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1 **Prooxidant modifications in the cryptome of beef jerky, the deleterious post-**
2 **digestion composition of processed meat snacks**

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29 **Key Words:** proteomics, jerky, beef, post-translational modification, advanced-
30 glycation end products, lipoxidation, DNA-binding.

32 **Abstract**

33 Snacking has traditionally been associated with consumption of foods rich in fats and
34 carbohydrates. However, new dietary trends switched to consumption of protein-rich
35 foods. This study investigates the impact of food processing on the cryptome of one
36 of the most widely consumed meat snacks, beef jerky. We have performed discovery-
37 driven proteome-wide analyses, which identified a significantly elevated presence of
38 reactive prooxidant post-translational modifications in jerky. We also found that these
39 protein decorations impact an important subset of in-silico predicted DNA binding
40 cryptides. Furthermore, we observed cell-dependent reduction in cell viability after
41 prolonged treatments with endogenous-like jerky digests. Collectively these findings
42 uncover the presence of prooxidant modifications in processed dried beef snacks and
43 associate their presence with cytotoxicity. Thus, the findings reported here can pave
44 the way for future studies aimed to establish appropriate dietary recommendations on
45 snacking trends.

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52 **1. Introduction**

53 Snacking is a dietary habit influenced by complex biopsychosocial factors estimated
54 to contribute to almost one third of dietary intake (Njike, Smith, Shuval, Shuval,
55 Edshteyn, Kalantari, et al., 2016). Although snacks have traditionally been associated
56 with consumption of foods rich in fats and carbohydrates with low nutritional profiles,
57 inclusion of whole foods and foods rich in proteins has changed the social perception
58 and outcomes of this habit (Farajian, Katsagani, & Zampelas, 2010).

59 Snacks rich in proteins tend to increase satiety, balance energy consumption and
60 reduce obesity among other significant health benefits (Chapelot, 2011; Marmonier,
61 Chapelot, & Louis-Sylvestre, 2000). However, potential effects of long-term
62 consumption of processed protein-rich snacks on human health remain poorly defined.
63 Proteins are an essential macronutrient required to sustain endogenous protein
64 synthesis and muscle mass (Phillips & Van Loon, 2011). Notwithstanding, the
65 influence of proteins and of their derivatives extend well beyond basic nutritional
66 values (Galvez & de Lumen, 1999; Koldovsky, 1989; Shah, 2000). Food bioactive
67 proteinaceous range from two amino acids to long peptides, and in the human body
68 are known to modulate hypertension, angiogenesis, diabetes, inflammation, immunity
69 and opioid synaptic synergy, to mention just a few (Chatterjee, Gleddie, & Xiao, 2018).
70 Although bioactive proteinaceous are naturally present in foods, a clear distinction
71 between protein molecules naturally present in foods and the cryptome must be
72 performed (Kitts & Weiler, 2003). Indeed, the cryptome consists in a subset of latent
73 functional units that are liberated from protein sequences under certain conditions,
74 such as proteolytic activity, which generates a novel cluster of bioactive peptides
75 commonly dubbed as cryptides (J Autelitano, Rajic, Ian Smith, Berndt, Ilag, & Vadas,
76 2006; Samir & Link, 2011).

77 All food molecules, including proteins, are required to pass through a digestive process
78 in the human body, and thus bioactivity of food proteinaceous becomes determined
79 by the molecular composition of the cryptome, as previously revised (Samir & Link,
80 2011). Endogenous enzymes in the digestive system generate a vast array of multi-
81 length proteolysed peptides that in some instances can become hard to link to the
82 parent protein, and that may possess bioactive abilities that surpass those of the
83 molecules naturally present in foods (Autelitano, Rajic, Smith, Berndt, Ilag, & Vadas,

84 2006). Thus, proteome-wide definition of cryptides and the cryptome of consumed
85 foods has been described in recent studies (Federica Iavarone, Claudia Desiderio,
86 Alberto Vitali, Irene Messana, Claudia Martelli, Massimo Castagnola, et al., 2018).
87 Although it is known that biological variables are able to modify the complex molecular
88 composition of the cryptome, and in turn to create a very singular bioactivity profile of
89 each consumed food in each individual (Samir & Link, 2011); other variables, such as
90 industrial processing and food composition seem to play fundamental roles in the
91 molecular composition and biofunctional outcomes of the cryptome. Industrial
92 processing is largely proven to alter the molecular composition and profile(s) of
93 functional molecules in foods (Capanoglu, Beekwilder, Boyacioglu, De Vos, & Hall,
94 2010; Escobedo-Avellaneda, Moure, Chotyakul, Torres, Welti-Chanes, & Lamela,
95 2011). Similarly, food composition refers to presence of components, such as lipids or
96 sugars, in the food composition, which alters the molecular profile(s) of functional
97 molecules in foods by addition of protein posttranslational modifications (PTMs) (A.
98 Serra, Gallart-Palau, See-Toh, Hemu, Tam, & Sze, 2016). Oxidative degradation of
99 both lipids and sugars generates highly reactive dicarbonyls and other molecules that
100 promote apparition of advanced glycoxidation end products (AGEs), and lipoxidation
101 end products (ALEs) (Vistoli, De Maddis, Cipak, Zarkovic, Carini, & Aldini, 2013).
102 These toxic protein decorations have been identified in food-derived nutraceuticals
103 and bioactive peptides as consequence of food processing (A. Serra, Gallart-Palau,
104 See-Toh, Hemu, Tam, & Sze, 2016). However, whether industrial processing and
105 molecular composition affect the profile(s) and biofunctional outcomes cryptome-wide
106 remains poorly uncovered.

