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1 Chemical reactivity of nitrite and ascorbate in a cured and cooked meat model
2 Implication in nitrosation, nitrosylation and oxidation

3

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14

15 Abstract

16 Nitrite, added to cured meat for its bacteriological and technological properties, is implicated
17 in the formation of nitroso compounds (NOCs), such as nitrosylheme, nitrosamines and
18 nitrosothiols, suspected to have a potential impact on human health. The mechanisms
19 involved in NOC formation are studied in regard with the dose-response relationship of
20 added nitrite and its interaction with ascorbate on NOC formation in a cured and cooked meat
21 model. The impact of a second cooking stage on nitrosation was evaluated. The addition of
22 nitrite in the cured and cooked model promoted heme iron nitrosylation and S-nitrosation but
23 not N-nitrosation. Nitrite reduced lipid oxidation without an additional ascorbate effect. The
24 second cooking sharply increased the nitrosamine content while the presence of ascorbate

25 considerably lowered their levels and protected nitrosothiols from degradation. This study
26 gives new insights on the chemical reactivity of NOCs in a cured meat model.

27

28

29 Key words

30 Nitrite, ham, nitrosamine, nitrosylheme, nitrosothiol

31

32

33 1. Introduction

34

35 Nitrite and nitrate are used in cured meat processing to prevent the growth of harmful
36 bacteria and develop the color and flavor of products. The ability of nitrite to limit the growth
37 of bacteria, especially *Clostridium botulinum* strains which produce toxins was formerly
38 demonstrated (Skibsted, 2011). Nitrite also promotes the conservation of cured meat thanks
39 to its antioxidant capacity (Berardo, De Maere, Stavropoulou, Rysman, Leroy, & De Smet,
40 2016).

41 Nevertheless, the use of nitrite in food has been debated for several years. The risk of nitrite
42 comes from its ability to form nitroso compounds (NOCs), some of which are mutagenic.
43 Indeed, nitrite can react with secondary amines (N-nitrosation) to form nitrosamines (De
44 Mey, De Maere, Paelinck, & Fraeye, 2017). Nitrosamines exist in two forms, volatile and
45 non-volatile, depending on their boiling point. Nitrosamines are considered as volatile when
46 they can be distilled from strong solutions of salts or caustic soda with a minimum recovery
47 of 80%. The involvement of nitrosamines in carcinogenesis is due to their enzymatic
48 degradation into reactive diazo compounds, which can cause damage to DNA (Tricker &
49 Preussmann, 1991). Nitrite can also react with heme iron to form nitrosylheme

50 (nitrosylation), responsible for the characteristic color of cured meat (Honikel, 2008), and
51 with free thiol groups (S-nitrosation) to form nitrosothiols (Kuhnle & Bingham, 2007). Both
52 nitrosylheme and nitrosothiols are known to release NO and can act as nitrosating agents
53 during digestion (Kuhnle & Bingham, 2007). Nitrate is not dangerous at the levels found in
54 meat products but it can be reduced into nitrite by certain microorganisms and in the human
55 body, in the oral cavity (van Maanen, van Geel, & Kleinjans, 1996).

56 Lipid oxidation in meat can lead to the formation of mutagenic aldehydes (Sottero,
57 Leonarduzzi, Testa, Gargiulo, Poli, & Biasi, 2019). Several epidemiological studies have
58 suggested that excessive consumption of cooked meat and cured meat is associated with an
59 increased risk of colorectal cancer (Larsson & Wolk, 2006). To limit oxidation and the
60 formation of nitrosamines, antioxidants such as sodium ascorbate are usually added to cooked
61 cured meat. Nonetheless, ascorbate has an ambivalent role, depending on the physico-
62 chemical environment. By reducing heme iron (Richards, 2013) and nitrous acid into nitric
63 oxide (Honikel, 2008; Skibsted, 2011), ascorbate contributes to heme iron nitrosylation. In
64 some conditions, ascorbate can also act as a pro-oxidant (Amaral, da Silva, & da Silva
65 Lannes, 2018).

66 Recent eating habits encourage consumers towards new cured meat consumption patterns and
67 people are consuming more and more ready-to-eat meals. These new practices have to be
68 taken into account when studying the chemical reactivity of nitrite. For instance, heating
69 pizzas will necessarily lead to a second cooking for pizza toppings such as ham, bacon bits,
70 and chorizo. The temperature applied is in most cases higher than 150°C. Yet such conditions
71 favor the formation of mutagenic compounds and especially nitrosamines in cured meat
72 (Herrmann, Duedahl-Olesen, & Granby, 2015a).

73 To prevent the risk of nitroso-compound production in cured meat, it is essential to
74 investigate the mechanisms involved in their formation. As a precursor of the formation of

75 NOCs, nitrite is suspected to be involved in carcinogenesis (Santarelli et al., 2010).
76 Nevertheless, the dose effect of nitrite, combined with the addition of ascorbate and their
77 interaction during cured meat processing are little documented (Herrmann, Granby, &
78 Duedahl-Olesen, 2015b; Higuero, Moreno, Lavado, Vidal-Aragon, & Cava, 2020) and, to our
79 knowledge, the second cooking was never investigated.

80 The aim of this study is thus to get a better understanding of the reactions and interactions of
81 the compounds involved in NOC formation in cured products. To this end, a model of cooked
82 cured meat was studied to assess the influence of composition, *i.e.* sodium nitrite doses and
83 ascorbate addition, and chemical interactions. Heat treatment at high temperature mimicking
84 a second cooking of products was performed to evaluate the chemical reactivity of nitrite and
85 ascorbate, through the formation of non-volatile nitrosamines and nitrosothiols.

86

87 2. Material and Methods

88 2.1. Reagents

89

90 All the reagents, *i.e.* ferrozine, ascorbate, acetone, hydrochloric acid, thiobarbituric acid,
91 ethanol, tempol, o-phenylenediamine (OPDA), and the Griess reagent kit for nitrite and
92 nitrate assays (ref : 23479-1KT-F) were purchased from Sigma Aldrich (Saint Louis, USA).
93 Vivaspin® 2 system (PES membrane with a cut-off of 5 kDa) (ref: VS0212) and the syringe
94 filters with 0.22 µm regenerated cellulose membranes (ref: 17761) were purchased from
95 Sartorius (Göttingen, Germany).

96

97 2.2. Preparation of cured meat samples

98 2.2.1. Cured and cooked meat model preparation

99

100 The experiment was carried out on thirty-three porcine shoulders from Piétrain x (Landrace x
101 Large White) pigs (100 kg weight), slaughtered in a commercial slaughterhouse after electric
102 stunning. The shoulders were defatted, derinded, denerved and deboned. To prevent inter-
103 animal and muscle variability, muscles of all animals were pooled and frozen at -20°C prior
104 to meat preparation.

