

Short cold exposures during incubation and postnatal cold temperature affect performance, breast meat quality, and welfare parameters in broiler chickens

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| 3 | Meat quality in cold-acclimated chickens |
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25 ABSTRACT

Cold stimulations during egg incubation were reported to limit the occurrence of ascites in broilers submitted to cold temperature after 14 d of age. However, data is lacking on the impacts of such strategy in case of cold temperature condition at chick starting. This study aimed to evaluate the effects of incubation and post-hatch cold challenge on performance, breast muscle integrity and meat processing quality in broiler chickens.

Ross 308 eggs were incubated under control temperature (I_0 , 37.6°C) or submitted to 15°C 31 during 30 min on d 18 and 19 of incubation (I_1). Chicks from each group were reared in floor 32 pens either at standard rearing temperature (T₀), from 32°C at d 0 to 21°C at d 21, or exposed 33 34 to colder rearing temperature (T_1) , from 29°C at d 0 to 21°C at d 21 of age. Birds were then all kept at 21°C until slaughter (d 40), when body weights (BW), feed conversion ratio 35 (FCR), breast muscle yield, meat processing quality and the occurrences of meat defects, 36 hock burns and pododermatitis were recorded. No significant impact of incubation conditions 37 on hatchability was observed. At d 40, BW was greater under T_1 than T_0 condition, with T_0 38 39 females (but not males) presenting greater BW after I₁ than after I₀ condition. On the whole period, T₁ chickens presented lower FCR than T₀, and greater breast meat yields at d 40. The 40 occurrence of white striping was greater in I_1T_1 males than in all other groups, except for the 41 42 I_0T_1 males. Hock burns were more frequent in I_1T_1 males than in all females and I_0T_0 males, whereas the occurrence of pododermatitis was lower in T₀ males than in other groups. Despite 43 some positive effects of I₁ incubation on growth after starting under low ambient temperature, 44 this study reveals the limits of such strategy concerning chicken health and welfare, 45 demonstrating that early thermal environment is a major component of the quality and 46 47 sustainability of chicken meat production.

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49 Key words: broiler, incubation, meat quality, white striping, welfare

INTRODUCTION

51 Chicken production is expected to increase continuously (+1.8%/year) for the next decades 52 (Alexandratos and Bruinsma, 2012), in view of the growing global demand in proteins from 53 animal source and the efficiency of this production (Petracci and Cavani, 2012). However, the 54 environmental, economic and social impacts of the production systems, including meat 55 quality and animal welfare are criteria of concern for consumers and citizens.

56 It was suggested that the genetic selection of fast-growing broilers on growth rate, feed efficiency and breast meat yield have simultaneously increased the sensitivity of these birds to 57 temperature variations (Piestun et al., 2008; Havenstein et al., 2003a,b; Zuidhof et al., 2014). 58 59 Because the thermoregulatory system of chicks is immature at hatching (Tzschentke, 2007), they remain very sensitive to postnatal cold temperatures during the first days of breeding 60 (Collin et al., 2003; Mujahid and Furuse, 2009). Environmental conditions during the 61 perinatal period are considered as critical for the chicken later performance and fitness 62 (Guilloteau et al., 2019). Especially, brooding chicks at low temperatures (26.7°C vs. 32.2°C) 63 64 results in a decreased feed efficiency and increased mortality rate (Renwick and Washburn, 1982). Fast-growing broilers exhibit greater ascites prevalence (Druyan et al., 2007) and leg 65 disorders (Yalçin et al., 2007; Zhang et al., 2014; Nyuiadzi et al., 2017) when they are 66 67 submitted to low rearing temperatures post-hatch.

Few studies showed that the exposure to cold temperature during embryogenesis reduced the prevalence of ascites and changed thermoregulatory mechanisms in the early life of chicks, helping them to cope with low rearing temperatures post-hatch (Shinder et al., 2011; Akşit et al., 2013). These authors showed that it decreased mortality rate, but it also increased the antioxidant activity of catalase in the liver of chicks at hatching (Loyau et al., 2014). Cold treatment in embryonic life can affect chick behavior (Bertin et al., 2018), but also performance. Van der Pol et al. (2013) showed that cold brooding temperature resulted in

lower BW than normal brooding temperature at 4 d of age. However, long-term positive 75 76 effects of embryonic and postnatal cold acclimation in broilers were reported on body weight (Shinder et al., 2009) and breast muscle yield (Shinder et al., 2011) when broiler chickens are 77 reared at standard temperature. Their treatment was applied in the last phase of 78 embryogenesis (days 18 and 19 of incubation), just before the blood concentrations of 79 triiodothyronine, a major hormone involved in thermoregulation and hatching process, is 80 reaching a peak (Reyns et al., 2003). The late phase of embryogenesis corresponds to the 81 switch of embryos from the ectothermic phase to the endothermic phase when a greater ability 82 to produce heat is acquired (Minne and Decuypere, 1984; Nichelmann and Tzschentke, 2002; 83 Tzschentke, 2007). Cold stimulations during embryogenesis were shown to induce long 84 lasting effects partially limiting the detrimental effects of postnatal cold temperatures on feed 85 conversion ratio in males reared in cages (Nyuiadzi et al., 2017). 86

87 In order to ensure the sustainability of poultry meat production, not only performance but also meat processing quality and animal welfare have to be considered. Metabolic diseases, 88 including ascites, but also muscle myopathies including white striping and wooden breast 89 have recently become a major source of loss for poultry meat production of fast-growing 90 broilers (Julian, 2005; Kuttappan et al., 2016). The rapid growth rate, the increase in breast 91 meat yield and gender are factors affecting the prevalence of white striping (Petracci and 92 Cavani, 2012; Alnahhas et al., 2016). The latter authors have pointed out genetics as the 93 major determinant of this meat defect. To a lesser extent, environmental and management 94 factors, including nutrition (Meloche et al., 2018) also contribute to the variance of the white 95 96 striping occurrence in breast muscle (Bailey et al., 2015). However, the impacts of rearing temperatures and early life environment on this defect has been poorly investigated, except in 97 the case of heat embryonic exposure (Clark et al., 2017). 98

To our knowledge no experimentation has been carried out on the effects of cold exposure during embryogenesis on meat processing quality and muscle defects when chickens are later submitted to standard or low environmental temperature. Thus, the objective of the present study was to assess the performance and meat quality at slaughter age in broilers exposed to low or standard temperature incubation in interaction with postnatal low or standard ambient temperature, while recording the impacts of these treatments on animal performance as well a as on some animal welfare criteria.