107 As recently unveiled, the raise in consumption of meat snacks has already overpassed
108 that of potato chips in United States (Nielsen, 2017). The main benefits of these
109 snacks, in which beef jerky is the most widely consumed (NHCS, 2018) are mainly
110 promoted on the basis of their high level of protein and their low level of fats. To further
111 investigate the impact that food processing and food composition exert on the
112 cryptome of meat snacks, we have characterized in this study the molecular
113 composition of cryptides from beef jerky and unprocessed beef meat by mass
114 spectrometry (MS)-based cryptomics (Samir & Link, 2011).

115

116 **2. Materials and methods**

117 **2.1. Reagents and chemicals**

118 All reagents used in this study were purchased from Sigma-Aldrich (St. Louis, MO,
119 USA) unless otherwise stated. Water and acetonitrile (ACN) used for mass MS
120 analysis were of high-performance liquid chromatography (HPLC) grade from Fisher
121 Scientific Inc. (U.S.A.). Complete Protease Inhibitor cocktail tablets were obtained
122 from Roche (Basel, Switzerland).

123

124 **2.2. Protein extraction from meat samples for shotgun proteomics study**

125 Lean raw beef cube steak (unprocessed meat) and beef jerky were purchased from a
126 local supermarket in Singapore. Ingredients listed on beef jerky label are: beef, sugar,
127 dextrose, maltodextrin, salt, soy protein, vegetable powders, species, flavour, acidity
128 regulator (331), vegetable oil, antioxidant (316), spice extracts, preservative (250).
129 Both products were of Australian origin. Homogenization of meat samples (~250
130 mg/sample) was performed with 2% sodium dodecyl sulfate (SDS), 100mM
131 ammonium bicarbonate (ABB) and protease inhibitor (1:50 v/w) (SDS extraction
132 buffer) using a ultrasonic processor with a 3 mm microtip probe (Sonics & Materials,
133 U.S.A.) at a 40% amplitude for 5-second pulses for five minutes, followed by 5-second
134 cool-down interval. Homogenization was performed on ice. After homogenization
135 samples were centrifuged at 10,000 x g for 10 minutes at 4°C and the supernatants
136 were collected. Pellets were further homogenized twice as described above and
137 supernatants were combined. Proteins were then precipitated and fat was removed
138 from 1 mL of meat homogenized supernatant by liquid/liquid extraction adding 5 mL
139 of 1:1 methanol/chloroform (v/v), vortexing and allowing to stand for an hour. The
140 mixture was then centrifuged at 5000 x g for 10 minutes at 4°C and liquids decanted.
141 Pellets were dissolved in a 1% sodium deoxycholate (SDC), 100mM ABB solution.
142 The experiment was done per triplicate using three samples obtained from three
143 packets of beef jerky or unprocessed meat.

144

145 **2.3. In-solution tryptic digestion of jerky and unprocessed meat proteins for**
146 **shotgun proteomics study**

147 Digestion of proteins from jerky and unprocessed meat samples was performed as
148 previously described (Gallart-Palau, Serra, Hase, Tan, Chen, Kalaria, et al., 2019;
149 Aida Serra, Gallart-Palau, Wei, & Sze, 2016; A. Serra, Zhu, Gallart-Palau, Park, Ho,

150 Tam, et al., 2016). Briefly, reduction of disulfide bonds was performed with 10 mM
151 dithiothreitol (DTT) at 60°C for 30 minutes, alkylation of cysteines was then performed
152 by adding 20 mM iodoacetamide and incubating at 37°C for 30 minutes. Samples were
153 then diluted 2.5-fold with 100 mM ABB containing 10 mM DTT and incubated for 30
154 min at 37°C. Protein concentration was determined with a bicinchoninic acid assay,
155 and 3 mg of protein from each sample were digested with trypsin at 1:20 enzyme-to-
156 protein ratio overnight at 37°C. Trypsin digestion was quenched and SDC was
157 precipitated by acidification to a final concentration of 0.5% formic acid (FA). SDC was
158 pelleted by centrifugation at 12,000 x g for 10 minutes at 4°C. The pellet was re-
159 suspended in 100mM ABB for further peptide recovery, and a second round of SDC
160 precipitation was performed. The supernatant from both rounds of centrifugation were
161 collected and combined. The peptides were desalted using a 1g Sep-pack C18
162 cartridge (Waters, U.S.A.). The eluted peptides were dried completely in a vacuum
163 concentrator (Eppendorf, Germany).