105 This model is based on the DCNO model (Dark Cooked with Nitrite and Oxidized), reported
106 by Santarelli et al. (2010), by its composition in heme iron and nitrite. After thawing, meat
107 was cut into large pieces (approximately 50mm), mixed with brine under vacuum, and then
108 churned for one night before cooking at a heat of 68.5°C for 48 min. Brine (containing
109 natural NaCl salt and sodium nitrite salt at various concentration, and dextrose) was mixed
110 with meat at 100 g/kg under vacuum conditions. The exact composition of the brine is
111 displayed in Supplementary Material 1. Different brine compositions in sodium nitrite were
112 used, leading to final rates of 40 ppm – the sufficient rate to add color and aroma (Alahakoon,
113 Jayasena, Ramachandra, & Jo, 2015), 80 ppm – the short-term goal of the profession, and 120
114 ppm – the maximum authorized by the French Code of Practice (Code des usages de la
115 charcuterie, de la salaison et des conserves de viande. *IFIP Ed.* Institut du Porc ; 2017). A
116 control without added nitrite was also performed. Sodium ascorbate was added at 300 mg per
117 kg of meat, and a control without ascorbate was performed with 120 ppm of sodium nitrite. In
118 the following article, 0ppm+A; 40ppm+A; 80ppm+A; 120ppm+A correspond respectively to
119 0, 40, 80 and 120 ppm of sodium nitrite added in the presence of ascorbate. Samples
120 120ppm-A correspond to 120 ppm of sodium nitrite added without ascorbate. Samples were
121 stored under vacuum at 4°C until analysis or a second cooking stage.

122

123 2.2.2. Second cooking of the cured and cooked meat model

124

125 The samples described in previous section were heated in an oven at 220°C for 6 min.
126 Moreover, to be more representative of domestic habits, samples treated with 120 ppm nitrite
127 with or without ascorbate were heated under two other temperature duration conditions:
128 180°C for 7 min and 260°C for 4 min. The experimental design of the cooking processes is
129 displayed in Supplementary Material 2.

130 The second cooking was performed on a thin water film (3-5 millimeters) to limit
131 dehydration and mimic what happens with the wet constituents of a pizza for instance. The
132 samples from the second cooking were frozen and stored at - 80°C until analysis.

133

134 2.3. Biochemical characterization

135 2.3.1. Determination of nitrite and nitrate content

136

137 Nitrite and nitrate were extracted using the method of Ionescu, Zara, Aprodu, Vasile, and
138 Carac (2006) , with slight modifications. Samples (2 g) were homogenized with a Polytron
139 PT 2100 (KINEMATICA AG, Switzerland) for 30 seconds in 10 mL of water and 2.4 mL of
140 2% NaOH. Then, 10 mL of water and 420 µL of 7.4% HCl, was added to samples to reach
141 pH 8. The samples were homogenized with a vortex and incubated at 50°C for 15 minutes.
142 Finally, the samples were deproteinized by centrifugation with a Vivaspin® 2 system with a 5
143 kDa cut-off threshold (16°C, 900g for 75 min with an SL 40R centrifuge from Thermo
144 Scientific, Waltham, USA). The supernatants were eliminated and the filtrates were used for
145 assays.

146 Nitrite and nitrate ion contents were then determined in the filtrates using the Griess reaction
147 with a Sigma-Aldrich colorimetric assay kit. Absorbance was measured on a microplate at
148 540 nm on a MULTISKAN SPECTRUM spectrophotometer from Thermo Scientific
149 (Waltham, USA). Residual nitrite and nitrate were expressed in mg/ kg of wet matter (ppm).

150

151 2.3.2. Determination nitrosothiol content

152

153 The filtrates (1 mL) prepared in section 2.3.1 were saturated with HgCl₂ (10 mg) to
154 specifically cleave the S-NO bonds of nitrosothiols (Gaston, 1999). The samples were filtered
155 on syringe filters with 0.22 µm regenerated cellulose membrane. Nitrite ion content was
156 measured using the Griess reaction with the Sigma-Aldrich colorimetric assay kit and the
157 difference between this measure and the level of nitrite ions initially present in the sample
158 (section 2.3.1) gave the nitrosothiol content (Gaston, 1999). Nitrosothiols were expressed in
159 mg S-NO/ kg of wet matter (ppm). A considerable loss of water was observed during the
160 second cooking stage (around 60%, data not shown). To prevent any artefact due to drip loss,
161 the nitrosothiol values were expressed in relation to dry matter in mg/ kg of dry matter.

162

163 2.3.3. Determination of non-volatile nitrosamine content

164

165 The filtrates prepared in section 2.3.1 were irradiated with a UV lamp (model LF 215.S, 254
166 nm, 2x15 W) from Uvitec (Cambridge, England) with a sample distance with source of d =
167 2cm. UV irradiation cleaves the S-NO bonds of nitrosothiols and the N-NO bonds of
168 nitrosamines and nitrosamides (Breider & von Gunten, 2017), giving a mixture of nitrite and
169 nitrate. C-nitroso and O-nitroso containing compounds, oximes, nitramines, and N-oxide are
170 stable under UV light and do not interfere in the assay (Breider & von Gunten, 2017).
171 Nitrosylheme iron does not interfere in the assays because of the withdrawal of nitroso-
172 myoglobin during Vivaspin centrifugation. If released, nitrosyl iron (Fe-NO) is rapidly
173 transformed, in oxidative medium, into more stable Fe[(OH)ONOO⁻]²⁻ (Fotiou, Fotiou, &
174 Deliconstantinos, 2009) with a UV resistant O-NO bond.

175 Irradiation kinetics were sampled at 1, 15, 30, 60 and 120 minutes of exposure. The content
176 of both nitrite and nitrate was measured using the Griess reaction. The concentration of $[\text{NO}^{2-}$
177 $+\text{NO}^{3-}]$ was plotted versus the inverse of time and extrapolation to $1/t = 0$, obtained by fitting
178 a second-degree polynomial function, giving the maximum rate of nitrite + nitrate released.
179 This measure gave the sum of nitrosothiol, non-volatile nitrosamine and nitrosamide content.
180 The subtraction of nitrosothiol content, obtained as described in section 2.3.2, gave the non-
181 volatile nitrosamine and the nitrosamide contents (Apparent Total N-Nitroso Compounds,
182 ATNC). Nevertheless, nitrosamides decompose rapidly and, contrary to nitrosamines, they do
183 not accumulate in food products, so their level is generally not detectable (Chow, Dhaliwal,
184 & Polo, 1984). Finally, non-volatile nitrosamines were expressed in mg N-NO/ kg of wet
185 matter (ppm) while, regarding nitrosothiols, the nitrosamine content after the second cooking
186 stage were expressed in relation to dry matter.

187

188 This method is described in more detail in Bonifacie et al. (submitted in MethodsX).

189

190 2.3.4. Determination of ascorbate content

191

192 Total ascorbate content and the two forms of ascorbate, *i.e.* oxidized and reduced, were
193 evaluated in the samples according to Vislisel, Schafer and Buettner (2007). The samples (0.8
194 g) were extracted in 4 mL of sodium acetate (2M, pH 5.5). After agitation and
195 homogenization, 1 mL of extract was mixed with a ratio of 1/5 in 250 μM EDTA prepared in
196 ethanol/ H_2O (90/10) (v/v). The samples were centrifuged at 4°C , 4000 rpm for 15 min. 100
197 μL of Tempol (4-hydroxy-2,2,6,6-tetramethylpiperidinyloxy) was added in 100 μL of the
198 samples. After agitation at 22°C at 600 rpm for 10 min, the ascorbate was oxidized into
199 dehydroascorbate (DHA). O-phenylenediamine (OPDA) (42 μL) was added and reacted with

200 DHA to form the fluorescent OPDA-DHA complex. In the absence of Tempol, only the
201 naturally oxidized ascorbate present in the medium can react with OPDA. The difference
202 between the two measurements gave the level of reduced ascorbate. The fluorescence of the
203 OPDA-DHA complex was measured in 96-well black polystyrene microplates with a Jasco
204 FP-8300 spectrofluorometer, (Jasco, Oklahoma City, USA) ($\lambda_{ex/em} = 345/425\text{nm}$). The
205 results were expressed in ppm.