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MATERIALS AND METHODS

All experimental procedures were approved by the Ethics Committee for Animal
Experimentation Val de Loire (CEEA Val de Loire, Tours, France, N°2014111809444741
(APAFIS#70).03).

111 Incubation Process

A total of 1,200 fertile Ross 308 broilers eggs with 3 d of storage from 36 wk-old breeder 112 flock were provided from a commercial hatchery (Couvoir Perrot, Pommerit-Jaudy, France). 113 Prior to incubation, average egg weight was determined (62.3 \pm 0.7 g) and eggs were 114 randomly divided in two treatment groups experiencing control (I_0 at 37.6°C and 56% RH) or 115 Poultry 116 cold incubation (\mathbf{I}_1) at PEAT INRA Experimental Facility (2018, 117 https://doi.org/10.15454/1.5572326250887292E12). I₁ consisted in exposing 3 trays of 132 eggs to 15°C and 81% RH during 30 min on d 18 and d 19 of incubation by transferring them 118 from the hatcher to a cold room, eggs being incubated at 37.6°C and 56% RH for the 119 120 remainder of incubation (Figure 1). This treatment resulted in a minimal eggshell temperature (\pm SD) of 26.9°C \pm 2.1°C and maximal eggshell temperature of 33.1 \pm 0.6°C, measured by 121 thermal imaging after 30 minutes of exposure. On d 7 and d 14 of incubation, unfertile and 122

undeveloped eggs were eliminated after candling. At d 18 of incubation, all eggs were moved to a common hatcher set at 37.6°C and 70% RH, with only the 30 min interruption at d 19 for the transfer of group I_1 to the cold room.

126 *Hatching*

Into the hatcher, the numbers of hatched chicks in control (I_0) and cold (I_1) incubation were 127 recorded at 480 h of incubation, and then at 504 hours (opening of the hatcher) to calculate 128 129 hatchability as the ratio of hatched chicks on fertile incubated eggs. Chicks from each treatment were weighed, wing-banded and sexed by cloacal observation. The body 130 temperature (Tb) of chicks was measured individually at hatching (on dry chicks after 131 opening the hatcher) by inserting an electronic thermometer (PX-TH519, Tex, Pelimex, 132 Ingwiller, France) in the cloaca. Fifty chicks from both treatment groups were randomly 133 134 selected for quality measurement according to Tona et al. (2003). Briefly, the score of 8 parameters including activity, down and appearance, retracted yolk, eyes, legs, navel area, 135 136 remaining membrane, and remaining yolk on the navel were measured to determine the chick 137 quality. Each score was summed and scored out of 100, with 100 being the highest score value for the best chick quality. 138

139 *Rearing Period*

Four-hundred chicks from each incubation group were divided randomly into 2 groups and were transferred to 2 identical controlled rooms under standard rearing conditions T_0 , decreasing from 32°C at d 0 to 21°C at d 21, or under cold rearing conditions T_1 , consisting in continuously decreasing temperature from 29°C at d 0 to 21°C at d 21, respectively (Figure 1). From d 22 to d 40, ambient temperature was maintained at 21°C in both conditions T_0 and T_1 . Chicks were reared in groups of same sex per condition with 4 repetitions per incubation treatment, sex and treatment group at the rate of 23 males or 27 females from d 0 to d 21,

reduced to 19 males or 22 females per group on the basis of the mean BW and standard error 147 of the group from d 22 to d 40 in order to limit the rearing density in pens at slaughter age. 148 Chicks were reared in floor pens (surface = 2.93 m^2) with wood shaving litter (1.5 kg/m^2). 149 Water and pelleted feed were supplied *ad libitum* until slaughter age (d 40). 150

Performance 151

Chickens were weighed at d 0, d 21, and d 40 of age. Feed consumption was recorded on each 152 153 floor pen from d 0 to d 21 and from d 22 to d 40. Feed conversion ratio (FCR) was calculated for both periods as the ratio of feed consumption by body weight (**BW**) gain per pen during 154 both periods and during the entire rearing period. 155

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Muscle and meat quality parameters

At d 40, 8 males and 8 females per floor pen per treatment group $(I_0T_0, I_0T_1, I_1T_0, I_1T_1)$ 157 representatives of their pen (average weight and SD) were slaughtered after 8 h of feed 158 withdrawal, in the experimental processing plant of INRA (UE PEAT, Nouzilly, France). 159 After evisceration, whole carcasses were stored at 4°C for 18 h until cut and deboned for meat 160 characteristics analyses. Abdominal fat, Pectoralis major and minor (breast) muscles were 161 weighed to calculate their yields in relation to body weight. The ultimate pH (pHu) of the 162 163 muscle was measured 24 h postmortem using a portable pH meter (model 506; Crison Instruments SA, Alella, Bercelona, Spain) by inserting the electrode in the left Pectoralis 164 165 major muscle as described by Berri et al. (2001). The color measurements of the muscle defined by 3 components, lightness (L*), redness (a*) and yellowness (b*) values according 166 to the CIE trichromatic (Girolami et al., 2013), were measured by using a Miniscan 167 Spectrocolorimeter (Hunterlab, Reston, VA) as described by Alnahhas et al. (2014; 2015). 168 The occurrence of meat quality defects (Kuttappan et al., 2012; Sihvo et al., 2014; Mudalal et 169 170 al., 2015), white striping (WS) and wooden breast (WB) were evaluated macroscopically in