164

165 **2.4. High-pressure liquid chromatography fractionation of jerky and** 166 **unprocessed meat peptides**

167 Desalted peptides were fractionated by high-pressure liquid chromatography (HPLC)
168 as previously described (Gallart-Palau, Serra, Lee, Guo, & Sze, 2017), with minor
169 modifications. Peptides were reconstituted in 200 µL of mobile phase A (85% ACN,
170 0.1% acetic acid) and fractionated using a PolyWAX LP 3.5 µm 4.6 × 250 mm column
171 (PolyLC, U.S.A.) with a Shimadzu Prominence UFLC system (Dionex, USA) with UV
172 monitoring of peptide intensities at 280 nm. Peptides were separated in a 70-min
173 gradient as follows: 0% B (0.1% FA) for 10 min, 0–20% B for 30 min, 20–65% B for
174 10 min, 65-100% B for 1 min, 100% B for 8 min, 100-0% B for 1 min, and 0% B for 10
175 min. Fractions were collected at 1 min intervals and combined in a total of 18 fractions
176 per sample according to the peak elution profile. Combined fractions were dried
177 completely in a vacuum concentrator.

178

179 **2.5. Liquid chromatography-mass spectrometry shotgun proteomics of jerky** 180 **and unprocessed meat peptides**

181 Dried fractionated peptide samples were reconstituted in mobile phase A (3% ACN,
182 0.1% FA). LC-MS/MS analysis of peptides was performed using a Dionex UltiMate
183 3000 UHPLC system coupled with an Orbitrap Elite mass spectrometer (Thermo

184 Fisher Inc., Bremen, Germany) as previously described with minor modifications
185 (Gallart-Palau, Serra, Qian, Chen, Kalaria, & Sze, 2015; Gallart-Palau, Serra, & Sze,
186 2016; Gallart-Palau, Serra, Wong, Sandin, Lai, Chen, et al., 2015; Aida Serra, Xavier
187 Gallart-Palau, Bamaprasad Dutta, & Siu Kwan Sze, 2018). Spray was generated
188 using a Michrom Thermo CaptiveSpray nanoelectrospray ion source (Bruker-Michrom
189 Inc., Auburn, USA) working at 1.5 kV. Peptide separation was performed using a
190 reverse-phase Acclaim PepMap RSL column (75 μm ID \times 15 cm, 2 μm particle size,
191 Thermo Scientific Inc.) maintained at 35°C and working at 300 nL/min. Eluents A (0.1%
192 FA) and B (90% ACN, 0.1% FA) were used to establish the following 60-min gradient
193 at a flow rate of 300 nL/min over a 60 minute period: 3% B for 1 min, 0–35% B for 47
194 min, 35–50% B for 4 min, 50–80% B in 5 seconds, 80% B for 78 sec, 80–3% B in 6
195 seconds, 3% B for 6.5 min. Orbitrap Elite mass spectrometer was set to positive mode
196 for data acquisition with Xcalibur 2.2 SP1.48 software (Thermo Fisher Scientific Inc.,
197 Bremen, Germany) software alternating between full Fourier transform-mass
198 spectrometry (FT-MS) (350–2000 m/z, resolution 60000, with 1 μscan per spectrum)
199 and FT-MS/MS (150–2000 m/z, resolution 30000, with 1 μscan per spectrum).
200 Fragmentation of the 10 most intense precursors with charge $>+2$ and isolated within
201 a 2 Da window was performed using high-energy collisional dissociation (HCD) mode
202 using 32% normalized collision energy. A threshold of 500 counts was enabled. For
203 Full FT-MS and FT-MS/MS automatic gain control was set to 1×10^6 .

204

205 **2.6. Protein extraction from meat samples for cell viability assay**

206 Proteins from 50 g of unprocessed meat or jerky were extracted with 200 mL of SDS
207 extraction buffer as detailed previously with minor modifications (A. Serra, X. Gallart-
208 Palau, B. Dutta, & S. K. Sze, 2018). Homogenization was performed with the ultrasonic
209 processor with a 6 mm probe (Sonics & Materials). Proteins were precipitated and fats
210 removed with 200 mL of 1:1 methanol:chloroform (v/v) as previously detailed. Pelleted
211 proteins (~1 g/sample) were dissolved in 8 mL of 4 mM hydrochloric acid (pH 2), and
212 homogenized with the ultrasonic processor using the 3 mm probe (40% amplitude, 10-
213 second pulses for 10 minutes, 10-second cool down, on ice). The meat samples were
214 further homogenized in a bullet blender (Next Advance, U.S.A.) with metallic beads
215 (1mm; Next Advance) at maximum speed for 30 minutes at 4°C. Samples were
216 centrifuged at 10,000 $\times g$ for 10 minutes at 4°C, and supernatants were collected.

217

218 **2.7. Preparation of endogenous-like unprocessed meat and jerky digests for**
219 **cell viability assay**

220 To simulate the sequential endogenous stomach, and duodenum human digestion,
221 pepsin and trypsin proteases were used in a static *in vitro* digestion strategy as
222 previously described (Nguyen, Bhandari, Cichero, & Prakash, 2015), with minor
223 modifications. Reduction and alkylation of unprocessed meat or jerky proteins were
224 performed as previously described for the shotgun proteomics study. Reduced and
225 alkylated proteins were digested with pepsin (1:20 w/w) at pH 2 and 37°C overnight.
226 Pepsin was inactivated by increasing the pH to 8 with ABB. Overnight digestion was
227 then performed with trypsin (1:40 w/w) at 37°C, and inactivated by adjusting to pH 3
228 with acetic acid. Peptides were then desalted using a 1g Sep-pack C18 cartridge
229 (Waters, U.S.A.). The eluted peptides were dried completely in a vacuum concentrator
230 (Eppendorf, Germany).

