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21 **Effect of different level of organic zinc supplementation on pork quality**

22

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34

35 **ABSTRACT**

36 This study investigated the effect of two levels of zinc chelate of glycine hydrate
37 (ZnGly) on performance, carcass characteristics and meat quality of growing-finishing pigs.
38 Thirty pigs (initial bodyweight: 61.0 ± 4.02 kg), were randomly assigned to three treatments
39 and fed *ad libitum* for 56 days with a basal diet supplemented with 0 mg/kg (control group,
40 CON), 45 mg/kg (Zn45 treatment) or 100 mg/kg (Zn100 treatment) of ZnGly. Both ZnGly
41 levels strongly reduced chilling carcass loss ($P < 0.001$). Micromineral content, color stability
42 and fatty acid profile of meat was not altered by ZnGly. Superoxide dismutase activity was
43 lowered by Zn45 compared to CON ($P = 0.007$); while, catalase activity was enhanced by
44 Zn100 ($P = 0.003$). Although ZnGly supplementation did not influence lipid oxidation in raw
45 meat and in meat homogenates incubated with pro-oxidant catalysts, Zn45 limited lipid
46 oxidation in cooked meat ($P = 0.037$). The results of this study demonstrated that
47 supplementing pigs with 45 mg/kg of ZnGly could improve the oxidative stability of pork
48 subjected to strong pro-oxidant conditions, but this effect need be further elucidated.

49

50 **KEYWORDS:** intramuscular fatty acids; antioxidant capacity; fat-soluble vitamins;
51 antioxidant enzymes; lipid oxidation; meat shelf-life.

52

53 **1. Introduction**

54 Zinc is an essential microelement for the regular growth and development of all animal
55 organisms (Sloup et al., 2017), as it plays a fundamental role in the activity of over 300 enzymes
56 and is required in more than 2,000 transcription factors involved in gene expression (Prasad
57 and Kucuk, 2002). Due to these multiple and important biological functions animal's diet need
58 to contain an adequate amount of this trace mineral in order to avoid deficiency. A zinc level
59 of about 50 mg/kg in the diet is recommended for growing-finishing pigs (NRC, 2012).
60 However, the innate zinc level of conventional swine diets is usually not sufficient to meet this
61 requirement, mainly due to the presence of phytic acid in cereals, which reduces zinc
62 availability by forming insoluble complexes (Oberleas et al., 1962; Sloup et al., 2017).
63 Therefore, zinc supplementation and/or the addition of the phytase enzyme, which releases zinc
64 from the phytate complex, are necessary practices to prevent zinc deficiency.

65 However, zinc supplementation also raises concerns related to environmental issues
66 (Monteiro et al., 2010) which justifies limitations in the levels of supplementation in different
67 countries (e.g., European Union regulation, 2016/1095). In this context, organic zinc sources,
68 such as amino acid chelates, increase the mineral bioavailability, thus allowing to reduce the
69 level of supplementation in the diet (van Heugten et al., 2003; Hill et al., 2014).

70 Although several studies have investigated the effect of zinc supplementation on growth
71 performance and carcass characteristics of growing-finishing pigs (Cemin et al., 2019;
72 Villagómez-Estrada et al., 2020), little information is available on the quality of the meat.
73 Indeed, only a few aspects of the pork quality, such as proximate composition, drip loss, pH,
74 color and fatty acids, have been marginally investigated when the effect of zinc
75 supplementation was assessed in growing-finishing pigs (Rekiel et al., 2005; Bučko et al.,
76 2013; Holen et al., 2017). However, the wide spectrum of biological activities of zinc may have
77 impacts on several meat quality traits mediated by the growth-promoting activity and/or related

78 to more direct effect. For example, zinc may exert a role in maintaining the integrity of cell
79 membranes and may act as an antioxidant factor through different mechanisms (Prasad, 1998;
80 Sloup et al., 2017). In particular, zinc could enhance the endogenous antioxidant defenses by
81 acting on antioxidant enzymes and inducing the synthesis of the metallothionein proteins which
82 are able to bind pro-oxidant metals or scavenge free radicals, such as hydroxyl radicals and
83 singlet oxygen (Prasad and Kucuk, 2002; Sloup et al., 2017). Moreover, zinc is believed to
84 decrease the formation of hydroxyl radicals, due to the competition with iron and copper (which
85 catalyze the production of hydroxyl radicals) to bind to the cell membrane (Oteiza et al., 1996).

86 Due to these multiple biological roles of zinc, in the present study, we hypothesized
87 that zinc supplementation may have some positive effect on the muscle antioxidant status and,
88 in turn, an effect on color and oxidative stability of pork. Therefore, the objective of this study
89 was to evaluate the effect of two levels (45 mg/kg and 100 mg/kg) of zinc chelate of glycine
90 hydrate (ZnGly) on growth performance, carcass characteristics and meat quality in growing-
91 finishing pigs.

92

93 **2. Materials and methods**

94 *2.1. Animals and experimental design*

95 The experimental trial was carried out at the facilities of the University of Catania and
96 the experimental protocol was approved by the animal welfare committee (OPBA) of the
97 University of Catania (No. 286946). Animals were handled by specialized personnel.

98 Thirty castrated male pigs (crossbred PIC × Piétrain) were selected for their age ($125 \pm$
99 5 day) and bodyweight (61.0 ± 4.02 kg) from a large range of barrows in a local commercial
100 farm. The animals were transported to the university facilities and allocated in individual pens
101 fitted with metal trough and nipple water dispenser. All the pens were placed in the same room
102 on a concrete floor and the barrows were bedded with wheat straw. Pigs were randomly

103 assigned to three dietary treatments (10 pigs/treatment) and fed *ad libitum* with a basal diet
104 supplemented with 0 mg/kg (control group, CON), 45 mg/kg (Zn45 treatment) or 100 mg/kg
105 (Zn100 treatment) of zinc chelate of glycine hydrate (ZnGly; Pancosma S.A., Rolle,
106 Switzerland). The ingredients and chemical composition of basal diet are shown in Table 1.
107 All the ingredients were ground, mixed thoroughly and pelleted. To ensure homogeneous
108 distribution, the zinc additive was pre-mixed with sepiolite and calcium carbonate (50:50 w:w)
109 and the obtained blend was then incorporated into the basal diet before pelleting in the ratio of
110 1 kg/1,000 kg. The carrier (i.e., sepiolite and calcium carbonate) without the addition of zinc
111 additive was included in the control group at the same dosage. Barrows were allowed free
112 access to feed and water during the entire experimental period. Offered concentrates and orts
113 were recorded for each pig in order to determinate the feed consumption. Individual
114 bodyweight was measured fortnightly during the trial. One pig from the Zn45 group died a few
115 days before the end of feeding trial for reasons unrelated to the experiment.

116

117 2.2. Slaughter procedure and samplings

118 After 56 days of feeding trial, all the pigs were weighed and conducted to a commercial
119 abattoir (transport duration of approximately 30 min) where they were slaughtered on the same
120 day by electric stunning and exsanguination according to the European Union Regulation
121 (council regulation (EC) No. 1099/2009). Liver samples were immediately collected after the
122 evisceration, vacuum packaged and frozen at -80 °C pending micromineral analysis. Hot
123 carcass weight (HCW) was recorded within 20 min from slaughter. Muscle pH was measured
124 at 45 min post-mortem directly in the *longissimus thoracis and lumborum* (LTL) muscle using
125 a pH-meter outfitted with a penetrating electrode (Orion 9106; Orion Research Incorporated,
126 Boston, MA). After 24 h of storage at 4 °C, carcasses were weighed to determinate the cold
127 carcass weight (CCW) and muscle pH was recorded as above. Then, a portion of LTL muscle

128 (between the 13th thoracic vertebra and the 3rd lumbar vertebra), together with backfat and rind
129 stuck on it, was excised from each right-side carcass, vacuum packaged and transported
130 refrigerated to the university laboratories. On the same day, muscle samples were divided into
131 three portions: *i*) an aliquot was stored vacuum-packed at -80 °C for micromineral and fatty
132 acid determination; *ii*) a second aliquot was chopped into small pieces, immediately frozen in
133 liquid nitrogen and stored at -80 °C for analyses of hydrophilic antioxidant capacity, fat-soluble
134 vitamins, cholesterol and antioxidant enzymes and *iii*) the third aliquot was aged vacuum-
135 packed at 4 °C for 24 h and then used for oxidative stability measurements.