Pectoralis major muscles on a scale of 3 classes according to Alnahhas et al. (2016). The 171 172 intramuscular lipid content was calculated by near-infrared spectroscopy using a Nirflex N-500 spectrometer (Buchi, Rungis, France) from samples of Pectoralis major collected at 40 d 173 of age (Alnahhas et al., 2016). The lipid peroxidation value was determined by the 174 quantification of thiobarbituric acid-reactive substance (TBARS) using the method developed 175 by Lynch and Frei (1993). Drip loss was determined according to Berri et al. (2007). The 176 177 cooking loss of the breast muscles was measured as previously described by Baéza et al. (2012) and Alnahhas et al. (2016). The average Warner-Bratzler shear force value (N/cm²) of 178 the meat was determined on 3 samples of cooked breast muscle $(1 \times 1 \times 3 \text{ cm})$ using an 179 180 Instron universal testing instrument (Instron 5543, Instron S.A., Guyancourt, France) as described by Honikel (1998) and Alnahhas et al. (2016) in order to assess the tenderness of 181 the meat. 182

183 Parameters related to chicken welfare

At slaughter (d 40), hock burn and pododermatitis occurrences were recorded on all slaughtered animals. At the end of the experiment, the litter of each floor pen was weighed. After homogenization of the litter amount in the pen at 40 days of age, 1 kg of litter from each floor pen was sampled and analyzed for dry matter (MS) to evaluate the litter moisture depending on the incubation and postnatal treatments.

189 Statistical Analysis

Data were analyzed using the statistics software Statview (version 5.0; SAS Institute, Cary, NC). The hatchability results were analyzed by using a Chi-square test. The effect of incubation treatment on body weight and body temperature at hatching was determined by one-way ANOVA followed by Student-Newman-Keuls test, while scores of chick quality were analyzed by a non-parametric Kruskal-Wallis test, because of of the heterogeneity of the variance of the data set. The effects of incubation condition, rearing condition and sex and their interactions on body weight, feed conversion ratio (n = 4 groups), carcass and meat quality (breast meat yield, abdominal fat yield, shear force value, pH_u, L*, a*, b*, cooking loss, drip loss, technological performance) and litter characteristics were performed by ANOVA followed by a Student-Newman-Keuls test. The frequencies of occurrence of pododermatitis, hock burn, wooden breast and white striping were analyzed by Chi-square tests.

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RESULTS

204 Effects of incubation and postnatal treatment on hatchability, chick quality and 205 performance

Hatchability of fertile eggs was not significantly affected by the cold incubation treatment (P 206 = 0.801) with hatchabilities over 95% in both groups in our experimental conditions (Table 207 1). However, at 480 h of incubation, i.e. 24 h before opening the hatcher, 43% of the I_0 chicks 208 were hatched, but only 36% of the I_1 chicks (P = 0.007). At hatching, neither the male chick 209 percentage nor the quality of chicks from cold-incubated eggs I_1 was different from that of 210 211 control chicks I_0 (P = 0.765 and 0.415, respectively; Table 1). Body temperature tended to be greater in I_1 than in I_0 chicks at hatching (P = 0.059), but BW (Table 1) was not influenced by 212 213 the incubation thermal treatment (P = 0.681). There were significant interactions of incubation condition, postnatal rearing temperature and sex on BW at d 21 (P = 0.008) and d 214 40 of age (P = 0.038; Table 2). At d 21, male BW were not different between I₀ and I₁ at 215 standard temperature T₀, while male BW was greater for I₁ than for I₀ group under cold 216 ambient temperature T₁ (P < 0.05; Table 3). On the contrary, I₁ females exhibited greater BW 217 at d 21 than I_0 ones at T_0 (P < 0.05), whereas there were no difference observed between 218

incubation conditions under cold ambient temperature T_1 . At slaughter age (d 40), body 219 220 weights of both males and females were greater under cold rearing temperature T_1 than under standard conditions T₀. In females (but not in males) under standard temperature T₀, I₁ group 221 exhibited greater slaughter BW than I_0 ones (P < 0.05; Table 3). The postnatal cold 222 temperature and the sex affected the FCR from d 0 to d 21 with greater FCR in females than 223 in males and in chickens reared under lower ambient temperature T_1 than at T_0 (P < 0.001 and 224 P < 0.001, respectively; Table 2). Between d 22 and d 40 (when ambient temperatures were 225 maintained at 21°C for both groups T₀ and T₁), FCR of chickens incubated in control 226 conditions I_0 was lower in T_1 than in T_0 group, while it did not differ depending on the 227 228 postnatal temperature in I₁-incubated chickens (Table 3). During this period, females exhibited higher FCR than males (P < 0.02). On the whole period from d 0 to d 40 (Table 2), 229 females exhibited greater FCR than males (P = 0.002) and T₁ chickens presented lower FCR 230 231 than T_0 chickens (P = 0.007).

Effects of incubation, postnatal treatment and sex on body composition and meat quality parameters

No effect of incubation conditions on breast meat yield and yellowness of the meat were 234 observed (P = 0.838 and P = 0.733, respectively). The breast meat yields were greater under 235 236 cold rearing temperature T_1 (P = 0.033) than under standard temperature T_0 , and females showed greater percentage of fillet than males (P < 0.001; Table 2). A greater intensity of 237 yellowness (b*) of the fillet was observed in males compared to females (P < 0.001) and in 238 chickens reared under cold temperature T_1 the first 21 days than under T_0 (P = 0.022; Table 239 3). Females exhibited lower thigh yield (P < 0.001) and lightness (L*; P = 0.026) but greater 240 abdominal fat percentage (P < 0.001), pHu (P < 0.001) and processing yield (P < 0.001) than 241 males (Table 2). The drip loss and the shear force of the cooked breast meat were not 242 influenced by the incubation treatment, rearing conditions or sex (P > 0.05, data not shown). 243

There was a significant interaction of incubation treatment and sex on fillet intramuscular lipids of the breast (P = 0.018). The highest lipid content was observed in I₀ male chickens and the lowest in both females I₀ and I₁ groups (Table 3). A significant interaction of incubation treatment and postnatal condition was measured on the redness intensity (a*; P =0.032) with a higher intensity of redness measured in I₀T₁ group than in the I₀T₀ and I₁T₀ groups, the I₁T₁ group presenting intermediate value (Table 3).