231

232 **2.8. Cell viability assay**

233 Cell viability was measured with a colourimetric tetrazolium dye, thiazolyl blue
234 tetrazolium bromide (MTT) assay. NIH 3T3 fibroblast and HCT116 cells (American
235 Type Culture Collection (ATCC), U.S.A.) were seeded per well in 96-well plates and
236 incubated overnight in Dulbecco's Modified Eagle's Medium (DMEM, GE Healthcare
237 Life Sciences, U.S.A.) supplemented with 10% fetal bovine serum. The endogenous-
238 like unprocessed meat or jerky protein digests were dissolved in 1 mL of 1x PBS,
239 diluted with DMEM and filtered with a 0.22 µm polyethersulfone filter (Sartorius AG,
240 Germany). The media was then exchanged with fresh media containing 0.25 to 1
241 mg/mL of endogenous-like digests and cell viability was tested via an MTT assay at
242 24, 48 and 72 hours according to manufacturer's protocol. Cells treated with DMEM
243 were considered as control. Absorbance was measured at 570nm and untreated
244 controls were set to 100% viability. The experiments were performed per triplicate.

245

246 **2.9. In-silico identification of cryptides with DNA-binding capacity**

247 For the in-silico characterization of cryptides with DNA-binding capacity the list of
248 unique peptides identified by shotgun proteomics after tryptic digestion of jerky and
249 unprocessed meat were converted to FASTA format (including post-translationally
250 modified and unmodified peptides) using an in-house program. Subsequently, the lists
251 were submitted to the DNA binding protein prediction (DNA-Binding) online platform

252 (Szilagyi & Skolnick, 2006). In-silico DNA binding capacity prediction was done from
253 peptide sequence at 15% false discovery rate (Szilagyi & Skolnick, 2006). Only
254 peptides with a prediction score confidence > 10 were considered as cryptides with
255 DNA binding capacity. Cryptides were in-silico predicted for every replicate (n=3).

256

257 **2.10. Bioinformatics and data analysis**

258 The raw MS/MS data were de-isotoped and converted into Mascot Generic Format
259 using the Thermo Proteome Discoverer software (version 1.4.1.14, Thermo Fisher
260 Scientific, U.S.A.), then searched against the *Bos taurus* reference proteome and
261 UniMod protein modification were identified using an in-house Mascot server using the
262 Mascot error tolerance search mode (version 2.6.1; Matrix Science, U.S.A.). The
263 search was performed with a 5 ppm precursor MS tolerance, 0.02 Da MS/MS fragment
264 tolerance. The missed cleavage sites was determined with Mascot search results.
265 GraphPad Prism 6 (GraphPad Software, Palo Alto, CA) was used for parametric
266 statistical analyses and for creating data plots. Data were analyzed by Student's t-test
267 at p-values < 0.05 and are reported as mean ± SD.

268

269 **2.11. Availability of data**

270 The data generated in this study are publicly available via the ProteomeXchange
271 consortium through the partner repository PRIDE (Vizcaino, Csordas, Del-Toro,
272 Dianes, Griss, Lavidas, et al., 2016).

273 *PRIDE Accession: PXD012112*

274

275 **3. Results and discussion**

276 **3.1. Protein post-translational modifications in jerky**

277 In this study we sought to investigate the presence of protein PTMs induced by food
278 processing and food composition in beef jerky using unprocessed meat as control.
279 Peptidome-wide characterization revealed a significantly higher number of total unique
280 peptides in unprocessed meat samples compared to jerky (**Supplementary Figure 1**
281 **and Supplementary datasets 1 and 2**), a fact that is consistent with previous reports
282 (Yu, Morton, Clerens, & Dyer, 2017), and that could be related with denaturation and
283 formation of proteinaceous aggregates of low solubility in jerky during production
284 processes. Beef jerky production process generally involves a drying process of the
285 marinated meat at about 71.1°C (160 degrees F) prolonged up to 12 h. The duration

286 of the drying process depends on specific recipe. On the contrary, drying temperature
287 of 71.1°C was recommended by the United States Department of Agriculture (USDA)
288 to achieve an immediate lethality of *Salmonella* and ensure an adequate
289 microbiological safety (USDA, 2007). This long drying process is often precluded by a
290 shorter post-drying heating step at about 135°C. In the first heating step meat is
291 cooked and microbial numbers is reduced whilst the second drying phase stabilizes
292 organoleptically the product and prevents microbial growth.

