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2
3 **The effects of superoxide dismutase-rich melon pulp concentrate on**
4 **inflammation, antioxidant status and growth performance of challenged post-**
5 **weaning piglets**

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20
21 Short title: Melon pulp concentrate fed to challenged piglets

22 **Abstract**

23 Piglets can often suffer impaired antioxidant status and poor immune response during
24 post-weaning, especially when chronic inflammation takes place, leading to lower
25 growth rates than expected. Oral administration of dietary antioxidant compounds
26 during this period could be a feasible way to balance oxidation processes and increase
27 health and growth performance. The aim of the trial was to study the effects of an
28 antioxidant feed supplement (melon pulp concentrate) that contains high concentration
29 of the antioxidant superoxide dismutase (**SOD**) on inflammation, antioxidant status and
30 growth performance of lipopolysaccharide (**LPS**) challenged weaned piglets. Forty-
31 eight weaned piglets were individually allocated to four experimental groups in a 2 x 2
32 factorial design for 29 days. Two different dietary treatments were adopted: a) Control
33 (**CTR**), fed a basal diet, b) Treatment (**MPC**), fed the basal diet plus 30g/ton of melon
34 pulp concentrate. On days 19, 21, 23 and 25 half of the animals within CTR and MPC
35 groups were subjected to a challenge with intramuscular injections of an increasing
36 dosage of LPS from *E. coli* (serotype 0.55:B5) (+) or were injected with an equal
37 amount of PBS solution (-). Blood samples were collected at the beginning of the trial
38 and under the challenge period for interleukin 1 β , interleukin 6, tumour necrosis factor
39 α , haptoglobin, plasma SOD activity, total antioxidant capacity, reactive oxygen
40 species, red blood cells and plasma resistance to haemolysis, and 8-oxo-7, 8-dihydro-
41 2'-deoxyguanosine. Growth performance was evaluated weekly. A positive effect of
42 melon pulp concentrate was evidenced on total antioxidant capacity, half-haemolysis
43 time of red blood cells, average daily gain and feed intake, while LPS challenge
44 increased proinflammatory cytokines and haptoglobin serum concentrations, with a
45 reduced feed intake and gain:feed. The obtained results show that oral SOD
46 supplementation with melon pulp concentrate ameliorates the total antioxidant capacity

47 and the half-haemolysis time in red blood cell of post-weaning piglets, with positive
48 results on growing performance.

49

50 **Keywords:** Cucumis melo, total antioxidant capacity, growth, lipopolysaccharide, pig.

51

52 **Implications**

53 Piglets can show impaired antioxidant status and inflammation processes during post-
54 weaning, leading to lower growth rates than expected. This is especially true when
55 occasional presence of pathogens in the farm causes a chronic inflammation status.
56 In this context superoxide dismutase administration could be a feasible way to
57 overcome inflammation, impaired antioxidant status, and low performance. The
58 present study demonstrates that oral supplementation with melon pulp concentrate,
59 rich in superoxide dismutase, is able to increase the total antioxidant capacity of
60 weaned pigs, contributing to sustain the health status and the growth rate of these
61 animals.

62

63 **Introduction**

64 Inflammation and oxidative stress are closely linked together (Kick *et al.*, 2012;
65 Carillon *et al.*, 2013a) and lead to decreased growth rates and feed efficiency in
66 weaned piglets. This is mainly due to the imbalance in oxidant/antioxidant equilibrium,
67 the decreased activity of major antioxidant enzymes, and the increase in radical-
68 mediated lipid peroxidation and protein and DNA oxidation (Campbell *et al.*, 2013;
69 Gessner *et al.*, 2017). In this context, the use of dietary antioxidant compounds after
70 weaning seems to be a feasible way to overcome impaired antioxidant status and poor
71 immune response in piglets (Bontempo *et al.*, 2014; Jiang *et al.*, 2015a,b). Nowadays

72 there is great interest in the therapeutic application of dietary superoxide dismutase
73 (**SOD**) (Carillon *et al.*, 2013a,b), as a primary antioxidant molecule. A specific
74 cantaloupe melon (*Cucumis melo L.*) from the *Cucurbitaceae* family, known to be
75 characterized by high SOD activity, showed antioxidant and anti-inflammatory
76 properties in *in vitro* and *in vivo* animal models such as rodents and horses (Vouldoukis
77 *et al.*, 2004a,b; Notin *et al.*, 2010).

78 Although some trials have been conducted on orally melon concentrate
79 administration in different species, at the present moment the effects of feeding SOD-
80 rich melon pulp concentrate on piglets subjected to stress are only reported in two
81 studies by Lallès *et al.* (2011) and Royer *et al.* (2016) on stress proteins along the
82 gastrointestinal tract and changes in blood oxidative stress biomarkers, respectively.
83 At experimental level, oxidative and inflammatory stress conditions can be mimed by
84 chronic immune system stimulation with an increasing dose of lipopolysaccharide
85 (**LPS**) challenge (Rakhshandeh and de Lange, 2012). The aim of the present study is
86 to evaluate the effects of oral supplementation of a SOD-rich melon pulp concentrate
87 in increasing antioxidant status, inflammation response, and growth performance in
88 LPS-chronically challenged weaning piglets.