The observation of breast meat for the detection of meat defects showed no case of wooden breast but a difference in the occurrence of white striping between experimental groups. Indeed, the occurrence of white striping was greater in males from the cold-incubated and exposed I_1T_1 chickens (with 58% occurrence) as compared to all other groups except the I_0T_1 males (Table 3). The lowest occurrences, below 18%, were observed in I_1T_0 males and females and in I_0T_1 females.

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257 Effects of incubation and postnatal treatment on chicken welfare

The occurrence of hock burns and pododermatitis were measured on carcasses at slaughter as 258 parameters related to welfare troubles depending on incubating and rearing conditions and 259 sex. Hock burns were more frequently observed in males experiencing cold both during 260 incubation and post hatching (I₁T₁ with 45% occurrence) as compared to all female groups 261 262 and males incubated and reared in standard conditions (P < 0.05). The male chickens with only one cold exposure (I_0T_1 and I_1T_0 males) presented intermediate values (Figure 2). The 263 occurrence of pododermatitis at slaughter was lower (below 28%) in males under standard 264 rearing conditions (in both incubation conditions) than in all other groups (Figure 3A), where 265 occurrences were over 64%. Greater occurence of pododermatitis were observed in I_1T_1 males 266 and females than in all other female groups (P < 0.05; Figure 3B). 267

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DISCUSSION

In an attempt for assessing the positive and negative impacts of perinatal thermal environment 270 271 of chicks, the main objective of this experiment was to study the effects of cold manipulations during incubation in male and female fast-growing chickens that were later exposed to 272 standard or to low postnatal temperature. The parameters considered consequently to these 273 274 treatments relate to growth performance and feed efficiency, but also meat processing quality and defects, and animal welfare criteria. Previous experiments had permitted to compare the 275 effects of two published methods of cold embryo manipulation (Shinder et al., 2011; Yalçin et 276 al., 2012) combined to continuous or fluctuating 4°C-colder or standard postnatal 277 temperatures (Nyuiadzi et al., 2017) during the first 21 d of age in Ross 308 broilers. These 278 279 studies conducted in pens for the two first and in cages for the latter allowed determining thermal conditions of incubation and postnatal rearing enhancing cold adaptation of chicks 280 281 that had little to no impact on performance in the long term. The present experiment, realized 282 in floor pens, confirms the positive effects of these treatments on performance parameters under normal temperature or posthach cold, however in a sex dependent manner. However, 283 the cumulative exposures to cold both during incubation and posthatching had some 284 detrimental effects on chicken welfare and meat quality parameters in the long term, 285 especially in males. 286

Concerning hatching parameters, on the one hand, no negative effect of incubation temperature I_0 or I_1 was observed on hatchability (over 95% in both conditions) or on body temperature neither body weight of hatched-chicks, as previously shown by Shinder et al. (2011) and Nyuiadzi et al. (2017). Nevertheless, cold treatment at the end of incubation slightly slowed down hatching process as demonstrated by the lower percentage of hatched

chicks in I₁ compared to I₀ at 480 h of incubation. This result is not surprising since a faster 292 293 hatching rate was observed following heat-stimulation during embryogenesis (Piestun et al., 2008), with half percentage of hatched chicks at 480 h in heat-treated groups relative to 294 controls, suggesting an impact of thermal treatments during incubation on hatching window. 295 On the other hand, no significant difference between I_0 and I_1 groups was observed on chick 296 297 quality, contrary to what was recently observed in the study of Nyuiadzi et al. (2017), where 298 the quality score related to the remaining membrane at the navel was lower in I_1 group than the one obtained with control incubation, probably due to a greater chick immaturity at 299 hatching. Sex-ratio was not affected by incubation treatment, whereas the male percentage at 300 301 hatching was previously reported to be increased by embryonic warm-stimulation during late incubation (Tzschentke and Halle, 2009). 302

As expected, both the sex and the postnatal temperature markedly affected growth and feed 303 efficiency, with, at d 21 and at d 40 of age, a significant interaction between prenatal and 304 postnatal thermal conditions and sex on the body weight of the birds. At d 21, just at the end 305 306 of the thermal treatments, the double cold exposure of group I_1T_1 was beneficial for growth in males, whereas female growth was stimulated by in ovo or post-hatch cold treatments or both. 307 This effect persisted even after the treatment, as shown by the greater body weights at 308 309 slaughter age exhibited by males reared under cold conditions and all females that have experienced at least one exposition to cold (*in ovo* and/or during rearing). The cold incubation 310 treatment seemed particularly beneficial in terms of growth for males under cold postnatal 311 temperature and in females under normal postnatal temperature. This may be the result of 312 metabolic or endocrine differences between sexes in the early growth phases. In particular, 313 314 there are sex-specific responses to hormonal factors or inhibitors and nutrients in the embryonic stage (Dainat et al.; 1991; Kocamis et al.; 1998, Bello et al., 2014) that may affect 315 316 growth in the long term, supporting this hypothesis. Whatever the postnatal temperature

applied, no detrimental effects of incubation condition on growth were observed. This result 317 318 is consistent with the previous studies reporting positive effects of cold incubation treatments (Shinder et al., 2011) and of postnatal cold-conditioning (Shahir et al., 2012) on BW at 319 320 slaughter age in males submitted to ascites-inducing conditions. The positive impact of cold on BW was even clearer in the present study compared to the transient effect already observed 321 with the same treatments applied in birds reared in cages until 21 d of age (Nyuiadzi et al., 322 2017). One hypothesis is that the birds exposed to cold have consumed more food for 323 sustaining lower post-hatching temperatures, chickens regulating their energy intake 324 depending on ambient temperature post-hatching. However, this is expected to lower feed 325 326 efficiency while increasing heat production. This was actually the case between d 0 and d 21 during the postnatal cold treatment, with greater feed conversion ratios observed in T₁ groups, 327 consistent with our previous results (Nyuiadzi et al. 2017) and those of Aksit et al. (2013) 328 329 recorded from d 0 to d 42 in chickens issued from old flocks and postnatally exposed to cold temperatures. However, because FCR became lower from d 22 onward, especially in the I₀ 330 groups submitted to T₁, broilers submitted to cold challenge during rearing were more 331 332 efficient on the overall period than those reared in standard conditions, whatever the incubation treatment considered. This result supports the hypothesis that chickens submitted 333 334 to slightly colder conditions during the first 3 weeks of age can gain efficiency by retaining more energy and tissue when placed in standard thermal conditions afterwards. This suggests 335 that metabolic adaptation might have occurred following the cold treatments at young age, in 336 the line of changes of the expression of metabolic enzymes previously reported by Loyau et 337 al. (2014) in case of embryonic exposure to 1°C lower incubation temperatures. 338

One other remarkable outcome of the present study was the 4% higher body weight observed in I_1 compared to I_0 females reared under standard postnatal conditions, whereas BW of males remains unaffected to cold treatment during incubation. This suggests a sex-specific response 342 of females to cold incubation temperature that would enhance their growth during the 343 postnatal period. The underlying mechanisms explaining this sex-difference remain to be 344 elucidated.