293 Protein PTMs significantly elevated in jerky were essentially derived from the reaction
294 between proteins with endogenous sugars, lipids or other components catalyzed by
295 high temperature treatment (**Figure 1 and Supplementary Table 1**). These subset of
296 PTMs, practically unidentifiable in unprocessed meat, included glycated –derived from
297 hexose, glucuronyl and galactosyl and oligosaccharides- (**Figure 1A**) and lypoilated
298 modified proteins (**Figure 1B**), as well as their degradation products, advanced
299 glycoxidation and lipoxidation end products (AGEs and ALEs, respectively) (**Figure**
300 **1C-D**). AGEs shown as a heterogeneous mixture of PTMs in jerky (**Figure 1E**). The
301 most abundant AGE-related PTM detected in this processed meat snack was the
302 highly reactive AGE precursor α -dicarbonyl 3-deoxyglucosone (3-DG)(Loughlin &
303 Artlett, 2010), which affects Arg residues, followed by the Lys modifications N ϵ -
304 carboxy-methyl-lysine (CML) and N ϵ -carboxy-ethyl-lysine (CEL) (**Figure 1E**). CML
305 and CEL have been subject of exhaustive study due to their capacity to interact with
306 human AGEs receptor (RAGE) (Xue, Rai, Singer, Chabierski, Xie, Reverdatto, et al.).
307 Additionally, Arg modifying AGE-related PTMs such as the methylglyoxal-derived
308 hydroimidazolone (MG-H1) and dihydroxyimidazolidine were also found significantly
309 elevated in jerky (**Figure 1E**).

310 Accumulation of AGEs in the mammal's body leads to apparition of oxidative stress,
311 inflammation, protein cross-linking and remains negatively associated with lifespan
312 (Gallart-Palau, Serra, & Sze, 2015; Gallart-Palau, Tan, Serra, Gao, Ho, Richards, et
313 al., 2019; Sharma, Kaur, Thind, Singh, & Raina, 2015). Although AGEs are
314 endogenously produced through the sugar metabolism in all tissues and fluids of the
315 body (Ahmed, Luthen, Haussinger, Sebekova, Schinzel, Voelker, et al., 2005), it was
316 shown that exogenous AGEs from foods possess the same abilities as endogenous
317 AGEs at the time to cause protein cross-linking and oxidative stress in cells (Cai, Gao,
318 Zhu, Peppia, He, & Vlassara, 2002). Similarly, it has been shown that dietary restriction
319 of AGEs in diabetes subjects and subjects with renal dysfunction has a significant

320 repercussion in lowering the circulatory levels of inflammatory markers and VCAM-1,
321 a specific marker of endothelial dysfunction (Uribarri, Peppas, Cai, Goldberg, Lu, He,
322 et al., 2003; Vlassara, Cai, Crandall, Goldberg, Oberstein, Dardaine, et al., 2002). As
323 demonstrated by Cai and colleagues (Cai, He, Zhu, Chen, Striker, & Vlassara, 2008)
324 and emphasized by Sharma *et al.* dietary reduction of AGEs in aging populations is
325 expected to exert therapeutic effects on the prevalence of several chronic age-related
326 diseases (Sharma, Kaur, Thind, Singh, & Raina, 2015). Presence of AGEs in snacks
327 has been found in chips, crackers and cookies (Sharma, Kaur, Thind, Singh, & Raina,
328 2015; Story, Hayes, & Kalina, 1996), however, their levels were very low compared to
329 meats (Sharma, Kaur, Thind, Singh, & Raina, 2015). Here, we demonstrate that the
330 levels of toxic glycans in the most widely consumed meat snack largely surpass those
331 in unprocessed meats and in other traditionally considered unhealthy snacks such as
332 potato chips.

333

334 **3.2. Food-ubiquitous bioactive motifs contain prooxidant modifications in** 335 **jerky**

336 Toxic glycans in bioactive peptides/motifs can alter their function(s) and molecular
337 interactions (A. Serra, Gallart-Palau, See-Toh, Hemu, Tam, & Sze, 2016). Similarly,
338 when the cryptome is affected by reactive PTMs (J Autelitano, Rajic, Ian Smith, Berndt,
339 Ilag, & Vadas, 2006), release of cryptome-derived peptides could become hindered
340 following endogenous enzymatic digestion (J Autelitano, Rajic, Ian Smith, Berndt, Ilag,
341 & Vadas, 2006). Thus, since reactive PTMs were only evident in jerky, we investigated
342 affection of food-ubiquitous bioactive peptides by reactive PTMs in this processed
343 food product. In depth analysis of post-translationally modified proteomes from jerky
344 led to identification of several bioactive peptides affected by AGE-related modifications
345 in (or surrounding) bioactive motifs as shown in **Table 1**. Of note, we found the beef
346 myosin-derived peptide ALGTNPTNAEVKK₁₃ modified in jerky by CML, CEL and Hex
347 at the residue K₁₃ (**Table 1**). This peptide contains the VKK region found in the
348 antihypertensive bioactive peptide VKKVLGNP originally discovered in pork myosin
349 (Katayama, Jamhari, Mori, Kawahara, Miake, Kodama, et al., 2007). Nevertheless,
350 this potentially bioactive peptide found in beef jerky displayed prooxidant modifications
351 at the bioactive motif VKK fact that, we hypothesize, may reduce or modify its
352 bioactivity. Stoichiometry analysis of the prooxidant modifications in the motif VKK
353 indicated that ~40% of the total peptide was modified by prooxidant modifications in

354 jerky. From these, ~29.3% of the peptide was modified by AGEs and ~9.6% was
355 glycosylated (Hex) (**Figure 2A**). On the contrary, no modified proteoform in this motif was
356 detected in unprocessed meat.