206

207 2.3.5. Determination of non-heme iron content

208

209 The non-heme iron content and the $\text{Fe}^{2+}/\text{Fe}^{3+}$ ratio were measured by the ferrozine assay,
210 according to Stolze, Dadak, Liu and Nohl (1996) and adapted to meat samples. Non-heme
211 iron was separated from heme iron by a centrifugation method using a Vivaspin® 2 system at
212 18°C , 4000 rpm for 75 min. The absorbance of the ferrozine-iron complex was measured at
213 562 nm on a multiskan spectrum from Thermo Scientific (Waltham, USA) and a standard
214 calibration curve is used to calculate the non-heme iron content in samples (Supplementary
215 Material 3). The results were expressed in μM .

216

217 2.3.6. Evaluation of heme iron nitrosylation

218

219 The selective extraction of nitrosylheme was carried out by adding acetone to the sample, at a
220 ratio of 4:1 (v/w) with 300 μL of water. After homogenization with a polytron, the samples
221 were filtered on folded filters (ref: 1202-150, Whatman). The level of nitrosylheme was
222 evaluated by measuring specific absorbance at 540 nm and using an absorption coefficient of
223 $11,3 \text{ mM}^{-1} \text{ cm}^{-1}$ (Hornsey, 1956) on a Jasco V-770 spectrometer (Jasco, Oklahoma City,
224 USA). In parallel, the total heme iron was estimated in the form of acid haematin by

225 extraction in acidic acetone. After homogenization, the samples were filtered and the total
226 heme iron was estimated by measuring specific absorbance at 512 nm with an absorption
227 coefficient of $9.52 \text{ mM}^{-1} \text{ cm}^{-1}$ (Hornsey, 1956). The nitrosylation of heme iron was expressed
228 as the percentage of nitrosylheme to total heme iron.

229

230 2.3.7. Lipid oxidation measurement

231

232 Lipid extraction was performed using the procedure of Folch, Lees and Sloane-Stanley
233 (1957) with slight modifications. To do this, the samples (1g) were ground in 4 mL of water
234 with a polytron. Ethanol-dichloromethane (1/2 (v/v)) was added to the sample extract at a
235 ratio of 3 (solvent):1 (sample) and the mixture was shaken vigorously. After evaporation the
236 concentration of lipids was calculated by weighing the dry residues. In parallel, lipid
237 oxidation was measured by the thiobarbituric acid reactive substances (TBARS) method
238 according to Mercier, Gatellier, Viau, Remignon, and Renerre (1998). The results of lipid
239 oxidation were then expressed as mg of malondialdehyde (MDA) per g of lipids.

240

241 2.3.8. Protein oxidation measurement

242

243 Protein oxidation was evaluated by the free thiols content measurement, with a modification
244 of Ellman's method using 2,2'-dithiobis(5-nitropyridine) (DTNP) (Winterbourn, 1990). The
245 adduct products of DTNP on free thiols were evaluated by their absorbance at 386 nm on a
246 Jasco V-770 spectrometer (Jasco, Oklahoma City, USA) using an absorption coefficient of 14
247 $\text{mM}^{-1} \cdot \text{cm}^{-1}$. In parallel, the total protein content was evaluated by the Biuret method. The
248 results were expressed as nanomoles of DTNP bound per milligram of protein.

249

250 2.3.9. Dry matter

251

252 Dry matter was evaluated by heating samples at 105°C for 16 hours and by weighing the dry
253 residues (da Silva, Ezequiel, Diesel, Lunedo, & Cleef, 2011).

254

255 2.4. Statistical analysis

256

257 The statistical analyses were carried out with STATISTICA software (version 13.3) from
258 TIBCO Software Inc. (Palo Alto, USA). The values for each experimental condition were
259 reported as the mean \pm standard error of the mean (SEM) of six independent repetitions. The
260 effect of nitrite and ascorbate treatment in the cooked model were assessed independently by
261 a one-way analysis of variance (ANOVA): (0ppm+A ; 40ppm+A ; 80ppm+A ; 120ppm+A)
262 and (120ppm+A ; 120ppm-A) respectively. The effect of second cooking conditions and
263 sodium nitrite doses were tested together by ANOVA (0ppm+A ; 40ppm+A ; 80ppm+A ;
264 120ppm+A at 220°C for 6 minutes). The effect of second cooking conditions and ascorbate
265 addition were tested together by ANOVA (120ppm+A ; 120ppm-A at 180°C for 7 minutes ;
266 220°C for 6 minutes; 260°C for 4 minutes). All ANOVA were performed using a confidence
267 level of 5%, and the *post hoc* test used was the Tukey test. A principal component analysis
268 (PCA) was also performed with STATISTICA software using centered reduced data.

269

270 3. Results and Discussion

271

272 In the cured and cooked meat model (section 3.1.) all the nitroso compounds and their
273 precursors, catalysts, and inhibitors were determined in relationship with sodium nitrite doses
274 and ascorbate. In the second cooking model (section 3.2.), only nitrosothiols and nitrosamines

275 were evaluated with regard to the doses of sodium nitrite added and the presence or not of
276 ascorbate. At high temperature, as used in the second cooking, nitrosylheme can lead to
277 various products, like hemochrome, hemichrome, oxo ferryl species and biliverdin, that are
278 impossible to evaluate because of their visible absorbance very close to that of nitrosylheme.
279 Moreover, iron or nitrosyl iron can be released from nitrosylheme at high temperature making
280 it difficult to evaluate the different forms of iron. Thus, nitrosylheme was not studied in the
281 second cooking stage. Concerning nitrite and nitrate, their levels were not affected by the
282 second cooking (results not shown).

283

284 3.1. Biochemical composition and molecular interactions in the cured and cooked 285 meat model

286 3.1.1. Impact of sodium nitrite and ascorbate added on residual nitrite and nitrate

287

288 Residual nitrate and nitrite are presented in Table 1. Overall, the increasing doses of added
289 sodium nitrite led to an increase of residual nitrate and nitrite. The basal level of residual
290 nitrate was higher than the values generally cited in the literature for fresh meat. For instance,
291 using the ion chromatography method, Iammarino and di Taranto (2012) found a maximum
292 level of 36.5 ppm of nitrate for red meat, while our results showed a value of 45.7 ppm in the
293 case of 0 ppm+A. The dose-response relationship of added sodium nitrite was similar
294 between 0, 40, and 80 ppm. Only a significant increase of residual nitrite was observed for
295 the 120 ppm+A condition ($p=0.001$).