345 Consistently with the higher body weight measured at slaughter age, our results showed + 0.4percentage points in breast meat yield in the groups having experienced postnatal cold 346 conditions. However, the increase in breast meat yields previously reported by Shinder et al. 347 348 (2011) following the same cold incubation treatment was not observed in the present experiment. This discrepancy could be due to different environmental rearing conditions of T_0 349 broilers in both experiments. The highest value of breast meat yield was obtained in female 350 351 groups, consistent with previous results (Collin et al., 2007). Breast meat of female also exhibited higher pHu, probably resulting of a lower muscle energy (glycogen) store (Le 352 Bihan-Duval et al., 2008). This result is in accordance with the higher L* values observed in 353 males than in females, this parameter having been reported to negatively correlate with pHu. 354 Unexpected lower values of the fillet redness a* were observed in I₀T₁ group compared to 355 356 other groups reared in standard postnatal conditions. The redness of the breast meat was previously associated to chicken behavior before slaughter and early rate of post-mortem 357 glycolysis in muscle, the most active chickens producing the most red meat (Berri et al., 2005; 358 359 Chabault et al., 2012). Our study suggested a possible impact of postnatal cold on meat redness, probably in relation with a modification of the chicken metabolic rate (Collin et al., 360 2003) or of muscle fiber metabolic type switch towards more oxidative one as already 361 suggested by Ueda et al. (2004). 362

White striping is described as white striations parallel to muscle fibers that occur in both pectoral and thigh muscles (Kuttappan et al 2013b) and correspond to inclusion of lipids between fibers, affecting many criteria including color, tenderness, cooked processing and nutritional value of meat (Kuttappan et al 2013a). In the recent study of Clark et al. (2017),

broilers experiencing cyclical heat exposures during incubation were less prone to exhibit 367 368 moderate to severe myopathic defects compared to controls. In this case, the lower rate of myopathy was associated to lower breast meat yields. In the present study, cold stimulations 369 370 during incubation (I_1) did not affect the occurrence of white striping under control postnatal temperature. In contrast, combining cold stimulation during incubation with cold post-natal 371 experience (I_1T_1) resulted in greater scores of white striping in males that were also 372 373 characterized by the greatest body weights at d 40 compared to all T₀ males and all female groups. This result points out the importance of the perinatal thermal environment on the later 374 development of muscle, at least in males. According to Kuttappan et al. (2012), a high growth 375 376 rate of chickens over a short period leading to heavier broilers would trigger the appearance of non-infectious quality defects in broilers such as white striping. This could explain the higher 377 occurrence of WS of I_1T_1 males, but not the lower occurrence observed in other I_1 birds. In 378 379 ovo temperature alteration and rearing temperature of chicks may influence the embryonic muscle development with consequences on both metabolism and structure. Several authors 380 have shown that embryonic heat-stimulation during incubation modified adipogenic and 381 growth factor expression, myogenesis and cells proliferation (Halevy et al., 2006; Piestun et 382 al., 2009; 2011; Al-Musawi et al., 2012). We can hypothesize that the combined exposure to 383 384 acute cold at the end of incubation (d 18 and d 19), resulting in a drop of eggshell temperature by 5 to 11°C (data not shown) and to postnatal cold rearing temperature, may have enhanced 385 the proliferation of adipogenic precursors and further lipid deposition in the breast muscle. 386

Unexpectedly, the higher white striping scores observed in the I_1T_1 male group was not associated with higher values of intramuscular fat content (IMF). Indeed, it had been previously reported that chickens with moderate to severe white striping scores highlighted higher IMF percentage than non-affected chickens (Alnahhas et al., 2016). However, we showed low IMF percentage in breast muscles of females compared to males, in accordance with previous results of Zerehdaran et al. (2004). The authors related this difference to the lower body weight or muscle fatty acid metabolism of females. Consistently with variations of IMF, in which the liposoluble pigments accumulate, meat of males appeared more yellow than that of females. The meat of broilers that experienced cold postnatal conditions T_1 also appeared more yellow compared to control T_0 . Altogether, these results highlighted the role of early life thermal experience in determining breast muscle development and meat quality, especially in male broilers exhibiting a high growth rate.

Beyond the observed effects on the quality of the meat, our study indicates that cold challenge 399 in ovo and/or after hatch negatively affects animal welfare evaluated through the occurrence 400 401 of hock burns and pododermatitis at slaughter. The combined cold treatments I_1T_1 increased the occurrence of hock burns in males and of pododermatitis in both sexes, whereas under 402 standard postnatal temperature, these welfare parameters were not affected by incubation 403 temperature. It is likely that hock burns increase was in part due to the increase in body 404 weights observed in I₁T₁ males at 42 d. Indeed, Baéza et al. (2012) demonstrated a negative 405 406 impact of increased age and body weight on these types of injuries in relation with the lower walking ability measured in broilers. However, the greater body weight cannot explain by 407 itself the higher occurrence of pododermatitis observed in both I₁T₁ males and females 408 409 compared to other groups, except I_0T_1 . Variations in the occurrence of pododermatitis between groups may also be influenced by environmental rearing conditions such as litter 410 moisture, which was significantly greater in T_1 than in T_0 male pens, and intermediate in all 411 female pens (data not shown). Indeed, low litter moisture percentages, for example in case of 412 low crude protein content of the diet (Li et al., 2018), resulted in lower rate of footpad 413 414 dermatitis in broilers or in turkeys (Mayne et al., 2007; Tullo et al., 2017). It is therefore possible that the lower post hatching temperature and the greater animal density of female 415 416 pens (22 vs. 19 for males to account for the body weight differences between sexes from d 22