357 Similarly the biofunctional motif RPR, originally identified in pork nebulin with
358 contrasted antihypertensive abilities (Escudero, Toldra, Sentandreu, Nishimura, &
359 Arihara, 2012), was found directly affected by a 3-DG modification and flanked by MG-
360 H1, 3-DG and Hex modifications in the peptide AGFAGDDAPRAVFPSIVGRPR in
361 jerky (**Table 1**). Stoichiometry of these prooxidant modifications in RPR peptide motif
362 indicated that 96.2% of the total peptide content was modified by AGEs and 3.8% was
363 modified by glycation (**Figure 2B**). Furthermore, 10.45% of the peptide containing the
364 biofunctional motif RPR presented an additional 3-DG modification in the Arg residue
365 R₁₀ (**Table 1**). None of these modifications were detected in the RPR motif in
366 unprocessed meat.

367 The bioactive motif FHG, proven to possess antihypertensive, antimicrobial and
368 cancer cytotoxic properties, was identified in jerky flanked by a CML modification in
369 the NDMAAQYKVLGFHG peptide from jerky (**Table 1**). Stoichiometry analysis of the
370 modified peptide indicated that 40.8% was modified by Hex and 17.5% was modified
371 by CML in jerky (**Figure 2C**). None of these modifications were detected in the FHG
372 motif in unprocessed meat.

373 We also observed that the functional peptides AGFAGDDAPRAVFPSIVGRPR and
374 NDMAAQYKVLGFHG, while were containing glycotoxins and prooxidant
375 modifications in jerky, occasionally presented trypsin miscleaved sites in their modified
376 potential digestion sites. On the contrary, no miscleaved sites were detected for these
377 peptides in unprocessed meat.

378

379 **3.3. Prooxidant modifications impair the release of cryptides from jerky**

380 Observation of missed cleavages in bioactive peptides of jerky prompted us to
381 investigate the presence of missed cleavage sites in jerky originated by a hindering
382 effect exerted by PTMs on normal protease activity due to their localization affecting
383 protease digestion sites. Trypsin is the active form of trypsinogen and represents one
384 of the main proteases in the digestive system (Antalis, Shea-Donohue, Vogel, Sears,
385 & Fasano, 2007; Eggermont, Molla, Tytgat, & Rutgeerts, 1971), thus it was used in
386 this study to investigate the presence of missed cleavage sites in jerky peptidomes
387 modified and unmodified by prooxidant modifications. As shown in **Figure 3A**, 74.6%

388 of the total unmodified peptides did not display any missed cleavage sites, whereas
389 the remnant ~25% was distributed in a range that included one to four missed
390 cleavage sites per peptide (**Figure 3A**). On the contrary, the presence of missed
391 cleavage sites in the prooxidant modified subset of peptides was significantly
392 upregulated compared to their unmodified counterparts affecting ~95% of total
393 modified peptides. From these, a big portion of 66.7% displayed one missed cleavage
394 site (**Figure 3A**). Hindering effects of PTMs on normal protease activity have been
395 demonstrated by acetylated lysines and other modifications (Zee & Garcia, 2012).
396 Thus, we found that presence of prooxidant PTMs in processed meat snacks
397 significantly reduces the endogenous protease efficiency of one of the main proteases
398 in the digestive system, which in turn affects the release and composition of the
399 resultant jerky-derived cryptome.

400

401 **3.4. Prooxidant modifications affect cryptides with DNA binding capacity in** 402 **jerky**

403 Biological functions of cryptides are incipiently uncovered and range from signaling
404 capacities to implication in regulatory mechanisms in target cells (Ueki, Someya,
405 Matsuo, Wakamatsu, & Mukai, 2007). These signaling capacities involve DNA
406 interaction (Ueki, Someya, Matsuo, Wakamatsu, & Mukai, 2007) and thus, identify
407 potential peptides with DNA binding capacity in highly consumed foods is required to
408 define and characterize the biofunctional outcomes of cryptides from foods. In this
409 study we have performed bioinformatics to obtain *in silico* predictions of the cryptides
410 with DNA binding capacity from whole peptidomes identified by shotgun proteomics in
411 jerky and unprocessed meat. DNA binding capacity is an important biological function
412 described *in vitro* in cryptome-released peptides (H Barkhudaryan, V Hunanyan,
413 Sarukhanyan, M Stepanyan, H Zakaryan, Grigoryan, et al., 2012; F. Iavarone, C.
414 Desiderio, A. Vitali, I. Messana, C. Martelli, M. Castagnola, et al., 2018) that can be
415 predicted in a peptidome-wide approach using in-silico bioinformatics strategies as
416 previously shown (Szilagyi & Skolnick, 2006).