136

137 2.3. Feed analyses

138 Representative feed samples were collected three times over the feeding period,
139 vacuum-packaged and stored at -20 °C pending analyses. Feed samples were ground in a
140 hammer mill fitted with 1-mm screen and analyzed for dry matter (DM), crude protein, crude
141 fat and ash according to the AOAC methods (1995). Neutral detergent fiber (NDF) was
142 determined according to Van Soest et al. (1991). The content of zinc, iron, copper and
143 manganese in the diet was determined by atomic absorption spectrometry as described in
144 Lombardo et al. (2017) with some adaptations. Approximately 1.3 g of oven-dried diet was
145 mixed with 2 drops of nitric acid and incinerated at 550 °C until a greyish-white ash was
146 obtained (48 h). The ashes were dissolved in 10 mL of 37% HCl and filtered through a paper
147 filter before analysis with the atomic absorption spectrometer.

148 Fatty acids of the basal diet were determined in a one-step extraction-transesterification
149 procedure using chloroform and methanolic sulfuric acid (Valenti et al., 2018). In brief, 1
150 mg/mL of internal standard (tridecanoic acid; C13:0) in hexane was placed into a glass tube
151 and the solvent was evaporated under nitrogen flow. An amount of 100 mg of ground feed
152 sample was weighted into the tube and 1.5 mL of chloroform and 2.5 mL of sulfuric acid (2%)

153 in methanol were added. Tubes were incubated for 2 h at 70 °C in a water bath. After cooling
154 to room temperature, 1.5 mL of chloroform and 2.5 mL of 6% K₂CO₃ were added. Samples
155 were centrifuged at 2,500 × g for 10 min at 4 °C and 1 mL of organic phase (bottom) was
156 collected and evaporated under nitrogen flow. The dried residue was dissolved with 1 mL of
157 hexane and analyzed through gas-chromatograph as later described for the analysis of
158 intramuscular fatty acids.

159 Tocopherols of the basal diet were extracted as reported by Rufino-Moya et al. (2020)
160 from 200 mg using 3 mL of methanol:acetone:petroleum ether (1:1:1, v:v:v) with BHT (0.01%,
161 w:v) and vortexing 1 min. The supernatant was collected after centrifugation at 1,000 × g for
162 5 min, repeating the extraction for a total of 3 times. The collected solvent was evaporated
163 under nitrogen flow and the residue was dissolved in 1 mL of methanol. Sample was filtered
164 by 0.22 µm PTFE filter and placed into a 2-mL vial. Tocopherols were determined through
165 ultra-high performance liquid chromatograph (UHPLC) as later detailed for the analysis of
166 muscle tocopherols.

167

168 2.4. Micromineral and fatty acid analyses

169 Micromineral content in liver and muscle samples were analyzed as described above
170 for the diet, with the only change being that 8 g of tissue was oven-dried and 8 drops of nitric
171 acid were used before ashing.

172 Intramuscular fat was extracted from 10 g of muscle using a mixture 2:1 (v:v) of
173 chloroform and methanol. Then, fatty acids were converted to fatty acid methyl esters (FAME)
174 by a base catalyzed transesterification using sodium methoxide in methanol. Methyl
175 nonadecanoate (C19:0) was used as internal standard. Fatty acids were separated through a gas-
176 chromatograph (model TRACE GC; Thermo Finnigan, Milan, Italy) in a 100-m high-polar
177 fused silica capillary column (25 mm i.d., 0.25-µm film thickness; SP. 24056; Supelco Inc.,

178 Bellefonte, PA) and identified by a flame ionization detector (FID). Gas-chromatography
179 conditions and identification of fatty acids was performed as reported in Natalello et al. (2019).
180 Atherogenicity (AI) and thrombogenicity (TI) indexes were calculated using the formulas
181 developed by Ulbricht and Southgate (1991), while the hypocholesterolemic to
182 Hypercholesterolemic ratio (h/H) was computed as reported in Fernández et al (2007), with
183 some minor changes as follows: $h/H = [(\text{sum of C18:1 } c9, \text{ C18:1 } c11, \text{ C18:2 } c9 \text{ } c12, \text{ C20:1 } c11,$
184 $\text{C18:3 } c9 \text{ } c12 \text{ } c15, \text{ C20:2 } c11 \text{ } c14, \text{ C20:3 } n-6, \text{ C20:3 } n-3, \text{ C20:4 } n-6, \text{ C22:4 } n-6, \text{ C22:5 } n-$
185 $3, \text{ C22:6 } n-3)/(\text{sum of C14:0 and C16:0})]$.

186

187 2.5. *Hydrophilic antioxidant capacity assays*

188 The antioxidant capacity in aqueous muscle extract was estimated with three different
189 assays in order to evaluate the radical scavenging activity and the reducing capacity as reported
190 in Luciano et al. (2017). In brief, muscle samples were deprived of visible fat and finely minced
191 using a knife. One gram of minced muscle was homogenized in 10 mL of distilled water for 1
192 min at 9,000 rpm (Diax 900, Heidolph Elektro GmbH & Co. KG, Kelheim, Germany). The
193 tubes containing samples were constantly kept in a water/ice bath during the homogenization.
194 Samples were centrifuged at $2,500 \times g$ for 20 min at 4°C and supernatant was filtered using
195 Whatman No. 541 filter paper. Filtrated samples were then aliquoted into 3 subsamples, each
196 for one of the following analyses, and stored at -80 °C.

197 The radical scavenging capacity was measured with the Trolox equivalent antioxidant
198 capacity (TEAC) assay, performed as described in Aouadi et al. (2014) with some
199 modifications. The ABTS⁺ solution was prepared by allowing an equal volume of 14 mM
200 aqueous ABTS [2,2-azinobis-(3-ethylbenzothiazoline 6-sulphonate)] and 4.9 mM potassium
201 persulphate to react for 16 h at room temperature. Twenty µL of filtered sample was mixed
202 with a 2 mL volume of ABTS⁺ solution and the absorbance at 734 nm was recorded after 60

203 min of incubation at 30 °C. A blank sample was prepared using 20 µL of distilled water instead
204 of the muscle sample in order to account for the spontaneous discoloration. The calibration
205 curve was made dissolving the Trolox standard (238813; Merck Life Science S.r.l., Milano,
206 Italy) in 4.9 mM potassium persulphate to get four points at concentrations ranging from 100
207 to 400 µg/mL.

208 Ferric reducing antioxidant power (FRAP) assay was performed as a measurement of
209 the reducing capacity, using the method developed by Benzie and Strain (1996) with minor
210 adaptations. In short, 50 µL of filtered muscle sample were mixed with 150 µL of distilled
211 water and 1.5 mL of a solution 10:1:1 (v:v:v) of 300 mM acetate buffer (pH 3.6), 10 mmol
212 TPTZ solution (2,4,6-tripyridyl-s-triazine in 40 mM HCl) and 20 mM aqueous ferric chloride.
213 The mixture was incubated at 37 °C for 60 min and the absorbance was then read at 593 nm as
214 well as a reagent blank. Aqueous ferrous sulfate heptahydrate ($\text{FeSO}_4 \times 7\text{H}_2\text{O}$) was used to
215 build a six-point calibration curve at concentrations ranging from 28 to 280 µg/mL.

216 Folin-Ciocalteu assay was performed as a further measurement of the reducing
217 capacity, using the method described by Makkar et al (1993). In brief, 0.5 mL of 1N Folin-
218 Ciocalteu reagent was mixed with 1 mL of filtered muscle sample previously diluted 1:2 with
219 distilled water. Then, 2.5 mL of sodium carbonate (20% w:v) was added to the mixture. After
220 40 min of incubation at room temperature in the dark, the solution was centrifuged at $2,500 \times$
221 g for 10 min at 4°C and the absorbance was read at 725 nm. A six-point calibration curve was
222 made using aqueous tannic acid at concentrations ranging from 10 to 100 µg/mL.

