudding.

Conflict of Interest and Authorship Conformation Form

Please check the following as appropriate:

All authors have participated in (a) conception and design, or analysis and interpretation of the data; (b) drafting the article or revising it critically for important intellectual content; and (c) approval of the final version.

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