417 Use of DNABind predict in this study (Szilagyi & Skolnick, 2006) revealed that jerky
418 peptidomes contained a total of 553 ± 47 peptides with DNA binding capacity whereas
419 unprocessed meat peptidomes contained 766 ± 39 peptides. These data were shown
420 in **Figure 3B** and whole lists of potential cryptides with DNA binding capacity from
421 jerky and unprocessed meat are included in **Supplementary dataset 3 and 4**. Further

422 bioinformatics analyses revealed that prooxidant modifications were affecting 11% of
423 the total cryptides with DNA binding capacities in jerky whereas these toxic
424 modifications only affected 0.6% of total DNA binding cryptides in unprocessed meat
425 **(Figure 3B)**.

426 Previous studies reported that prooxidant modifications affecting proteins with DNA
427 binding abilities can cause DNA damage (Guerrero, Vasudevaraju, Hegde, Britton, &
428 Rao, 2013; Padmaraju, Bhaskar, Prasada Rao, Salimath, & Rao, 2011), and it has
429 been hypothesized that this mechanism may play a central role in the pathology of
430 several major diseases in which cell apoptosis is crucially involved (Guerrero,
431 Vasudevaraju, Hegde, Britton, & Rao, 2013). Thus, based on the findings of this study,
432 further studies will be required to evaluate the abilities of the cryptome described with
433 DNA binding capacities in jerky and how these become modified by the effects of the
434 specific prooxidant modifications uncovered.

435

436 **3.5. Modified jerky cryptides impair cell line proliferation in fibroblasts**

437 To explore *in vitro* the findings described in the previous section and to investigate any
438 potential cytotoxic effect(s) of prooxidant modifications in the cryptome of beef jerky,
439 NIH3T3 fibroblasts and HCT116 cell lines were treated with endogenous-like jerky
440 digests. Digests from unprocessed meat in all conditions were used as control. Cells
441 were treated with concentrations of digests ranging from 0.25 to 1 mg/ml in different
442 time points (24h, 48h and 72h), and cell proliferation was evaluated by MTT assay.
443 Significant time and dose-dependent cytotoxic effects of endogenous-like jerky digests
444 were strikingly observed after 72h using 1 mg/ml of jerky digest on fibroblast cells (p-
445 value 0.049) **(Figure 4)**. Rest of time-points, cell line (data not shown) and lower
446 concentrations tested did not yield significant differences between jerky and
447 unprocessed meat **(Supplementary Figure 2)**. These findings highlight the expected
448 cell-dependent ability of prooxidant modified jerky cryptides to exert cytotoxicity. The
449 observed effects are apparently linked to long and abundant exposure of susceptible
450 cells to prooxidant modified jerky cryptides, suggesting that these detrimental effects
451 might be linked to interaction of variables like chronic/high level of consumption and
452 aging, a hypothesis that deserves further investigation.

453 Dose-dependent cytotoxic effects of AGEs, in susceptible cells, were encountered
454 several years ago by Li and colleagues (Li, Mitsuhashi, Wojciechowicz, Shimizu, Li,

455 Stitt, et al., 1996). The authors found that these pathogenic adducts stimulate the
456 presence of AGE-specific receptors (RAGEs) in target cells from tissues like renal,
457 central nervous system and vasculature (Li, et al., 1996) affecting the endothelium, a
458 basic target in the pathogenesis of several major diseases (A. Serra, Gallart-Palau,
459 Park, Lim, Lim, Ho, et al., 2018). Similarly, ALEs have also been proven to interact
460 with RAGEs in e.g. to dysregulate the levels of vasculature calcification (Ott, Jacobs,
461 Haucke, Navarrete Santos, Grune, & Simm, 2014). Prooxidant modifications are
462 eliminated from target cells by the renal system (Ott, Jacobs, Haucke, Navarrete
463 Santos, Grune, & Simm, 2014), and their potential implications as central and/or
464 comorbid factors in diseases that involve chronic failure of the renal system is under
465 intense scientific scrutiny (Rabbani & Thornalley, 2018). Although the uptake of
466 exogenous prooxidant modifications in the digestive system is somewhat limited
467 (Vlassara, et al., 2002), it was clearly shown that a reduction in the amount of
468 consumed prooxidant modifications has a significant impact on the levels of
469 inflammatory processes and protein PTMs that take place in the body (Cecil, Johnson,
470 Rediske, Lotz, Schmidt, & Terkeltaub, 2005; Vlassara, et al., 2002). Therefore, the
471 links between long-time consumption of foods that contain an elevated amount of
472 cryptides modified by prooxidant modifications, aging and human diseases, have to
473 be further explored in the biomedical and nutritional fields by crosstalk between these
474 disciplines. Similarly, we demonstrate here that definition of the spectrum of prooxidant
475 modifications from highly consumed foods cryptome-wide, as well as analysis of their
476 potential detrimental outcomes, becomes preferential to progress on the knowledge
477 and improvement of the ties linking nutrition and health.

478 **4. Conclusions**

479 We confirmed an abnormal presence of AGEs and ALEs in the peptidome of beef jerky
480 snacks, and shown that these reactive and toxic protein decorations directly affect
481 jerky-derived cryptides and modify the global composition of the jerky cryptome by
482 interfering with endogenous protease efficiency. Furthermore, we demonstrate that
483 endogenous-like digests from jerky significantly reduce cell viability in a cell-type and
484 dose-dependent manner under prolonged periods of exposure. Thus, our findings
485 indicate that current food processing procedures affect the molecular composition of
486 largely-consumed meat snacks. Based on these findings, additional research actions
487 will be required to establish the basis of a proper dietary recommendation on snacking

488 habits in line with current snacking trends, in which protein-rich snacks gradually
489 substitute the inclusion of carbohydrate- and fat-rich snacks in the diet.