89

90 **Material and methods**

91

92 *Animals housing and experimental design*

93 The present trial was performed at the Centro Clinico-Veterinario e Zootecnico-
94 Sperimentale d'Ateneo di Lodi, Università degli Studi di Milano. In total, 48 crossbred
95 female piglets (Topigs 40 x Topdelta) from the same herd were weaned at 24 ± 1 days
96 of age (BW 7.79 ± 0.17 kg) and divided in four homogeneous experimental groups of

97 twelve animals each in a 2 x 2 factorial arrangement. The piglets were placed in
98 individual pens (0.47 m²) and allocated in the same environmentally-controlled post-
99 weaning room on slatted floor. Each pen was equipped with one standard nursery pig
100 bite-style nipple drinker and a self-feeder to allow for *ad libitum* access to water and
101 feed. Room temperature and ventilation were electronically controlled over a 24 hours
102 period. Starting room temperature was 28°C with a ventilation of 10 m³/hour/piglet and
103 was decreased by 1°C/week until 25°C at the end of the trial.

104 The first factorial arrangement consisted of the administration of a basal diet or
105 the same basal diet plus melon pulp concentrate (Melofeed, Lallemand SAS, Blagnac,
106 France). The second factorial arrangement consisted on a LPS challenge, with
107 repeated increasing intramuscular injections of LPS from *E. coli* (serotype 055:B5,
108 Sigma-Aldrich Canada Ltd, Oakville, ON, Canada; cat. no. L2880) (+) to mimic chronic
109 inflammation, or the injection of an equivalent amount of PBS solution (-). The
110 challenge was performed starting on day 19 of the trial and subsequent injections were
111 performed on days 21, 23 and 25. The initial LPS dosage of 60 µg/kg of BW was
112 increased by 12% at each subsequent injection to reduce endotoxin tolerance
113 (Rakhshandeh and de Lange, 2012). Specifically, the applied concentrations of LPS
114 from the second to the fourth injection day were 67.2, 75.26, and 84.30 µg/kg of BW.
115 Individual body weight was determined prior to each LPS injection to calculate
116 individual total LPS amount to be injected.

117

118 *Experimental diets*

119 To avoid any other influencing stress factor besides the weaning and the
120 challenge, in the present trial pigs were fed with a mash wheat-based basal diet (Table
121 1) for all the trial period with (**MPC**) or without (**CTR**) the inclusion of 30 g/ton of melon
122 pulp concentrate. Melon pulp concentrate was provided in powder form and
123 incorporated through the premix at 1.5 kg/ton of feed to get the final content of 30g of
124 Melofeed/ton of feed (Lalleman SAS, Blagnac, France). The applied dosage of melon
125 pulp concentrate was based on the previous publication of Lallès *et al.* (2001) where
126 Promutase (former name of Melofeed) was supplemented at 5 and 20 g/ton feed.
127 Because a decrease of feed intake was expected due to the LPS challenge, in the
128 present trial the dosage was increased to 30 g/ton feed during the whole experimental
129 period.

130 The experimental diets were formulated to be isocaloric and isoproteic on net
131 energy and CP basis, and were produced with the same batches of feeds by
132 Tracciaverde S.R.L. (Bonemerse, Italy). Both CTR and MPC diets did not contain any
133 antimicrobials or growth promoters, being designed to meet or exceed the nutrient
134 requirements of weaned piglets recommended by National Research Council (NRC,
135 2012), with the exception of suboptimal concentration of tryptophan. In the present trial
136 no inclusion of tryptophan was performed besides its content in the feeds in order to
137 amplify the inflammation effect of the LPS challenge performed in the experimental
138 animals (Le Floc'h and Seve, 2007).

139

140 *Samples and data collection*

141 Blood samples for pro-inflammatory interleukin 1 β (**IL-1 β**), interleukin 6 (**IL-6**)
142 and tumour necrosis factor α (**TNF- α**) and haptoglobin were collected on days 0, 19,

143 21, 23, 25, and 29, while blood samples for plasma SOD activity, total antioxidant
144 capacity (**TAOC**), and reactive oxygen species (**ROS**) serum content were obtained
145 on days 0, 19, 21, 23, 25, 27, and 29. Each blood sample under the challenge period
146 was collected prior the LPS injection. For sampling procedures a 10 mL clot activator
147 vacutainer tube (VF-109SP, Venoject®, Terumo Europe N.V., Leuven, Belgium) was
148 used to yield serum from the cranial vena cava. Blood sample for plasma SOD activity
149 was collected separately using a 4 mL vacutainer tube containing EDTA (VF054STK,
150 Venoject®, Terumo Europe N.V., Leuven, Belgium). The serum was obtained by
151 allowing the whole blood to clot at room temperature for 30 min. The tubes were then
152 centrifuged at 1500 x g, 10 min at 4°C, the supernatant was placed in 2 mL Eppendorf
153 tubes (Eppendorf AG, Amburg, Germany) and subsequently stored at -80°C to prevent
154 changes in antioxidant and inflammation biomarkers. Blood samples for the whole
155 blood and red blood cells (**RBC**) resistance to haemolysis and plasma contribution by
156 Kit Radicaux Libres (**KRL**), and 8-oxo-7, 8-dihydro-2'-deoxyguanosine (**8-oxodGuo**)
157 were collected on days 19, 25, and 29 of the trial. Blood samples for KRL test were
158 collected by 4 mL vacutainer tubes containing EDTA (VF054STK, Venoject®, Terumo
159 Europe N.V., Leuven, Belgium) (Rossi *et al.*, 2013), immediately stored at 4°C,
160 processed within three hours from sampling, and analysed in the next 24 hours after
161 collection. Blood samples for 8-oxodGuo were collected with a 10 mL clot activator
162 vacutainer tube (VF-109SP, Venoject®, Terumo Europe N.V., Leuven, Belgium) and
163 serum was obtained by centrifugation at 1500 x g, 10 min at 4°C. Serum was then
164 placed in a 2 mL Eppendorf tube (Eppendorf AG, Amburg, Germany), and stored at -
165 80°C for pending analysis.