296 Overall, the levels of residual nitrate observed in our study were higher than those frequently
297 reported in the literature. This could be explained by the determination method, in particular
298 the extraction conditions we used. Indeed, (Hamano, Mitsuhashi, Tanaka, Matsuki, Oji, &
299 Okamoto, 2014) demonstrated that deproteinization by ultrafiltration, as performed in our

300 study, led to an increase of 2.3 times more extracted nitrate than classical chemical
301 deproteinization. Indeed, in chemical deproteinization nitrate can co-precipitate with proteins
302 leading to an underestimation of its concentration. Furthermore, the processing used to
303 prepare the cured and cooked model was chosen to optimize the study of these chemical
304 reactions, and is considered as very oxidative. Indeed, freezing/thawing as well as mixing and
305 cooking, increase oxidation in addition to the using of red meat (shoulder). These different
306 steps could lead to the release of ammonia from oxidized proteins and, in very oxidative
307 medium, ammonia can be oxidized into hydroxylamine by free oxygenated radicals and then
308 into nitrate (Fotiou et al., 2009). Moreover, according to Honikel (2008), products formulated
309 only with nitrite show residual nitrate concentrations (around 1/3 of the added nitrite) in the
310 finished product due to the oxidation of nitrite to nitrate.

311 Contrary to nitrate, no residual nitrite was observed in the 0 ppm+A model. This result is in
312 good agreement with that of Immarino and di Taranto (2012) and Higuero et al. (2020), who
313 observed nitrate but no nitrite in pork. Nitrite residual content increased significantly
314 ($p=0.001$) (Table 1) with the sodium nitrite added. This has also been reported by Higuero et
315 al. (2020), who described that the amount of residual nitrite and nitrate increased by
316 increasing the dose of nitrite/nitrate added.

317 In the cured model with 120 ppm of sodium nitrite, a significant increase of residual nitrite
318 and nitrate was observed when adding ascorbate ($p=0.001$) (Table 1). In the absence of
319 ascorbate, more nitroso compounds could be formed perhaps explaining the difference
320 observed here. This hypothesis will be confirmed by the results in section 3.1.5., which show
321 considerably more S-nitrosation in the absence of ascorbate.

322

323 3.1.2. Impact of sodium nitrite added to ascorbate chemistry

324

325 As mentioned in section 2.2.1., sodium ascorbate was added at the level of 300 mg per kg of
326 cured meat, and a control without ascorbate was performed with 120 ppm of sodium nitrite.
327 In our model, only oxidized ascorbate (dehydroascorbate) was observed (Table 1). This
328 oxidation of ascorbate might be explained by its implication in iron and nitrous oxide
329 reduction during the curing process. When no nitrite was added, *i.e.* 0 ppm+A, we observed
330 only 98 ppm of residual dehydroascorbate. This low level could be due to the strong
331 oxidation occurring during cooking and freezing, leading to the irreversible oxidation of
332 dehydroascorbate into deketogulonate (Vieira, Teixeira, & Silva, 2000). A significant
333 ($p=0.001$) effect of sodium nitrite addition was observed on dehydroascorbate, whose levels
334 varied from 312 to 357 ppm. These values were higher than the dose of 300 ppm added,
335 probably due to the natural ascorbate content in fresh meat and due to the natural
336 decomposition of ascorbate during cooking (Jutkus, Li, Taylor, & Mauer, 2015). For
337 instance, around 24 ppm of ascorbate was measured in fresh pork meat by Lahucky, Bahelka,
338 Novotna and Vasickova (2005). In our study, a value of 42 ppm was observed in the samples
339 without ascorbate added. This higher value, when compared to Lahucky et al. (2005) could
340 be explained by a drip loss during the process, leading to a concentration of ascorbate. This
341 result highlights the protective effect of nitrite against the oxidative degradation of
342 dehydroascorbate into non-vitamin compounds during the curing process.

343

344 3.1.3. Impact of sodium nitrite and ascorbate added in non-heme iron chemistry

345

346 Total non-heme iron was determined in the cured meat model and each form, *i.e.* reduced
347 (ferrous) and oxidized (ferric) were also quantified (Table 1). The content of non-heme iron
348 was significantly ($p=0.0000001$) higher when no sodium nitrite was added to the model, *i.e.* 0
349 ppm+A condition, compared to the conditions with nitrite, regardless of the concentration.

350 This result can be explained by the release of heme iron during the curing and cooking
351 process in the absence of added sodium nitrite, as will be described in section 3.1.4. PCA,
352 (Supplementary Material), showed that the variables free iron and heme iron were opposite
353 on the first axis and the correlation matrix, showing a significant (-0.81) negative correlation
354 between them and confirming this plausible mechanism. This mechanism implicates
355 oxygenated free radical chemistry, otherwise known as Fenton chemistry. Indeed, it has been
356 described that hydrogen peroxide can release iron from heme through the oxidative cleavage
357 of the porphyrin ring (Amaral et al., 2018). Hydrogen peroxide is naturally present in meat. It
358 can be formed by the oxidation of ferrous iron in the presence of oxygen ($Fe^{2+} + O_2 \rightarrow$
359 $Fe^{3+} + O_2^{\circ-}$) followed by the dismutation of superoxide ion ($2O_2^{\circ-} + 2H^+ \rightarrow H_2O_2 + O_2$)
360 (for a review on iron catalyst chemistry; Bechaux, de La Pomélie, Théron, Santé-Lhoutellier,
361 & Gatellier, 2018). In presence of nitrite, nitric oxide can react with superoxide radicals to
362 form a stable peroxynitrite ($NO^{\circ} + O_2^{\circ-} \rightarrow ONOO^-$) (Bechaux et al., 2018), thereby
363 preventing the formation of hydrogen peroxide and so increasing heme iron stability. The
364 effect of nitrite was mainly observed between 0 ppm+A and 40 ppm+A and a further increase
365 of sodium nitrite had only a slight effect on the non-heme iron content. The addition of
366 sodium nitrite in the model induced a significant increase of approximately 10% in ferrous
367 iron between 0 ppm+A and 40 ppm+A, which may be due to the ability of nitric oxide to
368 reduce ferric iron ($Fe^{3+} + NO^{\circ} \rightarrow Fe^{2+} + NO^+$) (Honikel, 2008). The addition of ascorbate
369 at the highest dose of nitrite did not impact the total non-heme iron content (Table 1).
370 However, in the absence of ascorbate, ferric iron is predominant (92%). These results are in
371 agreement with those of Realini et al. (2013) who reported that in red muscles of pigs, non-
372 heme iron is mainly present in ferric form (93%). Indeed, the addition of ascorbate induces
373 the reduction of ferric iron into ferrous iron according to the reaction $Fe^{3+} + Asc^- \rightarrow$
374 $Fe^{2+} + DHA$.

375

376 3.1.4. Impact of sodium nitrite and ascorbate added to heme iron

377

378 Total heme iron and nitrosylated-heme iron were quantified in our model and the percentage
379 of nitrosylation was calculated (Fig. 1). In the absence of sodium nitrite, the content of total
380 heme iron was significantly lower ($p=0.000001$) than in conditions where it was added,
381 regardless of its concentration. As described in section 3.1.3, these results can be explained
382 by the antioxidant capacity of nitrite that prevents heme iron release during the curing and
383 cooking process, supported by a positive correlation between heme iron and nitrite (0.65). We
384 can also hypothesize that this result was due to the stronger stability of nitrosylheme. Indeed,
385 the nitrosylation of heme prevents the release of iron from the porphyrin ring during
386 oxidative stress (Juckett et al., 1998). It is noteworthy that the sum of non-heme and heme
387 iron content led to approximately the same values, whatever the treatment of meat, showing a
388 notable conservation of the iron level in the model. Adding sodium nitrite significantly
389 increased ($p=0.000001$) nitrosylation without a dose effect. These results are in accordance
390 with Higuero et al. (2020) who described a significant increase in the rate of
391 nitrosomyoglobin in dry-cured loins of between 0 and 37.5 ppm of added nitrite/nitrate, not
392 significantly affected by higher nitrite doses.