223

224 2.6. *Fat-soluble vitamins and cholesterol*

225 Tocopherols, retinol and cholesterol were determined using the method developed by
226 Bertolín et al. (2018) with minor adjustments as follows. Lyophilized muscle samples (500 mg)
227 were placed in 15-mL plastic centrifuge tube together with 200 mg of L-ascorbic acid and 7.5

228 mL of saponification solution (10% w:v potassium hydroxide in 1:1 v:v ethanol:water
229 solution). Samples were saponified overnight at 22 °C in an incubator shaker (KS 4000 i
230 control; IKA®-Werke GmbH & Co. KG, Staufen, Germany) set at 250 rpm and protected from
231 light. The following day, 5 mL of 9:1 (v:v) hexane:ethyl acetate containing 25 µg/mL of BHT
232 was added and the tubes were vortexed for 60 s and then centrifuged at 2,000 × g for 5 min at
233 10 °C (Centrifuge 5810 R; Eppendorf s.r.l., Milano, Italy). The upper phase was collected and
234 placed in a glass tube. This extraction procedure was repeated twice. The collected organic
235 phases were evaporated under nitrogen flow at 40 °C using a sample concentrator and a block
236 heater (SBHCONC/1 and SBH130D/3 Stuart®; Cole-Parmer, Stone, United Kingdom). The
237 dry residues were dissolved in 1 mL of methanol (HPLC grade), warming tubes at 40 °C and
238 vortexing thoroughly. The samples were then filtered by 0.2 µm-13 mm PTFE syringe filters
239 and placed into 2-mL amber vials. The analytes were separated and quantified through a Nexera
240 UHPLC (Shimadzu Corporation, Kyoto, Japan) equipped with a C18 phase column (Zorbax
241 ODS; 25 cm × 4.6 mm, 5 µm; Supelco, Bellefonte, PA). A sample volume of 10 µL was
242 injected into the system and the isocratic mobile phase was methanol at the flow rate of 1.3
243 mL/min. The temperature of the samples (SIL-40C XS Autosampler, Shimadzu) and column
244 oven (CTO-40C, Shimadzu) were set at 25 °C and 40 °C, respectively. Retinol and cholesterol
245 were detected by a photodiode array detector (PDA; SPD-M40, Shimadzu) at the absorbance
246 of 325 nm and 220 nm, respectively. Tocopherols were detected using a spectrofluorometric
247 detector (RF-20AXS, Shimadzu) at 295 nm excitation wavelength and 330 nm emission
248 wavelength. The analytes were identified by comparison of retention times with those of pure
249 standards (Merck Life Science S.r.l., Milano, Italy). The quantification was achieved by
250 external calibration curves made with pure standards.

251

252 *2.7. Antioxidant enzymes*

253 Frozen muscle samples were deprived of visible fat and finely minced with a knife.
254 Five grams of minced muscle was placed in a 50-mL centrifuge tube with 10 mL of ice-cold
255 50 mM phosphate buffer [pH 7.0; disodium phosphate dodecahydrate ($\text{Na}_2\text{HPO}_4 \times 12\text{H}_2\text{O}$) and
256 KH_2PO_4] and homogenized at 13,000 rpm for 60 s (30 s + pause + 30 s) using a T18 digital
257 Ultra-Turrax® (IKA®-Werke GmbH & Co. KG, Staufen, Germany). During the
258 homogenization, the tubes containing samples were always kept in a water/ice bath to prevent
259 enzymatic reactions and oxidations. Tubes were then centrifuged at $2,800 \times g$ for 20 min at
260 4°C. Supernatants (1.8 mL) were collected in 2-mL microcentrifuge tubes and further
261 centrifuged at $10,000 \times g$ for 10 min at 4°C. The muscle extract thus obtained was aliquoted
262 (0.5 mL) in 3 microtubes and immediately stored at -80 °C until enzymatic analyses.

263 The catalase (CAT) activity was determined following the method of Jin et al. (2013)
264 with some modifications. An aliquot of muscle extract was thawed at room temperature for a
265 few min, vortexed and 30 μL was placed in a UV cuvette. A volume of 1.74 mL of H_2O_2
266 solution (11 mM H_2O_2 in 50 mM phosphate buffer) was added and the cuvette was immediately
267 capped and inverted 4 times. The changes in absorbance were monitored at 240 nm in kinetics
268 mode (model UV-1601; Shimadzu corporation, Kyoto, Japan) over 3 min with reading interval
269 of one second. A blank was prepared with 30 μL of 50 mM phosphate buffer and 1.74 mL of
270 H_2O_2 solution. The molar extinction coefficient of H_2O_2 ($39.5 \text{ M}^{-1} \text{ cm}^{-1}$) was used to calculate
271 the catalase activity and results were expressed as U/g of muscle. One unit (U) of catalase
272 activity was defined as the amount of muscle extract needed to decompose 1 μmol of H_2O_2 per
273 min.

274 Glutathione peroxidase (GSH-Px) activity was measured according to the procedure
275 developed by Flohé and Gunzler (1984) with some adaptations. Briefly, 500 μL of the assay
276 medium (100 mM potassium phosphate buffer (pH 7.0), 1 mM EDTA, 2 mM NaN_3), 100 μL
277 of muscle extract (previously diluted 4-fold in 50 mM phosphate buffer), 100 μL of glutathione

278 reductase (2.4 U/mL), 100 μ L of 10 mM L-glutathione and 100 μ L of NADPH solution (1.5
279 mM NADPH in 0.1% NaHCO₃) were directly placed into a UV cuvette and mixed by inverting
280 the cuvette. After 5 min of incubation at room temperature, the overall reaction was started by
281 adding 100 μ L of 1.5 mM H₂O₂ and the absorbance at 340 nm was monitored in kinetic mode
282 every second for 5 min. A blank was prepared with 50 mM phosphate buffer instead of muscle
283 extract. The extinction coefficient of 6.22 mM⁻¹ cm⁻¹ was used to calculate NADPH
284 concentration. One U of GSH-Px activity was defined as the amount of muscle extract needed
285 to oxidize 1 μ mol of NADPH per min.

286 Superoxide dismutase (SOD) activity was evaluated according to the method described
287 in Gatellier et al. (2004). In short, 760 μ L of 50 mM Tris-HCl buffer (8.2 pH) were located into
288 a UV cuvette. Then, 20 μ L of muscle extract and 20 μ L of 10 mM pyrogallol were added and
289 the cuvette was inverted. The absorbance at 340 nm was monitored in kinetic mode for 300
290 seconds with reading interval of 1 second. One U of SOD activity was defined as the amount
291 of muscle extract needed to inhibit the pyrogallol autoxidation by 50% through comparison
292 with a blank (20 μ L of 50 mM phosphate buffer in place of muscle extract). Results were
293 expressed as U of enzyme activity per g of muscle.

294

295 2.8. *Oxidative stability of backfat and meat*

296 Oxidative stability was evaluated in fresh and cooked meat over aerobic storage as
297 reported by Valenti et al. (2019). In brief, three 2-cm-thick slices were prepared from each LTL
298 muscle sample, placed in polystyrene trays and over-wrapped with 2-layers of domestic cling
299 film. Slices were stored in the dark at 4 °C for 0 (after 2 h of blooming), 3 and 6 days. After
300 each storage time, one of the 3 slices was used to determined color and then frozen pending
301 lipid oxidation analysis. Regarding the cooked meat, other 3 slices of muscle were prepared,
302 vacuum packaged and directly cooked at 70 °C for 30 min. Then, one slice was immediately

303 frozen pending lipid oxidation analysis (day 0), whereas the other two slices were stored in
304 aerobic conditions, as for the raw meat, for 2 and 4 days. After each storage time, one of the 2
305 slices was frozen pending lipid oxidation analysis. Furthermore, as described by Biondi et al
306 (2020), three slices of backfat were cut to a thickness of 2 cm and stored in the same conditions
307 as for the raw meat for 0, 3 and 6 days and used to measure the color descriptors.

308 Color stability of raw meat and backfat was measured by a Minolta CM 2022
309 spectrophotometer (d/8° geometry; Minolta Co. Ltd. Osaka, Japan) set to operate in the
310 specular components excluded (SCE) mode and to measure with the illuminant A and 10°
311 standard observer. Three measurements were taken on the meat and backfat surface on non-
312 overlapping areas and the mean value was calculated. The color descriptors L* (lightness), a*
313 (redness), b* (yellowness), C (saturation) and h_{ab} (hue angle) were measured in the CIE L* a*
314 b* color space. Total color change (ΔE) between 3 or 6 days of storage and the day 0 was
315 calculated as $\Delta E = [(\Delta L^*)^2 + (\Delta a^*)^2 + (\Delta b^*)^2]^{1/2}$. Where ΔL^* , Δa^* and Δb^* are the differences
316 in L*, a* and b*, respectively, between day 0 and day 3 or day 6. Furthermore, the reflectance
317 spectra from 400 to 700 nm wavelength were also recorded in order to calculate the 630/580
318 nm ratio as indicator of myoglobin oxidation.

319 Lipid oxidation was determined in both raw and cooked slices by measuring the 2-
320 thiobarbituric acid reactive substances (TBARS) at the end of each storage time, as reported in
321 Natalello et al. (2020) with some modifications. In brief, 5 g of frozen meat was firstly minced
322 with a knife and then homogenized with 15 mL of 7.5% (w/v) trichloroacetic acid (TCA). The
323 homogenate was filtered through filter paper (Whatman No. 1) and 4 mL of clear filtrate was
324 mixed with 4 mL of 0.02 M aqueous thiobarbituric acid (TBA). After incubation at 80 °C for
325 90 min, the absorbance was read at 532 nm (UV-1601; Shimadzu Corporation, Milan, Italy).
326 A calibration curve was prepared with TEP (1,1,3,3,-tetraethoxypropane) in distilled water at

327 concentrations ranging from 5 to 65 nmoles/4 mL and results were expressed as mg of
328 malondialdehyde (MDA) per kg of meat.