166 A sample of the basal diet was collected at the beginning of the trial and stored
167 at -20°C for pending analyses. Individual piglet BW was recorded on days 0, 8, 15, 19,

168 21, 23, 25, 26 and 29 with an electronic scale (ES100L, Ohaus, Switzerland), and
169 individual feed intake (**FI**) for 0-19 and 19-29 days periods was calculated by
170 subtracting the relative excreta to the total daily-administered amount of feed.
171 Subsequently average daily gain (**ADG**) and gain:feed (**G:F**) were calculated. The
172 health status of the piglets was daily checked and any sanitary treatment was recorded.
173 Morbidity and mortality were recorded. Rectal temperature was manually measured
174 before each LPS injection as a baseline measurement, after two hours from each LPS
175 injection, and on the last day of trial.

176

177 *Chemical analysis*

178 Serum pro-inflammatory cytokines IL-1 β , IL-6 and TNF- α were measured by
179 porcine-specific ELISA kit according to the recommendations of the manufacturer (R
180 and D Systems Inc., Abingdon Science Park, UK). The serum concentrations of
181 haptoglobin were determined by colorimetric assay (Tridelta Phasorange serum
182 haptoglobin assay, Cat. No. TP-801) and expressed on the basis of a standard curve
183 (Cooke and Arthington, 2013). All samples were assayed in duplicate. Intra-assay
184 coefficient of variation were 5.71%, 6.33%, 5.83%, and 7.41%, while inter-assay
185 coefficient of variation were 5.59%, 6.38%, 5.34%, and 6.18% respectively for IL-1 β ,
186 IL-6, TNF- α , and haptoglobin. All the intra- and inter-assay coefficients of variations
187 were within the range of values declared by the commercial product datasheet. Plasma
188 SOD activity and serum TAOC were measured using commercial available kits
189 according to the manufacturer's instructions (Sigma-Aldrich, Cat. No. 19160 and Cat.
190 No. CS0790, respectively), while ROS were evaluated by Cyt C reduction assay
191 (Sigma Aldrich, cat. No. C2506), as reported by Sartorelli *et al.* (2000). The total
192 antiradical activity of whole blood, RBC and the plasma contribution was determined

193 using the KRL biological test (Rossi *et al.*, 2013). Results were expressed as the time
194 (min) that is required to reach 50% of maximal haemolysis. Half haemolysis time for
195 total blood cells (**HT₅₀WB**) and for red blood cells (**HT₅₀RBC**) refers to the whole blood
196 and the red cell resistance to free radical attack, respectively. Lastly, the plasma
197 resistance to haemolysis (**HT₅₀PC**) was calculated by subtracting HT₅₀RBC from
198 HT₅₀WB. Serum concentration of 8-oxodGuo were determined using a competitive
199 ELISA method based on monoclonal antibody highly specific to 8-OHdG (Japan
200 Institute for Control Aging Fukuroi, Japan) following the manufacturer's instructions.
201 For 8-oxodGuo assays six different standard dilutions were used in triplicate for each
202 ELISA plate. Intra- and inter -assay coefficients of variation were respectively 5.56%
203 and 8.86%.

204 The chemical composition of the basal diet was analysed at the beginning of the
205 trial to determine DM (method 930.15), CP (method 984.13), ether extract (method
206 920.39A), ash (method 942.05), Ca (method 968.08) and P (method 946.06) content
207 following the relative Association of Official Analytical Chemists methods of analysis
208 (AOAC, 2005) (Table 1). Neutral detergent fibre content in the diet was determined as
209 reported by Van Soest *et al.* (1991).

210

211 *Statistical analyses*

212 Data relative to inflammatory and oxidative biomarkers, BW, and rectal
213 temperature were performed by ANOVA using the Proc MIXED for repeated
214 measurements in SAS (SAS Inst. Inc., Cary, NC), with the piglet as the experimental
215 unit. The statistical model included the effects of LPS challenge, dietary treatment
216 (CTR, MPC), time and their interaction as fixed effects. Average daily gain, FI and G:F
217 considered two different trial periods corresponding to days 0-19 (pre-challenge) and

218 19-29 (challenge and post-challenge) of the trial. Statistical analyses for these
219 parameters were performed by a GLM procedure in SAS with the piglet as the
220 experimental unit and the corresponding data relative to the pre-challenge period (0-
221 19 days) as covariate for 19-29 days period. Differences were considered significant
222 for $P < 0.05$. A tendency toward a significant difference between treatment means was
223 also considered at $P < 0.10$.