393 The addition of ascorbate did not affect nitrosylation (Fig. 1). Since the process was mainly
394 conducted under vacuum conditions, the formation of deoxymyoglobin ($Mb(Fe^{2+})$) might
395 have been favored, independently of the presence of ascorbate. Deoxymyoglobin is the form
396 in which myoglobin preferentially fixes nitric oxide to form nitrosomyoglobin ($Mb(Fe^{2+}) -$
397 NO) according to the following reaction $Mb(Fe^{2+}) + NO^{\circ} \rightarrow Mb(Fe^{2+}) - NO$ (Richards,
398 2013). Consequently, the experimental conditions, with a process mainly performed under
399 vacuum, probably made the reducing effect of ascorbate unobservable.

400

401 3.1.5. Impact of sodium nitrite and ascorbate added on nitrosation reactions

402

403 The products of nitrosation, *i.e.* nitrosothiols and nitrosamines, were measured in the cured
404 and cooked meat (Fig. 2). The non-volatile nitrosamines were not detected except at a low
405 value of 0.2 ppm for 40 ppm of added sodium nitrite. In nitrite-cured pork meat, Van Hecke,
406 Vossen, Vanden Bussche, Raes, Vanhaecke, & De Smet, 2014 did not show an NOC-induced
407 DNA adduct, demonstrating the very low level of nitrosation during the process. Herrmann et
408 al. (2015a) found a low level of nitrosamines in smoked ham, between 0.03 ppm of non-
409 volatile N-nitroso-methylaniline (NMA) and 2 ppm of non-volatile nitrosamine N-nitroso-
410 thiazolidine-4-carboxylic acid (NTCA). These authors also found up to 0.5 ppm of non-
411 volatile nitrosamine N-nitroso-thiazolidine-4-carboxylic acid (NTCA) in bacon and 4 ppm in
412 salami. Moreover, according to Massey, Key, Jones and Logan (1991) the Apparent Total N-
413 Nitroso Compounds levels in bacon are in the range of 0.4 to 6.8 ppm (N-NO). Nevertheless,
414 all these products underwent manufacturing processes (smoking, drying and salting) that
415 differed from our model, probably favoring the formation of nitrosamines, which could
416 explain these higher values.

417 On the contrary, nitrosothiol content increased with sodium nitrite addition, with a significant
418 effect observed at 80 ppm. Joosen et al. (2009) reported up to 22 ppm of nitrosothiols in
419 bacon, a value higher than those observed in our samples which ranged from 0.5 to 4 ppm.
420 The smoking step applied to the bacon could have promoted the formation of NOCs.

421 At 120 ppm of sodium nitrite added without ascorbate, a six-fold increase ($p=0.0000001$) of
422 nitrosothiols was observed when compared to the corresponding samples with ascorbate,
423 while no nitrosamines were observed. This result clearly shows a protective effect of
424 ascorbate against the nitrosation process. In an acidic condition and in the presence of

425 ascorbate, nitrosylation was found to predominate over nitrosation (Honikel, 2008; Skibsted,
426 2011). This was due to the formation of nitrous acid ($NO_2^- + H^+ \rightarrow HNO_2$) followed by its
427 reduction into nitric oxide ($HNO_2 + H^+ + e^- \rightarrow NO^\circ + H_2O$) while, in the absence of
428 ascorbate, nitrosation was favored by the formation of nitrosonium from nitrous
429 acid ($HNO_2 + H^+ \rightarrow NO^+ + H_2O$). Consequently, we can hypothesise that in our model,
430 nitrosothiols were predominantly formed by the reaction of nitrosonium with the free thiols.
431 Moreover, the correlation matrix (Supplementary Material), showed a negative correlation (-
432 0.52) between nitrosothiols and ascorbate in its oxidation form (dehydroascorbate), which
433 was the only form observed in the model. In addition, a possible increase of protective effect
434 of ascorbate against nitrosation during the process could be assumed. Indeed, both ascorbate
435 and dehydroascorbate can react with nitrous acid to form (di)nitroso-
436 ascorbic/dehydroascorbic acid intermediates (Skibsted, 2011), thereby preventing the reaction
437 of a nitrite derivate with free thiols.

438

439 3.1.6. Impact of sodium nitrite and ascorbate added in oxidation mechanisms

440

441 The lipid oxidation determination ranged from 6 μ g of MDA per g of lipids for the 120
442 ppm+A to 18 μ g of MDA for the no nitrite condition (Fig. 3). These values correspond to
443 approximately 0.5 and 1.5 mg MDA/kg of meat. These values are below the strong rancid
444 smell threshold generally reported in the literature, from 1.5 mg MDA/kg of meat (Tarladgis,
445 Watts, Younathan, & Dugan, 1960). So, all these products were available for consumption. In
446 the 0 ppm+A model, lipid oxidation was significantly ($p=0.0000001$) higher than in other
447 conditions. This difference can be explained by nitrite, which acts as a powerful antioxidant
448 in such products (Berardo et al., 2016). Our results are in line with those reported for dry-
449 cured loins (Higuero et al., 2020) and cooked cured meat products (Karwowska, Kononiuk &

450 Wojciak, 2019; Santarelli, et al., 2010). Also, as described in section 3.1.3., the absence of
451 nitrite leads to the increase of free iron. Free iron is known to promote lipid oxidation by the
452 Fenton reaction, leading to the formation of oxygenated free radicals, and by reacting with
453 lipoperoxides to form alcoxyle and peroxy free radicals ($Fe^{2+} + LOOH \rightarrow Fe^{3+} + LO^{\circ} +$
454 OH^{-}) and ($Fe^{3+} + LOOH \rightarrow Fe^{2+} + LOO^{\circ} + H^{+}$) (Bechaux et al., 2018) . Moreover, nitric
455 oxide reacts with superoxide radicals, as previously described in section 3.1.3., and lipid
456 radicals to form stable LONO and unstable LOONO that decomposes into stable LONO₂
457 ($NO^{\circ} + L(O)O^{\circ} \rightarrow L(O)ONO$) (Bechaux et al., 2018), thereby ending the oxidation process.
458 The addition of nitrite decreases lipid oxidation but no significant dose-response relationship
459 was observed. These mechanisms were confirmed by PCA and the correlation matrix,
460 (Supplementary Material), showing a positive correlation (0.78) of TBARS and non-heme
461 iron. On the contrary, a significant negative correlation (-0.73) was observed between
462 TBARS and residual nitrite. Van Hecke et al. (2014) described a considerable decrease of
463 lipid peroxidation in cured and cooked meat, with 120 ppm of nitrite added, when compared
464 to uncured meat. Moreover, a gradual decrease in TBARS values with the increase in amount
465 of sodium nitrite added (from 0 to 150 ppm) was previously described by Karwowska et al.
466 (2019) in cooked meat products. Free thiol content was also estimated. Contrary to lipid
467 oxidation, no nitrite effect was observed for thiols oxidation (results not shown). This was
468 probably due to considerable thiol oxidation in the previous steps of the process (freezing,
469 thawing and mincing).