to 40 of age) have increased litter moisture and hence pododermatitis occurrence in T₁ males 417 418 and all female groups. This might be explained by a higher water consumption in less efficient birds in this group, which remains to be explored, or to the lower drying of the litter 419 420 due to the lower ambient temperature. These observations are also consistent with our previous study in which a greater occurrence of leg problems was reported in birds reared in 421 cages at low post hatching temperature (28°C instead of 33°C at d 0, Nyuiadzi et al., 2017). It 422 is also in accordance with the recent work of Steenfeldt et al. (2019) showing that cold 423 temperature (15°C) during the second half of the growth period resulted in a colder and wetter 424 litter, resulting in more footpad pododermatitis than standard temperature (21°C) in fast-425 426 growing chickens. Nevertheless, the present study pointed to potential limitations of short and acute cold exposures at the end of incubation when chickens were later reared under postnatal 427 cold temperatures. These limitations could be relative to the timing, intensity, homogeneity 428 429 and duration of the treatment, possibly favoring the chicken muscle growth at the detriment of other health-related biological functions. 430

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432 Based on these results, it appears that short cold stimulations at the end of incubation has positive effects on growth in females, and no deleterious effects on feed efficiency, muscle 433 434 white striping and pododermatitis under normal postnatal temperatures. Despite the positive 435 effects of cold stimulations during embryonic incubation combined to low temperature at start on performance and feed efficiency in broiler chickens, the present study also revealed the 436 437 limits of such strategy to improve bird adaptive capacities to cold environment. Indeed, thermal manipulation during embryogenesis and rearing period have altered some aspects of 438 welfare and meat quality, with a recrudescence of hock burns, pododermatitis and white 439 440 striping defects in males exposed to cold both before and after hatching. Our results also confirmed that early management affects in the long term both animal welfare and product 441

quality, two important components of the image and competitiveness of the poultry production systems. It is now necessary to understand the cellular, molecular and pathological disturbances involved in the observed changes to propose early management strategies allowing efficient and sustainable production systems, limiting current meat defects in broilers and improving welfare.

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Table 1. Hatchability of fertile eggs, sex-ratio, quality score of chicks and body temperature at hatch depending on incubation temperature.

| | Incubation te | mperature ¹ | |
|---|-------------------------|------------------------|--------------------|
| Traits ² | I ₀ | I ₁ | P-value |
| Rate of hatched chicks at 480h of incubation, % of fertile eggs | <mark>43</mark> | <mark>36</mark> | <mark>0.007</mark> |
| Hatchability, % of fertile eggs | <mark>95.4</mark> | <mark>95.7</mark> | <mark>0.801</mark> |
| Male chick percentage, % | <mark>52.8</mark> | <mark>51.9</mark> | <mark>0.765</mark> |
| Total chick quality score ³ (/100) | <mark>93.3 ± 1.3</mark> | 94.3 ± 0.5 | <mark>0.415</mark> |
| Body weight at hatch, g | 42.7 ± 0.1 | 42.6 ± 0.1 | <mark>0.681</mark> |
| Body temperature at hatch, °C | 38.8 ± 0.1 | 39.0 ± 0.1 | <mark>0.059</mark> |

 $^{-1}$ Control incubation I₀: eggs were incubated until hatch at 37.6°C and 56% RH; I₁: eggs were incubated in the same conditions but exposed for 30 min to 15°C at d 18 and d 19 of incubation.

⁶⁶² ²For each parameter, values are presented as mean \pm standard error. Hatchability of fertile eggs was calculated for n = 581 I₀ and 576 I₁ fertile eggs. Body weight and body temperature were measured on n=548 and n = 50 per incubation treatment, respectively.

³Chick quality was calculated according to Tona et al. (2003).

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Table 2. Chicken performance and meat quality parameters and yields depending on incubation and postnatal rearing temperatures and sex at
 slaughter age, and measured on meat cut 24h postmortem¹.