224

225 **Results**

226

227 *Inflammatory and oxidative biomarkers status*

228 In the present trial LPS challenge induced a strong effect on inflammatory status
229 with increased serum concentration of IL-1 β (0.53 ng/mL vs. 0.21 ng/mL, $P < 0.01$), IL-
230 6 (6.99×10^3 ng/mL vs. 3.10×10^3 ng/mL, $P < 0.01$), TNF- α (2.56×10^{-2} ng/mL vs. 1.44
231 $\times 10^{-2}$ ng/mL, $P < 0.01$), and haptoglobin (8.96×10^5 ng/mL vs. 7.82×10^5 ng/mL,
232 $P < 0.05$) (Figure 1), while melon pulp concentrate supplementation did not induce any
233 significant change in these parameters. Time effect and the interaction between LPS
234 challenge and time were always significant ($P < 0.01$) (Figure 1). No significant
235 differences between challenged or not challenged piglets were found at the beginning
236 of the trial for IL-1 β , IL-6, and TNF- α , while a higher haptoglobin content was found in
237 piglets not subjected to LPS on day 0 (9.50×10^5 ng/mL vs. 5.62×10^5 ng/mL; $P < 0.01$).
238 Starting from day 21 challenged groups showed increased concentration of all
239 inflammation biomarkers until day 23 for haptoglobin or day 25 for interleukins and
240 TNF- α ($P < 0.01$).

241 Oxidative biomarkers were not affected by LPS challenge (Table 2), although
242 some trends were found for decreased TAOC (5.30 mM Trolox equivalent vs. 6.46 mM

243 Trolox, $P=0.08$), and increased HT₅₀WB (113.78 min vs. 109.14 min, $P=0.09$) and
244 HT₅₀PC (44.90 min vs. 40.89 min, $P=0.08$). Melon pulp concentrate supplementation
245 increased serum TAOC (7.22 mM Trolox equivalent vs. 4.54 mM Trolox equivalent,
246 $P<0.01$) and HT₅₀RBC (70.71 min vs. 66.41 min, $P<0.01$), but did not induce significant
247 effects on HT₅₀WB, HT₅₀PC, 8-oxo-dGuo concentration, SOD activity, and ROS.
248 Sampling time was always significant ($P<0.01$), with the exception of SOD activity and
249 HT₅₀PC, but no difference was detected considering the interaction between dietary
250 treatment, challenge and time.

251

252 *Growth performance and rectal temperature*

253 No significant effects of LPS challenge ($P=0.47$) or melon pulp concentrate
254 supplementation ($P=0.70$) and their interactions ($P=0.73$) were found on BW. The LPS
255 challenge impaired ADG (291 g/d vs. 490 g/d, $P<0.01$), FI (5.57 kg vs. 7.76 kg, $P<0.01$)
256 and G:F (0.52 vs. 0.62, $P<0.01$) from days 19 to 29. Average daily gain and FI were
257 increased in pigs fed melon pulp concentrate under the challenge and post-challenge
258 period (19-29 days on trial) (422 g/d vs. 359 g/d for ADG, $P<0.05$; 7.18 kg vs. 6.15 kg
259 for FI, $P<0.01$) (Table 3), but no differences were found during the first 19 days of the
260 trial and overall the experimental period. No significant differences were outlined for
261 G:F among the dietary treatments, and the diet x challenge interaction was found not
262 to affect growing performance.

263 Rectal temperature was always increased by LPS challenge at two hours after
264 injection ($P<0.01$, data not shown), while no effect of dietary melon pulp concentrate
265 supplementation was evidenced, with the exception of a significant reduction at two
266 hours after the fourth LPS injection (day 25 of the trial) (CTR= 40.25 ± 0.09 °C vs.
267 MPC= 40.12 ± 0.08 °C, $P=0.02$). No differences in piglet's rectal temperature at the

268 end of the trial (day 29) were found between CTR and MPC groups, also considering
269 the challenge effect or the diet x challenge interaction. During the trial no antibiotic
270 treatments were performed on experimental animals and no deaths occurred within
271 CTR and MPC groups.

272

273 **Discussion**

274 Weaning is a critical stress period for piglets and it is characterized by
275 decreased immune response, FI, and nutrient absorption (Niekamp *et al.*, 2007).
276 During this period, pigs are subjected to a number of stressors such as a sudden
277 separation from the sow, transportation and handling, different environment, liquid to
278 solid feed shift, establishment of a new social hierarchy, and increased exposure to
279 pathogens. All these factors lead to metabolism, immune system, and intestinal
280 functions alterations. Several studies report how serum pro-inflammatory cytokines
281 and their gene expression are increased at weaning (Pié *et al.*, 2004; Lallès *et al.*,
282 2011). In accordance with this, in the present trial initial high concentrations of IL-6 and
283 TNF- α were found in all the experimental subjects, although they were decreased just
284 before the LPS challenge (day 19). At the starting of the trial IL-6 showed half of the
285 concentration further evidenced during the challenge, while TNF- α concentration
286 detected on day 0 and during the LPS injection were comparable, outlining how
287 weaning is able to strongly affect the immune response of piglets.