470 The addition of ascorbate did not affect lipid oxidation (Fig. 3). It is more likely that the high
471 antioxidant power of nitrite, described above, masked the antioxidant effect of ascorbate. In
472 the conditions of the process applied, no greater effect of nitrite and ascorbate was observed
473 on lipid oxidation. These results are in agreement with those of Berardo et al. (2016) who did
474 not report a higher effect of nitrite and ascorbate when compared to the separate addition of

475 nitrite or ascorbate on lipid oxidation in dry fermented sausages. Moreover, ascorbate can act
476 as a pro-oxidant (Amaral et al., 2018), especially in the presence of a high amount of iron, by
477 regenerating the ferrous form, which acts in the reactions previously described. In addition,
478 ascorbate did not affect thiol oxidation (results not shown).

479

480 3.2. Effect of sodium nitrite doses and ascorbate on nitrosation during the second cooking 481 stage of the cured and cooked meat model

482

483 Cured meat products are sometimes recooked at high temperature before consumption, such
484 as ham in pizza toppings. Sausages and bacon are also often grilled or fried at high
485 temperature. Thus, it is of great interest to evaluate the extent of nitrosation that occurs
486 during these cooking processes. A significant increase of non-volatile nitrosamines was
487 observed after the second cooking stage ($p=0.0000001$) and sodium nitrite addition
488 ($p=0.00007$) (Fig. 4A). Nitrosamines increased significantly ($p=0.00007$) between 0 and 40
489 ppm of added sodium nitrite and the further increase was not significant. An high increase of
490 nitrosamine content in dried and fried sausages with increasing doses of nitrite from 0 to 350
491 ppm was already described by Herrmann et al. (2015b). Such high levels of nitrosamines
492 could negatively impact the health quality of cured meat. Thus, to prevent the formation of
493 these unsuitable products it is necessary to better understand the reactions involved in their
494 formation during the second cooking process. This is why different mechanisms are
495 described in this paper to better understand the effect of thermal processes on nitrosation.

496 The nitrosothiol content increased significantly ($p=0.000001$) when increasing the sodium
497 nitrite content (Fig. 4B). But, contrary to nitrosamines, no second cooking effect was
498 observed on nitrosothiols. The absence of a second cooking effect could be due to the low
499 stability of nitrosothiols at high temperature. Indeed, nitrosothiols begin to decompose from

500 130°C upwards to release NO (Melvin, Jones, Lutzke, Allison & Reynolds, 2019). In
501 contrast, most non-volatile nitrosamines are stable at high temperature; their boiling point
502 generally exceeds 200°C. For example, the most common non-volatile nitrosamines N-
503 nitrososarcosine (BP 221°C) and N-nitrosoproline (BP 263°C), are particularly thermostable.
504 Nitrosothiols are not described in the literature as dangerous by themselves, but
505 transnitrosation from nitrosothiols to secondary amines (Wainright, 1986) during digestion is
506 possible with, in this case, the risk of formation of mutagenic compounds (Kuhnle &
507 Bingham, 2007).

508 One of the ways described in the literature to reduce nitrosation is the addition of ascorbate to
509 products. This is why we decided to evaluate during the second cooking of cured meat the
510 impact of dehydroascorbate, which was the only form of ascorbate after the initial cooking
511 (Table 1). Ascorbate is thermosensitive, and its activity rapidly decreases after 95°C.
512 Nevertheless, dehydroascorbate, is more resistant to thermal treatment (Vieira et al., 2000)
513 and considerable residues were observed after the second cooking. These residues were
514 measured at 58%, 93% and 22% for the 180°C for 7 min, 220°C for 6 min and 260°C for 4
515 min treatments, respectively.

516 During the second cooking stage, dehydroascorbate limited the formation of nitrosamines
517 (Fig. 5A) especially for the condition 180°C for 7 min ($p=0.007$). These results were in
518 agreement with those of Mottram, Patterson, Rhodes & Gough (1975), who showed a
519 reduction in the production of N-nitrosodimethylamine (NDMA) in the presence of ascorbate.
520 In addition, Herrmann et al. (2015b) showed the inhibitory effect of antioxidants, such as
521 erythorbate and ascorbate, on the formation of certain nitrosamines during heating treatments.
522 The mechanisms involved in this protective effect have not yet been thoroughly described,
523 but ascorbate, as well as dehydroascorbate, could react with nitrous acid to form (di)nitroso-
524 ascorbate or (di)nitroso-dehydroascorbate intermediates (Skibsted, 2011), possibly preventing

525 N-nitrosation. This study did not show a significant effect between the three different
526 time/temperature pairs on the nitrosamine content. Therefore, our study does not distinguish
527 the best time/temperature combination of second cooking in terms of consumer health.

528 Nitrosothiol content was significantly lower ($p=0.000001$) in the absence of
529 dehydroascorbate for each second cooking condition (Fig. 5B). This was significantly
530 ($p=0.0057$) more pronounced at 180°C for 7 min. Hence, this result showed a
531 dehydroascorbate effect contrary to that observed on nitrosamines in the second cooking
532 stage, and also a contrary effect to that observed on nitrosothiols in the cured and cooked
533 meat model (section 3.1.5.). This could be explained by a mechanism differing from that
534 described previously. During second cooking, dehydroascorbate could protect nitrosothiols
535 from reaction with transition metals that cleave the S-NO bond. These transition metals are
536 released at high temperature: iron is released in large amounts from ferritin and myoglobin
537 while copper is released to a lesser extent from various enzymes such as (Cu/Zn)-Superoxide
538 dismutase and cytochrome C oxidase. Ascorbate and dehydroascorbate are iron and copper
539 chelators (Fisher & Naughton, 2004) and so they could stabilize nitrosothiols (Melvin et al.,
540 2019).

541 As previously observed with nitrosamines, the second cooking stage did not show a
542 difference between the three time/temperature pairs on nitrosothiol content.

543 We can also notice that the increase of the molar nitrosamine concentration in the absence of
544 ascorbate was not significantly different from the decrease of the molar nitrosothiol
545 concentration in the same condition (results not shown), so that global nitrosation (N + S
546 nitrosation) was not affected by ascorbate. This observation could suggest a possible transfer
547 of NO groups from nitrosothiols to nitrosamines during the second cooking stage in the
548 absence of ascorbate. Such transnitrosation was already observed in cured meat by Wainright
549 (1986).

550

551 4. Conclusion

552

553 This study gives new insights on the formation of nitroso compounds in cured cooked
554 meat products. The absence of nitrite leads to a decrease of nitrosylheme but promotes the
555 release of heme iron from myoglobin. The iron released is then implicated in the formation of
556 free radicals, involving the formation of oxidation products, some of which are mutagenic.
557 The addition of nitrite promotes the formation of nitroso compounds without a dose-response
558 effect. However, we also observed the role of nitrite as an antioxidant, preventing lipid
559 oxidation and to the ascorbate degradation during the curing process. A second cooking stage
560 promotes the formation of non-volatile nitrosamines. Ascorbate limits the formation of these
561 compounds at moderate high temperature (180°C) but at the same time protects the
562 nitrosothiols from degradation.

563 This study contributes to elucidating the chemical reactivity of NOCs in a cured meat
564 model. The results obtained here will help to better understand the behavior of cured meat
565 products during digestion, and more specifically NOCs. Indeed, the physicochemical
566 conditions of the digestive tract, like oxygen pressure, low gastric pH, and reducing
567 conditions, favor the formation of nitrite derivatives such as nitrosonium ions and nitric oxide
568 that promote the formation of nitroso compounds. We will also study ways of limiting the
569 formation of these compounds in the digestive tract.