329 Lipid oxidation was also assessed in meat homogenates incubated with Fe³⁺ and
330 ascorbate (Fe/Asc) as catalyst of oxidative reactions. The analytic procedures used were
331 adapted from Luciano et al. (2019). Briefly, 10 g of meat was homogenized with 40 mL of 2-
332 (N-morpholino)ethanesulfonic acid (MES) buffer (pH 5.7). The homogenate was equilibrated
333 at 4 °C and then 4 mL was collected for measuring the initial extent of lipid oxidation (0 min).
334 Subsequently, 40 µL of an equimolar solution of ferric chloride hexahydrate and L-sodium
335 ascorbate (45 mM) was added and samples were incubated at room temperature. After 30 and
336 60 min of incubation, 4 mL of sample was collected, mixed with 4 mL of 15% TCA and filtered
337 through Whatman No. 541 paper. The filtered sample (2 mL) was reacted with 2 mL of 0.02M
338 aqueous TBA at 80 °C for 90 min. The absorbance was read at 532 and a calibration curve was
339 prepared using standard solutions of TEP (2.5 to 32.5 nmoles/2 mL).

340

341 2.9. Statistical analysis

342 All the data were analyzed with the SPSS software (SPSS For Analytics, version 26;
343 IBM corporation, Armonk, NY). The single animal was used as statistical unit. The effect of
344 dietary additive on animal performance, carcass traits, micromineral content of liver and
345 muscle, intramuscular fat, meat fatty acid profile, overall antioxidant capacity, enzymatic and
346 non-enzymatic antioxidants in muscle was statistically analyzed using one-way ANOVA. Data
347 of color and lipid stability were analyzed using a mixed model for repeated measures. The
348 terms in the model were: dietary treatment (D), time of storage or incubation (T) and their
349 interaction (D × T) as fixed factors and each animal as random factor. When P-value was ≤ of
350 0.05, multiple comparisons were performed using the Tukey post hoc test.

351

352 **3. Results**

353 *3.1. Animal performance and carcass traits*

354 The animal performance and carcass traits of pigs fed diets supplemented with ZnGly
355 are shown in Table 2. The highest ZnGly supplementation (i.e., Zn100) reduced the average
356 daily gain (ADG; $P = 0.031$) or tended to reduce the daily intake ($P = 0.069$) and final body
357 weight ($P = 0.063$) compared to the CON treatment, while Zn45 group did not statistically
358 differ from the other two treatments. Conversely, the feed conversion ratio was not affected by
359 dietary Zn ($P > 0.05$). Hot carcass weight was greater in the control treatment compared only
360 to Zn100 treatment ($P = 0.042$), while the cold carcass weight was statistically comparable
361 between treatments ($P > 0.05$). Both levels of ZnGly supplementation reduced ($P < 0.001$)
362 carcass weight loss occurring within 24 h of chilling storage and tended to increase ($P = 0.094$)
363 the cold carcass yield as compared with CON treatment. Cooking loss, pH and color parameters
364 of muscle as well as color parameters of backfat were not influenced by the ZnGly
365 supplementation ($P > 0.05$).

366

367 *3.2. Liver and muscle microminerals and intramuscular fatty acids and cholesterol*

368 The contents of copper, iron, manganese and zinc in liver and muscle are reported in
369 Table 3. No differences were observed for all microminerals ($P > 0.05$) with the sole exception
370 of the iron concentration in the liver, which was higher in the Zn45 group than in CON and
371 Zn100 groups ($P = 0.006$).

372 Table 4 shows the effect of ZnGly supplementation on intramuscular fat content, fatty
373 acid (FA) profile and cholesterol of muscle. Dietary treatment did not affect the intramuscular
374 fat content and the individual fatty acids ($P > 0.05$), except for C22:5 *n*-3 that was lower in
375 both Zn45 and Zn100 compared to CON treatment ($P = 0.001$). In turn, the principal FA classes
376 (i.e., saturated, monounsaturated and polyunsaturated FA) and health indices (i.e.,

377 atherogenicity and thrombogenicity indices and hypocholesterolemic to hypercholesterolemic
378 ratio) were not altered by ZnGly supplementation ($P > 0.05$), as well as highly peroxidizable
379 polyunsaturated FA (HP-PUFA) with at least three double bonds and the peroxidability index.
380 Cholesterol concentration in muscle tended to be greater in CON group than Zn45 and Zn100
381 ($P = 0.06$).

382

383 *3.3. Antioxidant status of muscle*

384 As shown in Table 5, the highest dose of ZnGly supplementation (i.e., Zn100) increased
385 the catalase activity compared to CON and Zn45 treatments ($P = 0.003$), while Zn45 group
386 showed a lower SOD activity than CON, with an intermediate value for Zn100 treatment ($P =$
387 0.007). No differences between the three treatments were observed for the fat-soluble vitamins
388 (i.e., α -tocopherol and retinol) and the activity of GSH-Px ($P > 0.05$). Similarly, hydrophilic
389 antioxidant capacity, measured with TEAC, FRAP and Folin-Ciocalteu assays, were
390 statistically comparable for all the treatments ($P > 0.05$).

391

392 *3.4. Color stability of backfat and meat*

393 The effects of the mineral supplementation and the time of storage on color stability of
394 backfat and meat are reported in Table 6. Neither backfat color descriptors nor color descriptors
395 of raw meat were influenced by ZnGly supplementation ($P > 0.05$). All the color parameters
396 measured in both backfat and meat slices were affected by the time of storage, except for h_{ab}
397 in the backfat ($P = 0.174$). Specifically, L^* (lightness) values of backfat increased after 3 days
398 of storage and stabilized thereafter between day 3 and day 6. Instead, a^* (redness), b^*
399 (yellowness) and C^* (saturation) values were lower in the backfat after 6 days of storage
400 compared to day 0 and day 3 ($P < 0.001$). Consequently, ΔE was higher in backfat after 6 days
401 of storage than compared to 3-day storage time ($P = 0.001$). Concerning the meat color, L^*

402 values increased, while a*, b*, C* and 630/580 ratio decreased along the 6 days of refrigerated
403 storage ($P < 0.001$). The meat slices stored 3 days had higher values of h_{ab} compared to day 0
404 and day 6 ($P < 0.001$). Similar to backfat, ΔE values of meat were greater after 6 days of chilling
405 storage than after 3 days ($P < 0.001$). No significant interaction was found between the dietary
406 treatment and the storage time for all the color parameters ($P > 0.05$).

407

408 3.5. Meat lipid stability

409 There were no significant interactions between ZnGly supplementation and time of
410 storage/incubation for lipid stability analysis ($P > 0.05$). Therefore, figure 1 shows the main
411 effects of (a) the ZnGly supplementation and (b) the time of storage/incubation on meat lipid
412 oxidation as evaluated by TBARS assay. The dietary treatment did not significantly affect the
413 lipid oxidation in raw meat ($P = 0.250$) and homogenized meat incubated with pro-oxidant
414 catalysts ($P = 0.531$). Whereas the lower dose of ZnGly supplementation (i.e., Zn45) reduced
415 the lipid oxidation in cooked meat ($P = 0.037$; Fig. 1a). As expected, lipid oxidation increased
416 ($P < 0.001$) over the time of refrigerated storage or incubation (Fig. 1b), for the raw and cooked
417 meat or for the homogenized meat, respectively. In raw meat, TBARS values at day 0 and day
418 3 were similar and lower compared to day 6. In homogenized and cooked meat, the lowest and
419 the highest TBARS values were observed at the first and last time point, respectively, and
420 intermediate values for the center time point.

421

422 4. Discussion

423 4.1. Animal performance

424 Although high levels of zinc (Zn) supplementation have long been reported to stimulate
425 voluntary feed intake and weight gain in young pigs (Hahn and Baker, 1993; Case and Carlson,
426 2002; Barszcz et al., 2019), this effect seems less clear for growing-finishing pigs. Indeed, early

427 studies demonstrated that feeding growing pigs with increasing levels of Zn oxide (from 0 to
428 200 mg/kg) had no effect on growth performance (Poulsen and Larsen, 1995; Larsen and
429 Poulsen, 1996). Similarly, D'Souza et al. (2012) observed no improvement in feed intake and
430 daily gain when growing pigs were supplemented with 250 mg/kg of Zn. The reason for this
431 discrepancy between young and heavy pigs might be due to the different physiological period,
432 as the growing phase is a more stable and regular physiological period than post-weaning one,
433 being that weaning causes stress, reduces feed intake and increases susceptibility to infections
434 (Blaabjerg and Poulsen, 2017). As a result, growing-finishing pigs are less demanding than
435 post-weaning piglet in terms of Zn requirement which decreases from 100 to 50 mg/kg for
436 young to finishing pigs (NRC, 2012).