288 Pro-inflammatory cytokines IL-1 β , IL-6, and TNF- α are reported to be important
289 inducers of the synthesis of acute phase proteins, such as haptoglobin, among others
290 (Carroll *et al.*, 2004). In the present trial a higher concentration of haptoglobin in both
291 no-challenge groups compared to challenge groups was found at the beginning of
292 melon pulp concentrate supplementation. Although all the experimental animals were

293 selected, grouped and placed in the experimental facilities at the same time, this initial
294 difference seems to be unfortunately related to some un-accounted (not estimable)
295 stress factors (Salamano *et al.*, 2008; Cray *et al.*, 2009) rather than a direct effect of
296 high concentration of interleukins that could lead to increased levels of haptoglobin.

297 Besides the strong effect of weaning, occasional presence of pathogens in the
298 farm can causes a chronic inflammation status that can further impairs the immune
299 response and the antioxidant status of piglets. This last scenario can be efficiently
300 obtained from an experimental point of view applying a chronic challenge in the post-
301 weaning period. With this purpose, in the present trial a chronic challenge procedure
302 with LPS was adopted to mimic subclinical or mild clinical disease conditions that
303 frequently occur in the field due to the presence of pathogens that lead to immune
304 system stimulation (Rakhshandeh and de Lange 2012; de Ridder *et al.*, 2012). As a
305 result, in the present trial LPS was found to act on the immune system through
306 increased concentration of serum pro-inflammatory cytokines IL-1 β , IL-6, and TNF- α
307 and haptoglobin according to Carroll *et al.* (2001) and Frank *et al.* (2005), while the
308 administration of melon pulp concentrate did not lead to any significant variation on
309 immune response.

310 Inflammation and oxidative stress have been reported to be closely linked to
311 each other, since the pathways generating the mediators of inflammation are all
312 induced by oxidative stress. (Carillon *et al.*, 2013a). Oxidative stress decreases
313 antioxidant defences and induces elevated radical-mediated lipid peroxidation driven
314 by reactive oxygen and nitrogen species (Campbell *et al.*, 2013). At weaning, piglets
315 can be subjected to an accumulation of ROS that could exceed the antioxidant
316 capacities of the animal, leading to a high susceptibility to oxidative stress (Zhu *et al.*,
317 2012).

318 As a strong effect of the LPS challenge on immune response can be highlighted
319 in this trial, the general lack of significant changes in oxidative biomarkers, with the
320 exception of a trend for decreased serum content of TAOC, can be attributed to other
321 numerous factors. Moreover, independently from the applied dosage to mimic a
322 chronic inflammation in the present trial, it must be noted that the effects of an immune
323 challenge did not always lead to univocal results on oxidative stress biomarkers,
324 depending on the quantification of different antioxidant or pro-oxidant components of
325 oxidative stress (van de Crommenacker *et al.*, 2010). In the present trial oxidative
326 markers such as plasma SOD activity, ROS and 8-oxo-dGuo were not influenced by
327 melon pulp administration, while TAOC and HT₅₀RBC were improved in MPC groups.
328 If the obtained results on ROS can be attributed to their reported very short half-life
329 and high instability, the SOD activity could have been influenced by several factors and
330 can be interpreted in different ways, as the activity of major antioxidant enzymes (**AOX**)
331 is quite variable depending on the experimental design adopted, the challenge
332 performed or the animal species. Increased activity of AOX (SOD, CAT, GPx) can be
333 related both to the reaction of the organism to oxidative stress, or to the stimulation of
334 antioxidant defences when supplying antioxidant compounds in the diet. To
335 discriminate between these two opposite interpretations, additional biomarkers of
336 oxidative stress, such as oxidized proteins (e.g. protein carbonyls) or oxidized lipids
337 (e.g. TBARS, MDA, isoprostanes, lipid peroxides) could have been measured. In fact
338 a study by Royer *et al.* (2016) found that a decreased concentration of oxidative stress
339 biomarkers and an increased activity of AOX confirmed the efficacy of melon pulp
340 concentrate supplementation in pigs. Although dietary SOD mechanism of action is not
341 fully understood at the present moment, Carillon *et al.* (2013a) suggested that melon
342 pulp concentrate administration could induce increased antioxidant defence, as

343 outlined by TAOC and RBC blood resistance to haemolysis in the present trial, through
344 the activation of mRNA transcription or the regulation or induction of complex cellular
345 pathways, involving different transcription factors. These authors hypothesized that the
346 induction of antioxidant enzymes could be regulated at the transcriptional level through
347 the nuclear-factor-E2-related factor (Nrf2)/antioxidant response as demonstrated in
348 some other studies with different antioxidant supplements in humans and *in vitro*
349 (Muchová *et al.*, 2001; Zhang *et al.*, 2012).