570

571 Conflict of interest

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574

575

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581

582

583 References

584

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705

Figure caption

Table 1. Effect of added doses of sodium nitrite and ascorbate in the cured and cooked meat model on the concentration of residual nitrite, nitrate, ascorbate and free iron. 0ppm+A; 40ppm+A; 80ppm+A; 120ppm+A correspond respectively to 0, 40, 80 and 120 ppm of sodium nitrite added in the presence of ascorbate. Samples 120ppm-A correspond to 120 ppm of sodium nitrite added without ascorbate. The effect of nitrite and ascorbate treatment in the cooked model were assessed independently by a one-way analysis of variance with Tukey post hoc test.

Figure 1. Effect of added sodium nitrite doses and ascorbate on the concentration of nitrosylated heme iron and total heme iron, expressed in mM, in the cured and cooked meat model. 0ppm+A; 40ppm+A; 80ppm+A; 120ppm+A correspond respectively to 0, 40, 80 and 120 ppm of sodium nitrite added in the presence of ascorbate. Samples 120ppm-A correspond to 120 ppm of sodium nitrite added without ascorbate. The effect of nitrite and ascorbate treatment in the cooked model were assessed independently by a one-way analysis of variance with Tukey post hoc test.

Figure 2. Effect of added sodium nitrite doses and ascorbate on the concentration of nitrosation products, expressed in ppm (mg/ kg of wet matter): nitrosothiols and nitrosamines in the cured and cooked meat model. 0ppm+A; 40ppm+A; 80ppm+A; 120ppm+A correspond respectively to 0, 40, 80 and 120 ppm of sodium nitrite added in the presence of ascorbate. Samples 120ppm-A correspond to 120 ppm of sodium nitrite added without ascorbate. The effect of nitrite and ascorbate treatment in the cooked model were assessed independently by a one-way analysis of variance with Tukey post hoc test.

Figure 3. Effect of added sodium nitrite doses and ascorbate on the concentration of TBA-RS in the cured and cooked meat model, expressed in $\mu\text{g MDA/g lipids}$. 0ppm+A; 40ppm+A; 80ppm+A; 120ppm+A correspond respectively to 0, 40, 80 and 120 ppm of sodium nitrite added in the presence of ascorbate. Samples 120ppm-A correspond to 120 ppm of sodium nitrite added without ascorbate. The effect of nitrite and ascorbate treatment in the cooked model were assessed independently by a one-way analysis of variance with Tukey post hoc test.

Figure 4. Effect of added sodium nitrite doses in the cured and cooked meat model before and after second cooking (220°C/ 6 min) on nitrosamine (A) and nitrosothiol (B) concentrations expressed in ppm (mg/ kg dry matter). 0ppm+A; 40ppm+A; 80ppm+A; 120ppm+A correspond respectively to 0, 40, 80 and 120 ppm of sodium nitrite added in the presence of ascorbate. The effect of nitrite in the cooked model were assessed independently by a one-way analysis of variance with Tukey post hoc test. The effect of second cooking conditions and sodium nitrite doses were tested together by a two-way analysis of variance.

Figure 5. Effect of different second cooking conditions (180°C for 7 min, 220°C for 6 min, 260°C for 4 min), with ascorbate (in white) and without ascorbate addition (in grey) (with 120 ppm sodium nitrite) on nitrosamine (A) and nitrosothiol (B) concentrations expressed in ppm (mg/ kg dry matter). The effect of nitrite and ascorbate treatment in the cooked model were assessed independently by a one-way analysis of variance with Tukey post hoc test. The effect of second cooking conditions and ascorbate addition were tested together by a two-way analysis of variance.

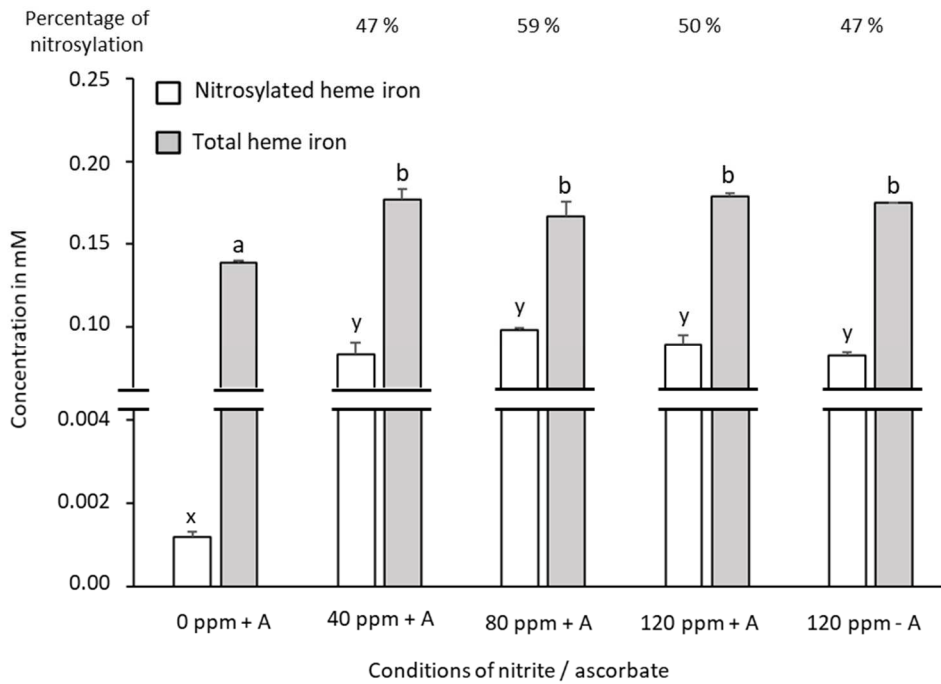


Figure 1. Effect of added sodium nitrite doses and ascorbate on the concentration of nitrosylated heme iron and total heme iron, expressed in mM, in the cured and cooked meat model. Values are mean \pm SEM of 6 independent determinations. Values without common superscripts, a, b for total heme iron and x, y for nitrosylated heme iron, differ significantly ($p < 0.01$).

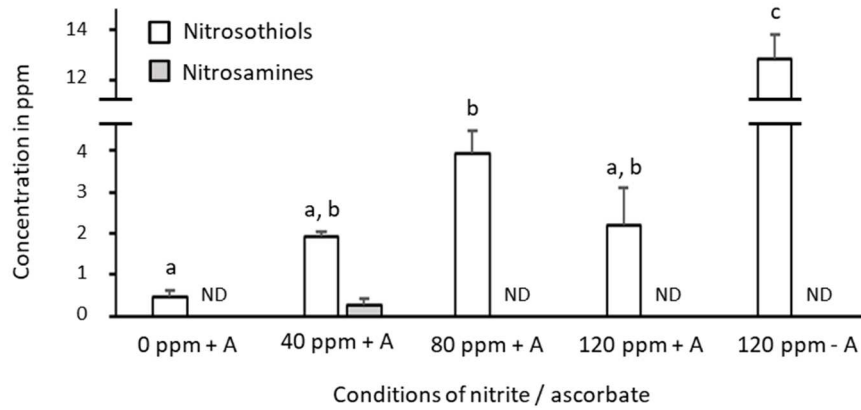


Figure 2. Effect of added sodium nitrite doses and ascorbate on the concentration of nitrosation products, expressed in ppm (mg/ kg of wet matter): nitrosothiols and nitrosamines in the cured and cooked meat model. Values are mean \pm SEM of 6 independent determinations. Values without common superscripts, a, b, c for total nitrosothiols, differ significantly ($p < 0.01$). ND for not detected.