| Factors ¹ | Incubation ten | nperature ² (I) | | Postnatal temperature ³ (P) | | | Sex (S) | | | <i>P</i> -values of interactions | | | ons |
|--|----------------------------|----------------------------|--------------------|--|-------------------------|------------------------|-------------------------|-------------------------|------------------------|----------------------------------|--------------------|--------------------|--------------------|
| | I ₀ | I ₁ | P-value | T ₀ | T ₁ | P-value | Male | Female | P-value | <mark>I×P</mark> | <mark>I×S</mark> | <mark>P×S</mark> | I×P×S |
| Performance and litter parameters ⁴ | | | | | | | | | | | | | |
| d 21 BW (g) | <mark>851±4</mark> | <mark>863±4</mark> | <mark>0.026</mark> | <mark>847±4</mark> | <mark>868±4</mark> | <mark><0.001</mark> | <mark>901±4</mark> | <mark>822±4</mark> | <mark><0.001</mark> | <mark>0.708</mark> | <mark>0.856</mark> | <mark>0.513</mark> | <mark>0.008</mark> |
| d 40 BW (g) | <mark>2500±16</mark> | <mark>2537±16</mark> | <mark>0.034</mark> | <mark>2462±16</mark> | <mark>2574±16</mark> | <mark><0.001</mark> | <mark>2740±14</mark> | 2337±11 | <mark><0.001</mark> | <mark>0.547</mark> | <mark>0.542</mark> | <mark>0.896</mark> | <mark>0.038</mark> |
| d 0 - d 21 FCR (g/g) | 1.38±0.01 | 1.38±0.01 | <mark>0.504</mark> | <mark>1.36±0.00β</mark> | <mark>1.39±0.00α</mark> | <mark><0.001</mark> | 1.37±0.00b | <mark>1.39±0.01a</mark> | <mark><0.001</mark> | <mark>0.940</mark> | <mark>0.076</mark> | <mark>0.050</mark> | <mark>0.490</mark> |
| d 22 - d 40 FCR (g/g) | 1.82±0.02 | 1.82 ± 0.01 | <mark>0.864</mark> | 1.85±0.01 | 1.79±0.01 | <mark><0.001</mark> | <mark>1.80±0.01b</mark> | <mark>1.84±0.01a</mark> | <mark>0.001</mark> | <mark>0.042</mark> | <mark>0.173</mark> | <mark>0.653</mark> | <mark>0.231</mark> |
| d 0 - d 40 FCR (g/g) | 1.65±0.01 | 1.66±0.01 | <mark>0.678</mark> | <mark>1.66±0.01α</mark> | <mark>1.64±0.01β</mark> | <mark>0.008</mark> | <mark>1.64±0.01b</mark> | <mark>1.67±0.01a</mark> | <mark>0.002</mark> | <mark>0.065</mark> | <mark>0.075</mark> | <mark>0.979</mark> | <mark>0.412</mark> |
| Litter moisture (%) | <mark>39.0±0.7</mark> | <mark>38.9±0.8</mark> | <mark>0.926</mark> | <mark>37.9±0.7</mark> | <mark>40.0±0.8</mark> | <mark>0.047</mark> | <mark>38.9±0.8</mark> | <mark>39.0±0.8</mark> | <mark>0.897</mark> | <mark>0.403</mark> | <mark>0.429</mark> | <mark>0.037</mark> | <mark>0.570</mark> |
| Meat yields and parameters of ca | rcass quality ⁵ | | | | | | | | | | | | |
| Breast meat yield (%) | 20.7±0.1 | <mark>20.7±0.1</mark> | <mark>0.838</mark> | <mark>20.5±0.1β</mark> | <mark>20.9±0.1α</mark> | <mark>0.033</mark> | <mark>20.2±0.1b</mark> | <mark>21.1±0.1a</mark> | <mark><0.001</mark> | <mark>0.978</mark> | <mark>0.960</mark> | <mark>0.547</mark> | <mark>0.609</mark> |
| Thigh Yield (%) | 22.1±0.1 | 22.1±0.1 | <mark>0.911</mark> | <mark>22.0±0.0</mark> | <mark>22.2±0.0</mark> | <mark>0.213</mark> | 22.5±0.0b | <mark>21.8±0.0a</mark> | <mark><0.001</mark> | <mark>0.240</mark> | <mark>0.413</mark> | <mark>0.945</mark> | <mark>0.902</mark> |
| Abdominal Fat (%) | <mark>2.2±0.0</mark> | 2.2 ± 0.0 | <mark>0.236</mark> | <mark>2.2±0.0</mark> | 2.2 ± 0.0 | <mark>0.155</mark> | 2.1±0.0b | <mark>2.3±0.0a</mark> | <mark><0.001</mark> | <mark>0.968</mark> | <mark>0.861</mark> | <mark>0.618</mark> | <mark>0.461</mark> |
| Processing yield (%) | 82.3±0.5 | 82.5±0.5 | <mark>0.855</mark> | 82.1±0.6 | <mark>82.8±0.5</mark> | <mark>0.281</mark> | <mark>80.6±0.5b</mark> | <mark>83.9±0.5a</mark> | <mark><0.001</mark> | <mark>0.989</mark> | <mark>0311</mark> | <mark>0.978</mark> | <mark>0.765</mark> |
| Ultimate pH | 5.84±0.01 | 5.83±0.01 | <mark>0.882</mark> | <mark>5.84±0.01</mark> | <mark>5.83±0.01</mark> | <mark>0.609</mark> | <mark>5.80±0.01b</mark> | <mark>5.87±0.01a</mark> | <mark><0.001</mark> | <mark>0.147</mark> | <mark>0.696</mark> | <mark>0.289</mark> | <mark>0.192</mark> |
| Yellowness, b* | 10.2 ± 0.1 | 10.2±0.1 | <mark>0.733</mark> | 10.0±0.1β | 10.4±0.1α | <mark>0.014</mark> | 10.70±0.11a | <mark>9.77±0.11b</mark> | <mark><0.001</mark> | <mark>0.159</mark> | 0.277 | <mark>0.664</mark> | <mark>0.302</mark> |
| Redness a* | -0.38±0.05 | -0.39±0.05 | <mark>0.808</mark> | -0.50±0.05 | -0.27±0.05 | <mark>0.003</mark> | <mark>-0.32±0.06</mark> | -0.45±0.05 | <mark>0.092</mark> | <mark>0.038</mark> | <mark>0.525</mark> | <mark>0.629</mark> | <mark>0.165</mark> |
| Lightness L* | <mark>50.9±0.2</mark> | <mark>50.8±0.3</mark> | <mark>0.825</mark> | <mark>50.8±0.3</mark> | <mark>50.9±0.2</mark> | <mark>0.944</mark> | <mark>51.2±0.3a</mark> | <mark>50.5±0.2b</mark> | <mark>0.026</mark> | <mark>0.313</mark> | <mark>0756</mark> | <mark>0.674</mark> | <mark>0.319</mark> |
| Fillet intramuscular lipids (%) | 1.17±0.09 | 1.12±0.08 | <mark>0.298</mark> | 1.11±0.07 | 1.18±0.10 | <mark>0.576</mark> | 1.42±0.10 | <mark>0.92±0.04</mark> | <mark><0.001</mark> | <mark>0.853</mark> | <mark>0.018</mark> | <mark>0.548</mark> | <mark>0.167</mark> |

⁶⁷² ¹ Statistical effects of Sex (S), Postnatal temperature (P) and Incubation temperature (I) and of their interactions were analyzed by ANOVA. a, b: different letters correspond

673 to significant differences (P < 0.05) between sexes and α , β : different letters correspond to significant differences (P < 0.05) between postnatal temperatures.

² Control incubation I_0 : eggs were incubated until hatch at 37.6°C and 56% RH; I_1 : eggs were incubated in the same conditions but exposed for 30 min at 15°C at d 18 and d 19 of incubation.