350 In the present study, decreased FI and impaired growth performance during
351 LPS challenge period were observed according with Gessner *et al.* (2017). Lower
352 performance under the challenged corresponded to a higher concentration of
353 inflammation markers and an increased rectal temperature of the piglets at two hours
354 after each LPS injection as a sign of the presence of a systemic inflammation (Ceciliani
355 *et al.*, 2012). When an infection occurs, metabolic shifts are characterized by the
356 redistribution of nutrients away from the growth processes toward the immune system
357 function, with a subsequent decrease in feed efficiency for growth (Gessner *et al.*,
358 2017). Moreover the suboptimal concentration of tryptophan supplied in the present
359 trial (approximately 78% of the requirements on total basis for piglets ranging from 7
360 to 25 kg BW) (NRC, 2012) could have represent an amplification factor for the
361 effectiveness of the applied challenge, since tryptophan metabolic demand is strongly
362 increased by the synthesis of acute phase proteins during immune stimulation, with a
363 consequent lower availability for growth (de Ridder *et al.*, 2012).

364 Besides the inflammation effect of the applied challenge, the administration of
365 melon pulp concentrate led to increased performance, which instead was not observed
366 by Lallès *et al.* (2011). It must be outlined that the total duration of administration in the
367 work of Lallès *et al.* (2011) was 5 or 12 days, after an initial two-days period of fasting

368 to induce greater stress protein concentrations. Animals were then slaughtered on
369 days 7 and 14 after weaning. The lack of improved performance in the work of Lallès
370 *et al.* (2011) can be probably due to the short duration of melon pulp concentrate
371 supplementation to translate any positive effect in the gastrointestinal tract or on
372 immune response into improved growth performance.

373 In conclusion, the present study demonstrates that oral supplementation with
374 melon pulp concentrate rich in SOD during the post-weaning period can enhance the
375 total antioxidant capacity, half-haemolysis time of red blood cells and growth rate of
376 post-weaning piglets when oxidative stress and/or inflammation are increased.

377

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381

382 **Declaration of interest**

383 The authors declare no conflicts of interest

384

385 **Ethics statement**

386 The protocol for care, handling, and sampling of animals defined in the present
387 study was reviewed and approved by the Università degli Studi di Milano Animal Care
388 and Use Committee (Protocol No 82/14).

389

390 **Software and data repository resources**

391 Data or models from the present work are not deposited in an official repository

392

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497 2581–2589.

498 **Tables**499 **Table 1** *Ingredients (% w/w) and chemical composition (g/100g dry matter) of the basal diet*500 *fed to the experimental piglets*

501

Ingredients	Feeds
Wheat meal	29.47
Barley meal	23.12
Wheat flaked	14.00
Soybean meal, 48.0% CP	17.00
Sweet whey powder	6.00
Soybean oil	3.00
Corn gluten meal	2.50
Dextrose monohydrate	1.50
Dicalcium phosphate	1.30
L-lysine, 78% Lys	0.57
Calcium carbonate	0.50
Sodium chloride	0.30
Vitamin-mineral premix ¹	0.25
L-Thr	0.23
DL-Met	0.18
Flavour	0.05
Sweetener ²	0.01
Zinc oxide	0.01
Cu sulphate	0.01
Chemical composition³ (g/100g DM)	
DM	88.70
CP	18.01

Ether extract	4.52
Ash	5.08
NDF	14.59
Ca	0.75
Total P	0.57
Digestible energy (Mcal/kg DM)	3.44
Net energy, (Mcal/kg DM)	2.45
Lys	1.25
Met+Cyst	0.80
Thr	0.85
Trp	0.21

502 Lys = lysine; Thr = threonine; Met = methionine; Cyst = cysteine; Trp = tryptophan

503 ¹The vitamin-mineral premix provided the following quantities of vitamins and micro minerals per
504 kilogram of complete diet: vitamin A, 10,500 IU; vitamin D3, 2500 IU; vitamin E, 15 mg; vitamin B1, 1.5
505 mg; vitamin B2, 3.8 mg; vitamin B12, 0.025 mg; vitamin B6, 1.6 mg; calcium pantothenate, 12 mg; nicotinic
506 acid, 15 mg; biotin, 0.15 mg; folic acid, 0.5 mg; vitamin K3, 3 mg; Fe, 100 mg; Cu, 6 mg; Co, 0.75 mg;
507 Zn, 150 mg; Mn, 65 mg; I, 0.75 mg; Se, 0.3 mg; ethoxyquin, 150 mg.

508 ²The sweetener (Optisweet®) was provided by Nutriad (Dendermonde, Belgium)

509 ³Feeds were analysed for DM, CP, ether extract, ash, NDF, Ca and total P according to AOAC (2005).

510 All other values were calculated from NRC (2012).

511 **Table 2** Mean concentration of oxidative biomarkers^{1,2} of post-weaning piglets supplemented with melon pulp concentrate in the diet and subjected
 512 to a chronic LPS challenge^{3,4}.