437 In the present study, the Zn supplementation did not improve growth performance. On
438 the contrary, the highest Zn level (i.e., Zn100) reduced the average daily gain (ADG) and
439 tended to reduce feed intake and final body weight compared to control treatment. A possible
440 explanation for our results may be due to the fact that pigs from control group tended to increase
441 feed intake to compensate for the lack of Zn supplementation and thus to meet their daily Zn
442 requirement. In turn, this may have led to an increase of ADG. This hypothesized compensatory
443 effect may not have been observed in the studies of Poulsen and Larsen (Poulsen and Larsen,
444 1995; Larsen and Poulsen, 1996) because growing pigs were fed restrictively (i.e.,
445 approximately 10% below the expected maximum intake) and thus not allowing them to
446 increase their intake to meet Zn requirements. On the other hand, some studies provided
447 evidences that omitting the supply of microminerals, including Zn, in diets of pigs had no
448 negative impact on animal growth performance, even when growing pigs were fed *ad libitum*
449 (Shelton et al., 2004; Gowanlock et al., 2013). Nevertheless, it should be stressed that, unlike
450 young pigs, only a few studies have investigated the Zn supplementation in heavy pigs. Hence,
451 further investigations are required to fully explicate the effect of Zn supplementation in heavy

452 pigs, especially in commercial farm conditions, where there are likely to be several challenges
453 for pigs.

454

455 *4.2. Carcass traits and meat quality*

456 Interestingly, cold carcass weight (CCW) was comparable between the three
457 treatments, despite the fact that the hot carcass weight (HCW) was reduced by the highest level
458 of Zn supplementation. The difference for the HCW could be a direct consequence of the
459 observed trend for the final body weight, which were slightly lowered by the highest Zn level
460 (i.e., Zn100). While the lack of effect on CCW was clearly due to the lower carcass weight loss
461 for the Zn treatments that occurred during the 24 hours post-mortem of refrigerated storage.
462 Indeed, chilling carcass loss was reduced by more than one percentage point by dietary Zn
463 additive and, in turn, cold carcass yield tended to be higher in Zn pigs.

464 Moisture loss during refrigerated storage reduces the overall weight of the carcass,
465 which contributes to economic loss by reducing the weight of the salable product. In addition,
466 water loss may have a huge effect on meat quality as it is closely related to color, taste,
467 tenderness and juiciness (Warner, 2017). The rate and extent of pH fall are the main factors
468 affecting the ability of the muscle to retain fluids; indeed, a rapid decline of pH after slaughter
469 or a high ultimate pH usually lead to defects in raw meat, such as PSE (pale, soft, and
470 exudative) or DFD (dark, firm, and dry) meats (Warner, 2017). In the present study, muscle
471 pH values measured at 45 minutes or 24 hours post-mortem were not affected by Zn
472 supplementation and can be considered as normal values for pork (Matarneh et al., 2017).
473 Therefore, it would appear that the reduction of chilling carcass loss should be due to other
474 mechanisms not related to muscle pH.

475 The effect of Zn supplementation on pork quality has been rarely investigated.
476 However, the few studies conducted so far have reported no effect of Zn on pork water loss

477 (Bučko et al., 2013; Gowanlock et al., 2013; Holen et al., 2018). While, there are a number of
478 poultry studies that have observed a positive effect of Zn supplementation on meat moisture
479 retention. For instance, Liu et al. (2011) demonstrated that different Zn levels in broiler diets
480 were able to reduce drip loss of both breast and thigh muscles regardless the Zn source.
481 Likewise, cooking loss of breast and thigh was linearly reduced by increasing inclusions of Zn
482 bearing palygorskite in a broiler diet, but drip loss was not affected (Yang et al., 2016).
483 Moreover, meat from quails fed diets supplemented with 40 and 80 mg/kg of Zn showed lower
484 drip and cooking losses (Rouhalamini et al., 2014). Similar to our findings, in a recent study
485 (Chang et al., 2021) a reduction of breast water loss was reported after 24 h of refrigerated
486 storage and no difference for pH values when ducks received a basal diet supplemented with
487 increasing levels of zinc glycine chelate. To explain these findings, most of the above studies
488 on poultry meat (Liu et al., 2011; Yang et al., 2016; Chang et al., 2021) speculated that Zn
489 supplementation may have enhanced the antioxidant capacity of muscle and consequently
490 reduced lipid oxidation, which has a negative influence on the integrity of cell membranes and
491 thus on the ability to retain intracellular fluids. In the present study, we might rule out this
492 hypothesis as no significant effects on the lipid oxidation of raw meat were observed. Indeed,
493 as better explained in the following paragraphs, lipid oxidation in raw meat developed to a
494 limited extent even after 6 days of storage in aerobic conditions and was not affected by the
495 dietary treatment. Therefore, it seems implausible that during the first 24 hours after slaughter
496 there was such a strong lipid oxidation able to significantly influence the chilling carcass loss.
497 Further studies are needed to elucidate the biological mechanisms involved in reducing meat
498 water loss when animals are supplemented with Zn additives.

499 In the current study, the Zn supplementation did not influence neither color of muscle
500 and backfat nor meat fatty acid profile. To the best of our knowledge, there are hardly any
501 studies in which the effect of dietary Zn on pork color and fatty acid have been evaluated.

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502 Bučko et al. (2013) observed that fresh meat, from pigs supplemented with 66 mg/kg of organic
503 Zn, showed lower values of L* (lightness), b* (yellowness) and polyunsaturated FA (PUFA).
504 Whereas, Rekiel et al. (2005) did not find any significant effect of 0.05% Zn oxide on the main
505 FA classes (i.e., saturated, monounsaturated and polyunsaturated FA) and Holen et al. (2018)
506 noticed no meaningful effect on color characteristics of pork when pigs were supplemented
507 with organic and inorganic Zn under crowded housing conditions.

508 In our experiment, Zn supplementation tended to reduce the cholesterol content of pork,
509 which was in agreement with Dukare et al. (2021) who reported decrease of thigh and breast
510 cholesterol content in broiler meat by supplementing different sources of Zn up to 80 mg/kg
511 level. However, differently from our findings, these authors reported also a reduction of fat
512 percentages of thigh and breast by dietary Zn that might explain the lower content of
513 cholesterol.

514 It is well known that liver is the main storage site of Zn (Hill et al., 2014) and thus it is
515 mostly responsible for Zn homeostatic regulation. Indeed, several studies have found higher
516 Zn content in liver compared to other organs (van Heugten et al., 2003; Untea et al., 2017). In
517 line with that, we observed here a greater concentration of Zn in the liver than muscle tissue.
518 A previous study has shown that liver Zn content of nursery pigs increased linearly as the level
519 of Zn supplementation increased from 25 to 100 mg/kg (Hill et al., 2014); while Case and
520 Carlson (2002) detected higher Zn content in the liver of pigs supplemented with an excessively
521 high level (i.e., 3,000 mg/kg) of Zn but not in pigs supplemented with lower levels (i.e., 150
522 and 500 mg/kg). In the present study, no differences in hepatic Zn content were observed when
523 pig diets were supplemented with 45 and 100 mg/kg of Zn. From our results it would appear
524 that the supply of Zn at low levels has no meaningful effects on Zn accumulation in liver of
525 growing-finishing pigs. It should be recalled that Zn can compete with other microminerals,
526 such as iron (Fe) and copper (Cu), for absorption and high levels of dietary Zn could lead to

527 deficiencies of these elements (Jensen-Waern et al., 1998). An early study of Cox and Hale
528 (1962) demonstrated that Fe concentration in pig liver was lowered by Zn supplementation at
529 the level of 4,000 mg/kg but not with 2,000 mg/kg. Whereas, Carlson et al. (1999) observed no
530 difference in Fe concentration of liver from pigs supplemented with 3,000 mg/kg. Interestingly,
531 in our study, a significant higher concentration of Fe was found in the liver from Zn45
532 treatment. Hence, our finding seems to suggest that low levels of zinc glycinate (ZnGly) have
533 not limited the absorption and the accumulation of micromineral, but rather Fe content in the
534 liver was enhanced by 45 mg/kg of ZnGly.

535 In the present study, micromineral content in the muscle was not affected by Zn
536 treatments. Our results indicated that pork from pigs given ZnGly up to 100 mg/kg should not
537 raise any concerns for consumption as muscular Zn concentration was comparable with control
538 treatment. This was in accordance with a previous report (Jensen-Waern et al., 1998), in which
539 the supplementation of Zn did not influence microelement contents in pig muscle despite the
540 high dosage used (i.e., 2500 mg/kg).