490

491

492

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498

499 **Competing interests**

500 The authors declare that they have no competing interests.

501

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504 **References**

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662

663

664 **Figure captions:**

665 **Figure 1. Relative quantitation of the significantly elevated PTMs in beef jerky**
666 **compared to beef unprocessed meat. A.** Comparison of the total number of
667 matched glycated peptides (glycation category includes hexosylation [+162Da at KN],
668 hexosamine [+161 at KT], heptosylation [+192 Da at KNQRST], lactosylation [+324
669 Da at KR], glucuronylation [+176 Da at S] and galactosylation [+178 Da at K]); **B.**
670 Comparison of the total number of matched lipoilated peptides [+188 Da at K]; **C.**
671 Comparison of the total number of matched advanced lipoxidation end products
672 (ALEs)-containing peptides. In this category the modification nonanedioic (azelaic)
673 acid cross link in oxidized lipoproteins [+175 Da at K] was considered. **D.** Comparison
674 of the total number of matched advanced glycoxidation end products (AGEs)-
675 containing peptides. This category included carboxymethylation [+58 Da at CKW],
676 carboxyethylation [+78 Da at K], dihydroxyimidazolidine [+72 Da at R],
677 carboxymethylation [+58 Da at CKW] methylglyoxal-derived hydroimidazolone (MG-
678 H1) [54 Da at R] and 3-deoxyglucosone [+144 Da at R]. **E.** Categorization of AGEs in
679 jerky.

680

681 **Figure 2. Stoichiometry of bioactive peptides containing prooxidant PTMs**
682 **significantly elevated in jerky compared to beef unprocessed meat. A.** Beef
683 myosin-derived peptide ALGTNPTNAEVKK# overlaps with the partial sequence VKK
684 of the bioactive peptide VKKVLGNP. The unmodified proteoform of this peptide was
685 detected in unprocessed meat. **B.** The gamma enteric smooth muscle actin peptide
686 AGFAGDDAPRAVFPSIVGRPR which contains the bioactive sequence RPR. This
687 peptide was not present in unprocessed meat. **C.** The beef myoglobin-derived peptide
688 NDMAAQYK#VLGFHG which contains the bioactive sequence FHG. This peptide was
689 not present in unprocessed meat. # denotes the modified site. Bioactive motifs are
690 underlined.

691

692 **Figure 3. Peptidome-wide analysis of jerky and beef unprocessed meat. A.**
693 Presence of missed cleavage sites expressed in percentage in tryptic digested jerky
694 and unprocessed meat. *means significant differences between jerky and
695 unprocessed meat assessed by chi-square test ($p < 0.05$). **B.** Peptidome-wide
696 characterization of prooxidant post-translationally modified peptides with DNA binding
697 capacity. A total of 766 and 553 peptides displayed potential DNA-binding capacity in
698 unprocessed meat and jerky, respectively. Prooxidant modified peptides represented
699 the 11% of the total peptides that displayed DNA-binding capacity in jerky. DNA-
700 binding capacity of tryptic peptides was predicted *in silico*.

701

702 **Figure 4. MTT assay results for NIH 3T3 exposed to 1 mg/mL of endogenous-**
703 **like jerky or unprocessed meat digests after 24 h, 48 h and 72 h treatment.** Data
704 are expressed as mean \pm SEM. Significant differences were assessed by ANOVA
705 with uncorrected Fisher's LSD multiple comparisons test ($p < 0.05$).

Table 1. Peptides identified in the beef jerky that contained sequences corresponding to entire or partial known bioactive peptides. Locations of the PTM have been bolded, while the sequence matching known bioactive peptides have been underlined. ^bACE, refers to angiotensin-I converting enzyme. Reported peptides were identified in at least two replicates.

Accession number	Protein	Identified Peptide	Reported bioactive motif	PTM	Bioactivity	Reference
A0JNJ5	Myosin light chain	ALGTNPTNAE <u>VKK</u>	VKKVLGNP	CML	ACE ^b -inhibitory	
		ALGTNPTNAE <u>VKK</u>		CEL		
		ALGTNPTNAE <u>VKK</u>		Hexose		
F1MKC4	Actin, gamma enteric smooth muscle	AGFAGDDAPRAVFPSIVGR <u>RPR</u>	RPR	MG-H1	Antihypertensive	
		AGFAGDDAPRAVFPSIVGR <u>RPR</u>		3-DG		
		AGFAGDDAPRAVFPSIVGR <u>RPR</u>		Hexose		
		AGFAGDDAPRAVFPSIVGR <u>RPR</u>		3-DG, 3-DG		
P02192	Myoglobin	NDMAAQYKVLGF <u>FHG</u>	FHG	CML Hex	ACE-inhibitory, antimicrobial, cancer cell cytotoxic	