	LPS challenge				SEM ⁵	P-values		
	-		+			Diet	Challenge	Diet X Challenge
	Dietary treatments ²							
	CTR	MPC	CTR	MPC				
N. of piglets ⁶	12	12	12	12				
SOD (IU/mL)	54.50	54.50	54.28	55.54	3.69	0.66	0.77	0.65
TAOC (mM Trolox equivalent)	4.97	7.94	4.11	6.49	1.71	<0.01	0.08	0.66
ROS (Abs)	0.16	0.15	0.15	0.15	0.03	0.73	0.62	0.74
8-oxo-dGuo (ng/mL)	0.89	0.84	0.87	0.91	0.06	0.99	0.40	0.26
HT ₅₀ WB (min)	106.05	112.23	113.38	114.17	4.60	0.20	0.09	0.32
HT ₅₀ RBC (min)	65.31	71.19	67.51	70.23	2.40	<0.01	0.66	0.26
HT ₅₀ PC (min)	40.74	41.03	45.87	43.94	3.89	0.72	0.08	0.62

513 LPS = lipopolysaccharyde; SEM = standard error of the means; SOD = superoxide dismutase; TAOC = total antioxidant activity, ROS = reactive oxygen species;
 514 8-oxo-dGuo = 8-oxo-7, 8-dihydro-2'-deoxyguanosine, HT₅₀WB = half haemolysis time of whole blood, HT₅₀RBC = half haemolysis time of red blood cells; HT₅₀PC
 515 = plasma contribution to half haemolysis.

516

517 ¹SOD and TAOC were performed on plasma samples; ROS and 8-oxo-dGuo were performed in serum; HT₅₀WB and HT₅₀RBC were performed on whole
518 blood. HT₅₀PC was obtained as the difference between HT₅₀WB-HT₅₀RBC.

519 ²Total antioxidant activity was performed by KRL (Kit Radicaux Libres) biological test (Rossi *et al.*, 2013). Results are expressed as the time (min) required to
520 reach 50% of maximal haemolysis (HT₅₀), which refers to the whole blood, red blood cells and plasma resistance to free-radical attack.

521 ³The challenge was performed from day 19 to 25 of the trial with increasing dosages of lipopolysaccharide (LPS from *E. coli* serotype 055:B5). Subsequent
522 intramuscular injections of LPS were performed on days 19, 21, 23 and 25. Initial concentration of LPS was 60 µg/kg of BW and the dosage was increased by
523 12% at each subsequent injection to reduce endotoxin tolerance and mimic a chronic inflammation in piglets. The applied concentrations of LPS from the second
524 to the fourth injection day were 67.2, 75.26, and 84.30 µg/kg of BW. Individual body weight was determined prior each LPS injection to calculate individual total
525 LPS amount to be injected.

526 ⁴Dietary treatments: CTR- = piglets fed the basal diet and not subjected to the LPS challenge; CTR+ = piglets fed the basal diet and subjected to the LPS
527 challenge; MPC- = piglets fed the basal diet added with 30g/ton of melon pulp concentrate (Melofeed, Lallemand SAS, Blagnac, France) and not subjected to
528 LPS challenge; MPC+ = piglets fed the basal diet added with 30g/ton of melon pulp concentrate (Melofeed, Lallemand SAS, Blagnac, France) and subjected to
529 LPS challenge.

530 ⁵SEM = pooled SEM. Means are presented as least square means.

531 ⁶Piglets were reared in individual pens (0.47 m²) with *ad libitum* access to feed and water.

532 **Table 3** Body weight (BW), average daily gain (ADG), feed intake (FI) and gain:feed (G:F) of post-weaning piglets supplemented with melon
 533 pulp concentrate in the diet and subjected LPS challenge¹

	LPS challenge				SEM ⁴	P-values		
	-		+			Diet	Challenge	Diet X Challenge
	Dietary treatments ²							
	CTR	MPC	CTR	MPC				
N. of piglets ³	12	12	12	12				
BW (kg)								
Day 0	7.79	7.79	7.78	7.79	0.71	0.70	0.47	0.73
Day 29	17.94	18.48	16.98	17.37				
ADG (g/d)								
Days 0 to 19	298	297	343	328	51.98	0.79	0.13	0.78
Days 19 to 29	464	517	255	327	60.69	0.05	<0.01	0.75
Days 0 to 29	350	368	317	331	56.34	0.50	0.14	0.91
FI ⁴ (kg/period)								
Days 0 to 19	8.56	8.55	9.50	9.31	1.24	0.87	0.18	0.89
Days 19 to 29	7.31	8.22	4.99	6.15	0.67	<0.01	<0.01	0.73

Days 0 to 29	15.74	16.63	14.65	15.56	1.75	0.31	0.22	0.10
G:F								
Days 0 to 19	0.64	0.65	0.69	0.66	0.04	0.71	0.12	0.32
Days 19 to 29	0.62	0.63	0.51	0.53	0.05	0.64	<0.01	0.72
Days 0 to 29	0.64	0.64	0.63	0.61	0.03	0.69	0.18	0.41

534 LPS = lipopolysaccharide; SEM = standard error of the means

535 ¹The challenge was performed from d 19 to 25 of the trial with increasing dosages of lipopolysaccharide (LPS from *E. coli* serotype 055:B5). Subsequent
536 intramuscular injections of LPS were performed on days 19, 21, 23 and 25. Initial concentration of LPS was 60 µg/kg of BW and the dosage was increased by
537 12% at each subsequent injection to reduce endotoxin tolerance and mimic a chronic inflammation in piglets. The applied concentrations of LPS from the second
538 to the fourth injection day were 67.2, 75.26, and 84.30 µg/kg of BW. Individual body weight was determined prior each LPS injection to calculate individual total
539 LPS amount to be injected.