541

542 *4.3. Oxidative stability*

543 Among the endogenous defense systems against oxidative processes in muscle, SOD is
544 one of the most important enzymes and is responsible for the dismutation of superoxide anions
545 into hydrogen peroxide (Bekhit et al., 2013). In mammals, SOD enzyme contains different
546 cofactors and exists in three different forms: CuZn-SOD, present in the cytoplasm and
547 containing Cu and Zn; Mn-SOD, present exclusively in the mitochondrial spaces and
548 containing Mn; EC-SOD, containing also Zn and Cu but operates in the extracellular space
549 (Zelko et al., 2002). In pigs as well as in other mammal species, CuZn-SOD shows much
550 greater activity than other forms (Marklund, 1984). Although some studies have observed an
551 increase in SOD activity in the serum or tissues of animals supplemented with increasing levels

552 of Zn (Bun et al., 2011; Dukare et al., 2021), it is believed that SOD activity is mainly regulated
553 by the availability of Cu rather than Zn (Harris, 1992; Carlson et., 1999). In the present study,
554 we measured the total activity of the three forms of SOD as whole and observed a lower total
555 SOD activity in the muscle of pigs supplemented with 45 mg/kg of ZnGly. This result might
556 be derived from some interaction of the Zn with the other cofactors of SOD (i.e., Cu and Mn),
557 since Zn – competing for adsorbent and for binding to cell membranes – can reduce the
558 availability of Cu and Mn. Similarly, Zn supplementation may also have some interactive effect
559 with Fe, which is a cofactor of catalase enzyme (Nicholls, 2012). Indeed, in the present
560 experiment, catalase activity was higher in the muscle of pigs supplemented with 100 mg/kg
561 of Zn. Consistently, Wen et al. (2011) showed that muscle from Pekin ducks supplemented
562 with 240 mg/kg of Zn had a higher catalase activity compared with 0, 15 and 30 mg/kg.
563 However, this hypothesis seems to contrast our result of micromineral contents of muscle,
564 which were unaffected by Zn supplementation.

565 On the other hand, an increase in enzyme activity is often interpreted as an adaptive
566 response to oxidative stress (Renerre et al., 1996). This belief is supported by the studies of
567 Bun et al. (2011) and Stukelj et al. (2013), which reported increased hepatic or serum SOD
568 activity in broilers challenged with a pathogen or pigs infected with a virus, respectively.
569 Therefore, our finding may indicate that the lowest Zn supplementation led to a better oxidative
570 condition status of animal and therefore a lower need to increase the response of the enzymatic
571 defenses. Nevertheless, it should be underlined that the pigs of the present study were not
572 subjected to any type of stress, since the animals were raised in ideal environmental and
573 nutritional conditions throughout the experiment as well as kept in individual pens, thus
574 avoiding any competitions problem between pigs. Moreover, it is worth mentioning that the
575 muscle from the present study had a remarkable concentration of α -tocopherol (approximately
576 3 μ g/g of muscle) regardless the dietary treatment. Hence, considering that α -tocopherol

577 (vitamin E) is a powerful antioxidant in muscle systems (Pettigrew and Esnaola, 2001; Bekhit
578 et al., 2013), its high content may have adequately protected the muscle against natural
579 oxidative processes and consequently limiting the enzymatic defense response.

580 In the current study, color stability of meat and backfat was not affected by Zn
581 supplementation. Whereas, the effect of the storage days on the color parameters was evident
582 as expected. Yet, it is important to emphasize that the color variations over time were
583 numerically small, indicating that the backfat and muscle of the present experiment were not
584 prone to oxidation even after 6 days of aerobic storage. This low propensity to oxidation was
585 consistent with the values of secondary products of lipid oxidation assessed by the TBARS
586 assay. Indeed, the TBARS values of raw meat did not statistically differ between meat at time
587 0 and after 3 days of storage, but increased only after 6 days of aerobic storage at 4 ° C and in
588 any case with values always lower than 0.1 mg MDA per kg of meat (Fig.1b). It should be
589 remembered that these values were far below the threshold at which consumers may detect off-
590 flavors in pork (Sheard et al., 2000). Even when the meat was subjected to a strong oxidative
591 stressor, such as incubation with pro-oxidant catalysts (Fe^{3+} and ascorbate), the observed values
592 were always below 0.9 mg MDA per kg of meat (Fig.1b). A plausible explanation for this
593 scarce propensity of pork to lipid oxidation could be due to the fairly high concentration of α -
594 tocopherol in the muscles of the present study regardless the dietary treatment, as already
595 mentioned above; indeed, vitamin E has been proven to be particularly effective in delaying
596 lipid oxidation and reducing secondary lipid oxidation products in pork as well reviewed by
597 Pettigrew and Esnaola (2001).

598 However, when pork slices were subjected to a much stronger oxidative challenge, such
599 as cooking, TBARS values from cooked pork slices increased to nearly 1.8 mg MDA/kg after
600 4 days of refrigerated aerobic storage (Fig.1b). Interestingly, cooked pork from pigs
601 supplemented with 45 mg/kg of ZnGly showed a higher resistance to lipid oxidation than

602 control treatment, with intermediate TBARS values for Zn100 treatment (Fig.1a). Thus, it may
603 be speculated that the effect of Zn on lipid oxidation was partially hidden by the strong
604 protection exerted by vitamin E when the pork was exposed to low or medium oxidized
605 stressors (i.e., refrigerated aerobic storage or incubation with pro-oxidants), while this effect
606 was evident in highly stressful conditions.

607 These results demonstrated that ZnGly may enhance the oxidative stability of pork
608 under strong oxidative challenge, even if the antioxidant action mechanism of Zn remains to
609 be elucidated. Indeed, according to our findings, the Zn supplementation would appear to have
610 no effect on the fat-soluble vitamins or on the hydrophilic antioxidant capacity – evaluated by
611 TEAC, FRAC and Folin-Ciocalteu assays – of the muscle. Moreover, the lack of response on
612 the SOD activity would seem more an indirect effect on the overall oxidative state of animal
613 than a direct effect since the lowest Zn level decreased the activity of this Zn-containing
614 enzyme. Although in the present study a series of analyzes was carried out to better study the
615 antioxidant effects of Zn supplementation on pork oxidative stability, further research is
616 essential to clarify the antioxidant effect of dietary Zn by also evaluating other aspects related
617 to Zn such as its role in the induction of metallothionein proteins and in the inhibition of
618 NADPH oxidase (Prasad and Kucuk, 2002). Furthermore, future studies should be designed to
619 evaluate the effect of Zn in oxidative challenge conditions such as diets deficient in vitamin E
620 or supplemented with polyunsaturated FA, or physiological stress conditions.

621

622 **5. Conclusions**

623 So far, little has been done to assess the effects of Zn supplementation on the
624 performance and quality of the carcass and meat for pigs in the growing-finishing stage. The
625 results obtained in this study suggest that supplementing the diet of growing-finishing pigs with
626 45 or 100 mg/kg of ZnGly may reduce chilling carcass loss and cholesterol content of meat.

627 Even though the highest level of Zn supplementation slightly lowered the average daily gain,
628 cold carcass weight did not statistically differ between treatments, indicating that ZnGly has
629 no detrimental effect on economic productivity for swine industries. Both levels of Zn
630 supplementation did not alter the micromineral content, fatty acid profile, fat-soluble vitamins
631 and hydrophilic antioxidant capacity of muscle. Muscle from pigs supplemented 45 mg/kg of
632 Zn showed a lower SOD activity compared to control muscle, while the highest level of Zn
633 increased the catalase activity. This result certainly needs further investigation to establish the
634 role of Zn in regulation of the activity of these enzymes.

635 Our results showed that the Zn supplementation level of 45 mg/kg improved the meat
636 oxidative stability when pork was subjected to strong oxidative challenges such as cooking,
637 while no effects were observed under aerobic storage of fresh meat or incubation of meat
638 homogenates with pro-oxidant catalysts. Therefore, in these experimental conditions, an
639 antioxidant effect of Zn supplementation may be not evident under milder oxidizing conditions
640 and may be hidden by the effective protection of vitamin E against lipid oxidation. Future
641 studies should be addressed to examine the Zn supplementation in different breeding and meat
642 storage conditions to better capture the antioxidant potential of Zn and its influence on the
643 shelf-life of pork.

644

645

646 **Conflict of interest**

647 The authors Hajer Khelil-Arfa, Mieke Zoon and Alexandra Blanchard are employees at
648 Pancosma S.A. and they played no role in collection, analysis and interpretation of data. All
649 authors declare no potential conflicts of interest.