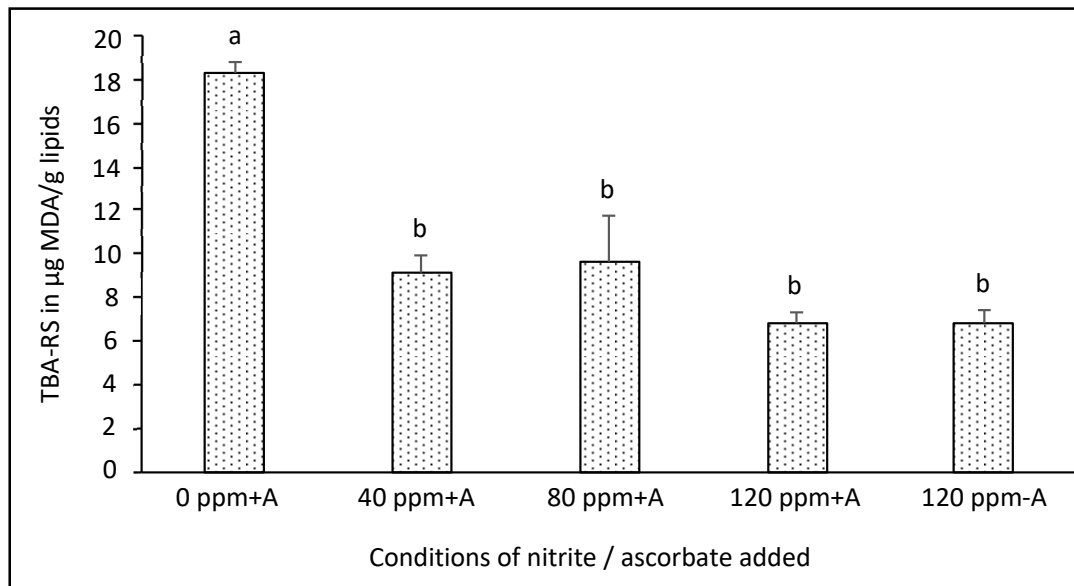


Figure 3. Effect of added sodium nitrite doses and ascorbate on the concentration of TBA-RS in the cured and cooked meat model, expressed in µg MDA/ g lipids.

Values are mean \pm SEM of 6 independent determinations. Values without common superscripts differ significantly ($p < 0.01$).

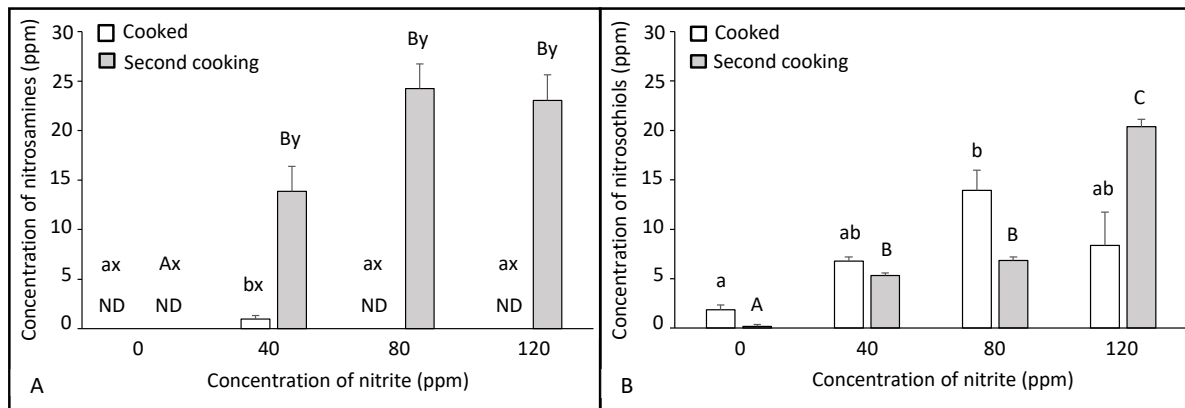


Figure 4.

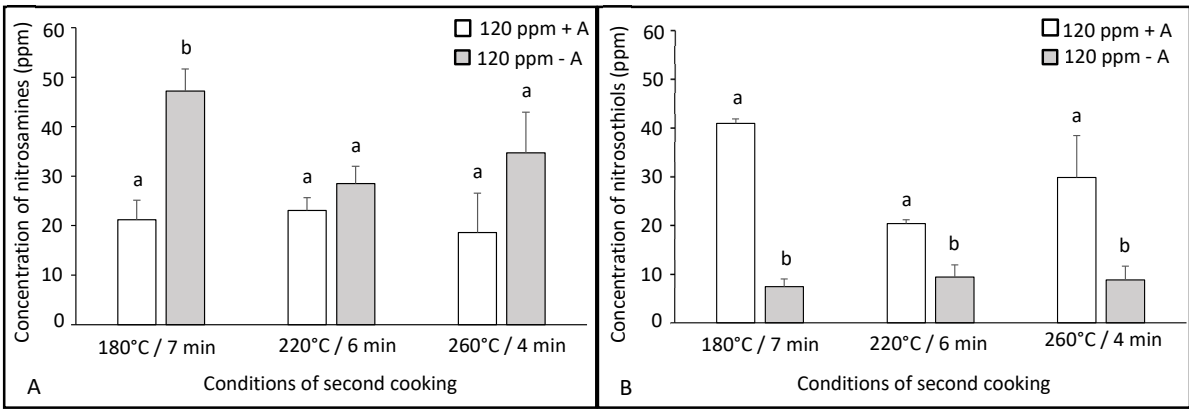


Figure 5.

Table 1. Effect of added doses of sodium nitrite and ascorbate in the cured and cooked meat model on the concentration of residual nitrite, nitrate, ascorbate and free iron.

	0 ppm+A	40 ppm+A	80 ppm+A	120 ppm+A	120 ppm-A
Residual NO ₂ (ppm)	0.00 ^a ± 0.00	7.74 ^b ± 0.16	18.33 ^c ± 0.64	34.28 ^c ± 1.75	29.91 ^d ± 1.43
Residual NO ₃ (ppm)	45.68 ^a ± 14.04	61.60 ^a ± 3.87	47.52 ^a ± 7.55	120.86 ^c ± 6.78	84.16 ^b ± 10.74
Oxidized ascorbate (ppm)	98.30 ^a ± 20.98	312.39 ^b ± 19.90	357.09 ^b ± 19.82	340.19 ^b ± 12.57	41.93 ^a ± 12.95
Free iron total (μM)	82.07 ^c ± 1.73	28.45 ^a ± 1.61	26.66 ^a ± 0.87	34.95 ^b ± 1.20	35.49 ^b ± 0.78
Ferrous iron Fe ²⁺ (%)	60.73 ^b ± 0.60	71.25 ^c ± 5.46	66.87 ^c ± 1.26	72.85 ^d ± 0.34	7.96 ^a ± 4.12
Ferric iron Fe ³⁺ (%)	39.27 ^c ± 0.62	28.75 ^b ± 5.46	33.13 ^b ± 1.26	27.15 ^a ± 0.34	92.04 ^d ± 4.12