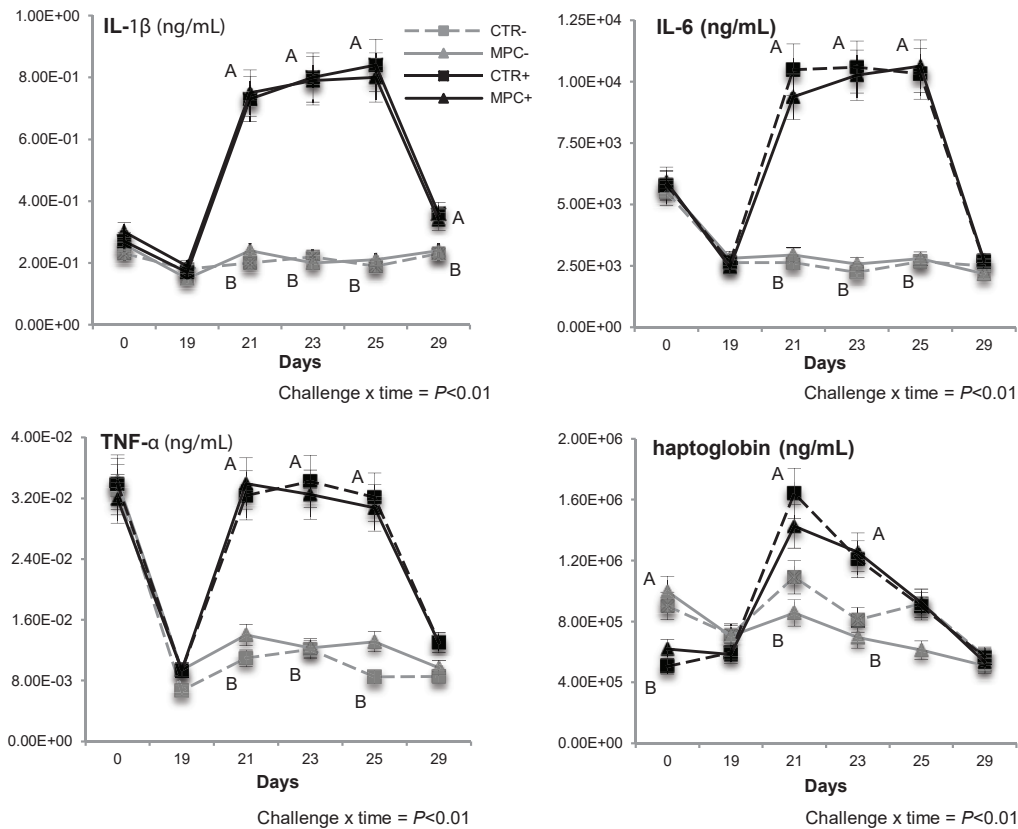
540 ²Dietary treatments: CTR-= piglets fed the basal diet and not subjected to the LPS challenge; CTR+= piglets fed the basal diet and subjected to the LPS
541 challenge; MPC-=piglets fed the basal diet added with 30g/ton of melon pulp concentrate (Melofeed, Lallemand SAS, Blagnac, France) and not subjected to
542 LPS challenge; MPC+= piglets fed the basal diet added with 30g/ton of melon pulp concentrate (Lallemand SAS, Blagnac, France) and subjected to LPS
543 challenge.

544 ³Piglets were reared in individual pens (0.47 m²) with *ad libitum* access to feed and water.

545 ⁴SEM = pooled SEM. Means are presented as least square means.

546 **List of figure captions**

547 **Figure 1** Challenge effect on Interleukin 1 β (IL-1 β), Interleukin 6 (IL-6), Tumour Necrosis Factor α (TNF- α) and haptoglobin serum
548 concentration in post-weaning piglets supplemented with melon pulp concentrate¹ in the diet and subjected to chronic LPS challenge^{2,3}



549

550 ¹Dietary treatments: CTR-= piglets fed the basal diet and not subjected to the LPS challenge; CTR+= piglets fed the basal diet and subjected to the LPS
551 challenge; MPC-=piglets fed the basal diet added with 30g/ton of melon pulp concentrate (Melofeed, Lallemand SAS, Blagnac, France) and not subjected to
552 LPS challenge; MPC+= piglets fed the basal diet added with 30g/ton of melon pulp concentrate (Melofeed, Lallemand SAS, Blagnac, France) and subjected to
553 LPS challenge.

554 ²The challenge was performed from day 19 to 25 of the trial with increasing dosages of lipopolysaccharide (LPS from *E. coli* serotype 055:B5). Subsequent
555 intramuscular injections of LPS were performed on days 19, 21, 23 and 25. Initial concentration of LPS was 60 µg/kg of BW and the dosage was increased by
556 12% at each subsequent injection to reduce endotoxin tolerance and mimic a chronic inflammation in piglets. The applied concentrations of LPS from the second
557 to the fourth injection day were 67.2, 75.26, and 84.30 µg/kg of BW. Individual body weight was determined prior each LPS injection to calculate individual total
558 LPS amount to be injected.

559 ^{3A,B}Different letters refer to significant differences between challenged and not challenge piglets for $P<0.01$.