650

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656

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855 **Table 1.**

856 Ingredients and chemical composition of the basal diet.

	Basal diet
<i>Ingredients, g/kg as fed basis</i>	
Maize	420
Barley	200
Wheat bran	152
Soybean meal (48% crude protein)	134
Fava bean	74
Vitamin premix ¹	10
Sodium carbonate	8
Amino acid premix ²	2
<i>Chemical composition, g/kg DM</i>	
DM, g/kg as fed	905
Crude Protein	170
Crude Fat	32.9
Neutral detergent fiber	165
Ash	41.7
<i>Fatty acids, g/kg DM</i>	

C16:0	2.52
C18:0	0.49
C18:1 <i>c</i> 9	3.50
C18:2 <i>c</i> 9 <i>c</i> 12	7.61
C18:3 <i>c</i> 9 <i>c</i> 12 <i>c</i> 15	0.58
<i>Tocopherols, mg/kg DM</i>	
α-Tocopherol	7.59
γ-Tocopherol	21.5
δ-Tocopherol	6.42
<i>Microminerals, mg/kg DM</i>	
Copper	16.8
Iron	94.6
Manganese	1.17
Zinc	22.3

857

858 ¹ One kg of premix contained: vitamin A (650,000 U.I.); vitamin D3 (200,000 U.I.); vitamin E
859 (7,000 mg); vitamin K3 (250 mg); vitamin B1 (250 mg); vitamin B2 (450 mg); vitamin B6
860 (350 mg); vitamin B12 (3 mg); niacinamide (2,500 mg); calcium D-pantothenate (2,000 mg);
861 folic acid (100 mg); choline chloride (50,000 mg).

862 ² One kg of premix contained: lysine (80,000 mg); threonine (280,000 mg); methionine
863 (240,000 mg); tryptophan (120,000 mg); L-valine (240,000 mg).

864

865 **Table 2.**

866 Effect of zinc glycinate (ZnGly) supplementation on animal performances and carcass traits.

	Dietary treatment ¹			SEM ²	P-value
	CON	Zn45	Zn100		
<i>Growth performance</i>					
Average daily intake, kg/d	3.17	3.06	2.95	0.041	0.069
Final body weight, kg	115	111	107	1.380	0.063
ADG ³ , kg/d	0.97 ^a	0.88 ^{ab}	0.83 ^b	0.022	0.031
FCR ³	3.29	3.52	3.56	0.065	0.179
<i>Carcass traits</i>					
HCW ³ , kg	92.7 ^a	89.8 ^{ab}	86.3 ^b	1.080	0.042
CCW ³ , kg	88.2	86.4	83.2	1.000	0.107
Carcass yield (hot), %	80.6	81.0	80.4	0.261	0.612
Carcass yield (cold), %	76.6	78.0	77.5	0.258	0.094
Chilling carcass loss, %	4.85 ^a	3.70 ^b	3.59 ^b	0.156	<0.001
Cooking loss, %	29.7	28.8	26.9	0.588	0.126
Muscle pH at 45 min	6.18	6.21	6.27	0.032	0.509
Muscle pH at 24 h	5.50	5.49	5.50	0.026	0.986
<i>Muscle color descriptors</i>					
L* (lightness)	52.5	53.2	52.0	0.583	0.700
a* (redness)	7.73	8.12	7.56	0.210	0.567
b* (yellowness)	7.98	8.44	7.72	0.270	0.562
C* (saturation)	11.1	11.7	10.8	0.334	0.551
h _{ab} (hue angle)	45.6	46.1	45.3	0.379	0.736
<i>Backfat color descriptors</i>					
L* (lightness)	75.1	74.5	76.0	0.431	0.361
a* (redness)	5.73	6.22	5.72	0.205	0.546
b* (yellowness)	6.35	7.16	6.57	0.257	0.438
C* (saturation)	8.57	9.50	8.72	0.322	0.476
h _{ab} (hue angle)	47.7	48.9	48.8	0.44	0.469

867
868 ¹ CON, control; Zn45, diet containing 45 mg/kg of ZnGly; Zn100, diet containing 100 mg/kg
869 of ZnGly.870 ² SEM, standard error of the mean.871 ³ ADG, average daily gain; FCR, feed conversion ratio (daily intake/ADG); HCW, hot carcass
872 weight; CCW, cold carcass weight.

873

874 **Table 3.**
 875 Effect of zinc glycinate (ZnGly) supplementation on micromineral content in liver and muscle
 876 (mg/kg fresh tissue).

	Dietary treatment ¹			SEM ²	P-value
	CON	Zn45	Zn100		
<i>Liver</i>					
Copper	10.8	11.7	11.1	0.217	0.281
Iron	281 ^b	335 ^a	296 ^b	7.510	0.006
Manganese	3.53	3.61	3.63	0.027	0.286
Zinc	58.4	58.1	59.1	0.336	0.467
<i>Muscle</i>					
Copper	1.88	1.90	1.93	0.027	0.757
Iron	22.9	24.7	27.2	1.130	0.302
Manganese	0.20	0.20	0.20	0.002	0.573
Zinc	19.4	19.6	19.8	0.299	0.867

877
 878 ¹ CON, control; Zn45, diet containing 45 mg/kg of ZnGly; Zn100, diet containing 100 mg/kg
 879 of ZnGly.

880 ² SEM, standard error of the mean.

881

882 **Table 4.**
883 Effect of zinc glycinate (ZnGly) supplementation on intramuscular fat, cholesterol and meat
884 fatty acid profile.
885

	Dietary treatment ¹			SEM ²	P-value
	CON	Zn45	Zn100		
Intramuscular fat, g/100 g	1.65	1.68	1.92	0.105	0.534
Cholesterol, mg/g	0.60	0.52	0.54	0.015	0.060
<i>Fatty acids, mg/100 g</i>					
C10:0	2.22	2.06	2.29	0.143	0.820
C12:0	1.49	1.51	1.83	0.122	0.451
C14:0	20.2	21.2	24.2	1.470	0.524
C16:0	387	399	449	25.00	0.571
C17:0 <i>anteiso</i>	4.76	4.82	5.93	0.367	0.348
C16:1 <i>c9</i>	51.0	54.8	56.5	3.450	0.809
C17:0	2.35	2.81	2.73	0.201	0.613
C18:0	200	196	230	13.30	0.526
C18:1 <i>t9</i>	2.23	2.47	2.35	0.175	0.865
C18:1 <i>c9</i>	650	676	778	46.80	0.505
C18:1 <i>c11</i>	65.4	68.8	74.8	4.150	0.654
C18:2 <i>c9 c12</i>	177	168	195	10.20	0.570
C20:0	2.69	2.74	3.31	0.232	0.484
C20:1 <i>c11</i>	11.0	12.1	14.2	0.856	0.308
C18:3 <i>c9 c12 c15</i>	6.35	5.19	6.70	0.573	0.557
C20:2 <i>c11 c14</i>	5.10	5.23	6.90	0.480	0.231
C20:3 <i>n-6</i>	4.48	4.51	4.38	0.185	0.963
C20:3 <i>n-3</i>	0.80	0.82	0.93	0.110	0.884
C20:4 <i>n-6</i>	29.5	28.4	28.6	0.804	0.832
C22:4 <i>n-6</i>	4.37	4.56	4.72	0.181	0.743
C22:5 <i>n-3</i>	4.20 ^a	3.06 ^b	2.86 ^b	0.171	0.001
C22:6 <i>n-3</i>	0.86	1.24	0.98	0.130	0.500
SFA ³	621	630	719	40.20	0.552
MUFA ³	779	814	925	54.80	0.531
PUFA ³	233	221	251	12.00	0.612
PUFA <i>n-6</i>	221	211	240	11.40	0.601
PUFA <i>n-3</i>	12.2	10.3	11.5	0.662	0.526
PUFA <i>n-6/n-3</i>	18.4	21.5	21.4	0.765	0.174
AI ⁴	0.46	0.47	0.46	0.005	0.626
TI ⁵	1.08	1.09	1.10	0.014	0.822
h/H ⁶	2.38	2.32	2.39	0.027	0.527
HP-PUFA ⁷	50.6	47.8	49.2	1.58	0.782
Peroxidability index ⁸	328	312	342	14.1	0.700

886
887 ¹ CON, control; Zn45, diet containing 45 mg/kg of ZnGly; Zn100, diet containing 100 mg/kg
888 of ZnGly.

889 ² SEM, standard error of the mean.
890 ³ SFA, saturated fatty acids; MUFA, monounsaturated fatty acids; PUFA, polyunsaturated fatty
891 acids.
892 ⁴ Atherogenicity index = (C12:0 + 4 × C14:0 + C16:0)/(MUFA + PUFA *n*-6 + PUFA *n*-3).
893 ⁵ Thrombogenicity index = (C14:0 + C16:0 + C18:0)/(0.5 × C18:1) + (0.5 × other MUFA) +
894 (0.5 × PUFA *n*-6) + (3 × PUFA *n*-3) + (PUFA *n*-3/PUFA *n*-6).
895 ⁶ hypocholesterolemic to hypercholesterolemic ratio = [(sum of C18:1 *c*9, C18:1 *c*11, C18:2 *c*9
896 *c*12, C20:1 *c*11, C18:3 *c*9 *c*12 *c*15, C20:2 *c*11 *c*14, C20:3 *n*-6, C20:3 *n*-3, C20:4 *n*-6, C22:4
897 *n*-6, C22:5 *n*-3, C22:6 *n*-3)/(sum of C14:0 and C16:0)].
898 ⁷ Highly peroxidizable-PUFA = calculated as the sum of PUFA with three or more unsaturated
899 bonds.
900 ⁸ Calculated as: peroxidability index = (Σdienoic × 1) + (Σtrienoic × 2) + (Σtetraenoic × 3) +
901 (Σpentaenoic × 4) + (Σhexaenoic × 5).
902

903 **Table 5.** Effect of zinc glycinate (ZnGly) supplementation on the overall antioxidant capacity,
 904 enzymatic and non-enzymatic antioxidants in muscle.

	Dietary treatment ¹			SEM ²	P-value
	CON	Zn45	Zn100		
<i>Antioxidant enzymes, U/g</i>					
Catalase (CAT)	141 ^b	150 ^b	170 ^a	3.84	0.003
Glutathione peroxidase (GSH-Px)	0.26	0.28	0.28	0.011	0.835
Superoxide dismutase (SOD)	132 ^a	111 ^b	121 ^{ab}	2.76	0.007
<i>Fat-soluble vitamins</i>					
α -tocopherol, μ g/g of meat	3.16	2.96	3.09	0.085	0.646
retinol, ng/g of meat	15.1	14.9	14.1	0.672	0.828
<i>Hydrophilic antioxidant capacity</i>					
TEAC ³	49.9	60.5	61.3	4.42	0.512
FRAP ⁴	32.7	34.3	31.8	1.24	0.726
Folin–Ciocalteu ⁵	0.69	0.66	0.66	0.014	0.468

905
 906 ¹ CON, control; Zn45, diet containing 45 mg/kg of ZnGly; Zn100, diet containing 100 mg/kg
 907 of ZnGly.

908 ² SEM, standard error of the mean.

909 ³Trolox equivalent antioxidant capacity (TEAC). Expressed as mg of Trolox equivalents/g of
 910 muscle.

911 ⁴Ferric reducing antioxidant power (FRAP). Expressed as mg of Fe⁺² equivalents/g of muscle.

912 ⁵Expressed as mg of tannic acid equivalents (TAE)/g of muscle.

913 **Table 6.**

914 Effect of zinc glycinate (ZnGly) supplementation on color stability of backfat and meat.

	Dietary treatment (D) ¹			Time (T) ²			SEM ³	P-value		
	CON	Zn45	Zn100	Day 0	Day 3	Day 6		D	T	D x T
<i>Backfat color descriptors</i>										
L* (lightness)	75.9	76.4	77.0	75.2 ^b	76.7 ^a	77.4 ^a	0.233	0.107	<0.001	0.059
a* (redness)	5.00	5.38	5.23	5.88 ^a	5.44 ^a	4.28 ^b	0.126	0.521	<0.001	0.768
b* (yellowness)	5.47	6.22	6.02	6.68 ^a	6.66 ^a	4.34 ^b	0.186	0.123	<0.001	0.381
C* (saturation)	7.48	8.27	7.99	8.91 ^a	8.62 ^a	6.18 ^b	0.212	0.266	<0.001	0.582
h _{ab} (hue angle)	57.6	54.1	48.4	48.5	50.4	61.2	2.97	0.451	0.174	0.527
ΔE ⁴	4.41	4.23	3.35	-	3.25	4.73	0.293	0.421	0.001	0.956
<i>Meat color descriptors</i>										
L* (lightness)	55.0	55.0	53.9	52.5 ^c	54.6 ^b	56.7 ^a	0.332	0.399	<0.001	0.752
a* (redness)	5.85	5.93	5.55	7.79 ^a	5.28 ^b	4.23 ^c	0.187	0.475	<0.001	0.692
b* (yellowness)	6.41	6.46	5.97	8.03 ^a	6.28 ^b	4.5 ^c	0.196	0.436	<0.001	0.616
C* (saturation)	8.69	8.78	8.16	11.2 ^a	8.22 ^b	6.19 ^c	0.267	0.439	<0.001	0.618
h _{ab} (hue angle)	47.5	47.5	47.0	45.6 ^b	49.7 ^a	46.7 ^b	0.290	0.678	<0.001	0.877
630/580 ratio	1.24	1.24	1.23	1.34 ^a	1.21 ^b	1.15 ^c	0.009	0.617	<0.001	0.927
ΔE ⁴	5.98	5.91	5.81	-	4.68	7.12	0.267	0.966	<0.001	0.740

915 ¹ CON, control; Zn45, diet containing 45 mg/kg of ZnGly; Zn100, diet containing 100 mg/kg of ZnGly.

916 ² Days of refrigerated aerobic storage.

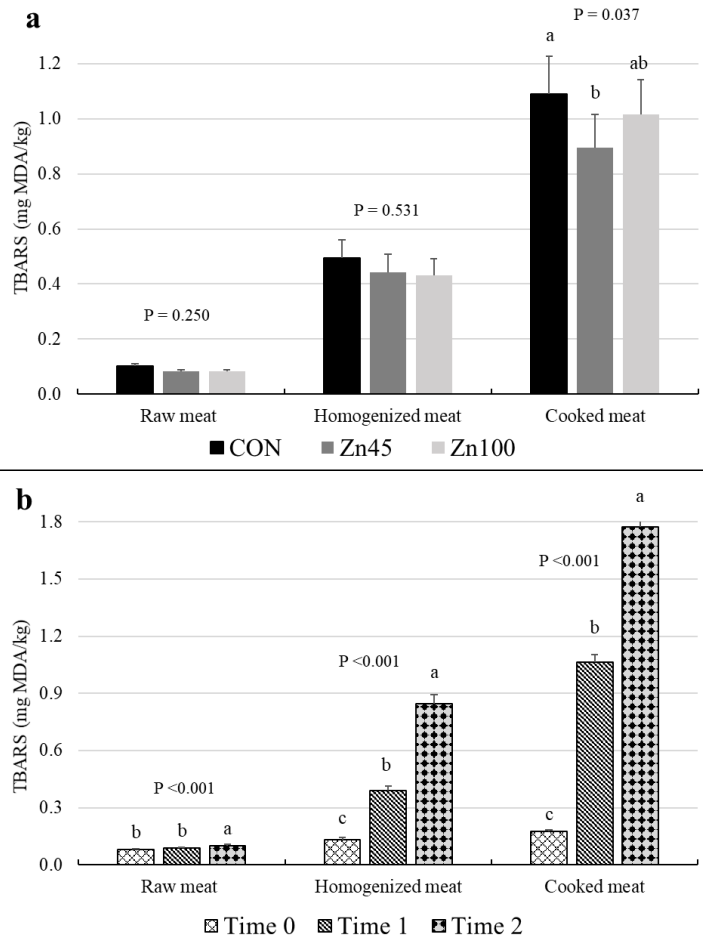
917 ³ SEM, standard error of the mean. ⁴ Total color change between each day of storage and the day 0. Calculated as $\Delta E = [(\Delta L^*)^2 + (\Delta a^*)^2 + (\Delta b^*)^2]^{1/2}$. Where, ΔL^* , Δa^* and Δb^* are the differences in L*, a* and b*, respectively, between day 0 and day 3 or 6.

918

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920

921



922
 923
 924 Fig. 1. Main effect of (a) zinc glycinate (ZnGly) supplementation or (b) time of storage/incubation
 925 on lipid oxidation (TBARS assay) measured in raw and cooked meat slices stored aerobically at
 926 +4 °C or in homogenized meat incubated with Fe³⁺ and ascorbate. Columns are the mean values
 927 and error bars represent the standard error of the mean.
 928 CON, control; Zn45, diet containing 45 mg/kg of ZnGly; Zn100, diet containing 100 mg/kg of
 929 ZnGly.
 930 0, 1, 2 correspond to: days 0, 3, 6 (raw meat); minutes 0, 30, 60 (homogenized meat incubated
 931 with Fe³⁺ and ascorbate); days 0, 2, 4 (cooked meat).
 932 ^{a, b, c} Different superscript letters, within each class (raw meat or homogenized meat or cooked
 933 meat), indicate differences between means ($P \leq 0.05$) tested using the Tukey's adjustment for
 934 multiple comparisons.