676 ³ Control temperature T_0 from 32°C at d 0 to 21°C at d 21 or Cold temperature T_1 from 29°C at d 0 to 21°C at d 21, then T_0 and T_1 were maintained at 21°C from d 22 to d 40.

 4 Body weights (BW) were measured on n = 92 to 108 chickens per group at d 21 and 76 to 88 at d 40, and for Feed conversion ratio (FCR) and litter moisture percentage,

678 values are presented as mean \pm standard error (n = 4 per group).

⁵ Parameters were measured on n = 29 to 35, except for processing yield measured on n = 11 to 13, and for Fillet intramuscular lipid % measured on n = 6 to 9.

681 **Table 3**. Performance of chickens and meat quality parameters depending on incubation and postnatal rearing temperatures and sex when at least

- one interaction is significant between these factors. Body weights were measured at d 21 and d 40 of age. Feed conversion ratio (FCR) was
- calculated between d 22 and d 40.

| Sex (S) | | M | ales | | | Fe | emales | | | | | | | | |
|--|-------------------------------|----------------------------------|---|-----------------------------|--------------------------------------|---------------------------------------|---|-----------------------|--------------------|------------------------|------------------------|--------------------|--------------------|--------------------|--------------------|
| Postnatal temperature ¹ (P) | T ₀ T ₁ | | T ₀ | | T _i | | Statistical et | | | effects ³ | | | | | |
| Incubation temperature ² (I) | I _o | I. | Io | I. | I ₀ | I ₁ | | I ₁ | I | P | S | <mark>I×P</mark> | <mark>I×S</mark> | <mark>P×S</mark> | I×P×S |
| <mark>d 21 BW (g)</mark> | $\frac{892 \pm 8^{b}}{2}$ | <mark>893 ± 9^b</mark> | $\frac{896 \pm 7^{b}}{2}$ | 920 ± 7^{a} | 797 ± 7^{d} | <mark>823 ± 7°</mark> | $\frac{835 \pm 6^{\circ}}{2}$ | $830 \pm 6^{\circ}$ | <mark>0.026</mark> | <mark><0.001</mark> | <mark><0.001</mark> | <mark>0.708</mark> | <mark>0.856</mark> | <mark>0.513</mark> | <mark>0.008</mark> |
| <mark>d 40 BW (g)</mark> | 2685 ± 31^{b} | $\frac{2686 \pm 27^{b}}{2}$ | $\frac{2769 \pm 24^{a}}{2769 \pm 24^{a}}$ | $\frac{2818 \pm 27^{a}}{2}$ | $\frac{2240 \pm 18^{\text{e}}}{100}$ | 2330 ± 23^{d} | $\frac{2388 \pm 19^{\circ}}{2388 \pm 10^{\circ}}$ | $2390 \pm 21^{\circ}$ | <mark>0.034</mark> | <mark><0.001</mark> | <mark><0.001</mark> | <mark>0.547</mark> | <mark>0.542</mark> | <mark>0.896</mark> | <mark>0.038</mark> |
| d 22 - d 40 FCR (g/g) | $\underline{1.85\pm0.03}$ | $\underline{1.80\pm0.01}$ | 1.76 ± 0.02 | 1.78 ± 0.01 | 1.87 ± 0.01 | 1.88 ±0.01 | 1.79 ± 0.02 | 1.82 ± 0.02 | <mark>0.864</mark> | <mark><0.001</mark> | <mark>0.001</mark> | <mark>0.042</mark> | <mark>0.173</mark> | <mark>0.653</mark> | <mark>0.231</mark> |
| Litter moisture (%) | 37.0 ± 1.1 | 36.4 ± 1.0 | $\frac{41.6 \pm 1.3}{1.0}$ | 40.5 ± 1.2 | 38.0 ± 1.1 | $\frac{40.1 \pm 1.8}{1.8}$ | 39.3 ± 1.6 | 38.6 ± 2.2 | <mark>0.926</mark> | <mark>0.047</mark> | <mark>0.897</mark> | <mark>0.403</mark> | <mark>0.429</mark> | <mark>0.037</mark> | <mark>0.570</mark> |
| Redness a* | -0.45 ± 0.12 | -0.37 ± 0.12 | -0.22 ± 0.09 | -0.24 ± 0.13 | -0.66 ± 0.10 | -0.47 ± 0.10 | -0.16 ± 0.08 | -0.47 ± 0.08 | <mark>0.808</mark> | <mark>0.003</mark> | <mark>0.092</mark> | <mark>0.038</mark> | <mark>0.525</mark> | <mark>0.629</mark> | <mark>0.165</mark> |
| Fillet intramuscular lipids (%) | 1.63 ± 012 | 1.13 ± 0.09 | 1.59 ± 0.23 | 1.40 ± 0.27 | 0.80 ± 0.05 | $\frac{1.06 \pm 0.10}{1.06 \pm 0.10}$ | 0.92 ± 0.06 | 0.93 ± 0.12 | <mark>0.298</mark> | <mark>0.576</mark> | <mark><0.001</mark> | <mark>0.853</mark> | <mark>0.018</mark> | <mark>0.548</mark> | <mark>0.167</mark> |

 1 Control temperature T₀ from 32°C at d 0 to 21°C at d 21 or Cold temperature T₁ from 29°C at d 0 to 21°C at d 21, then T₀ and T₁ were maintained at 21°C from d 22 to d 40.

 2 Control incubation I₀: eggs were incubated until hatch at 37.6°C and 56% RH; I₁: eggs were incubated in the same conditions but exposed for 30 min at 15°C at d 18 and d

686 19 of incubation.

687 ³ Statistical effects of Sex (S), Postnatal temperature (P) and Incubation temperature (I) and of their interactions were analyzed by ANOVA.

688 For FCR and litter moisture, values are presented as mean \pm standard error (n = 4 per group).

689 ^{a,b,c,d} Different letters correspond to significant differences (P < 0.05) between treatments groups.

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- 691
- 692

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FIGURE LEGENDS

Figure 1. Experimental design. Number of birds per treatment are reported in the Materialsand Methods section.

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697 Figure 2. Effects of sex, incubation condition and postnatal rearing condition on the occurrence of white striping on the breast meat observed 24 h after slaughter at d 40. Control 698 incubation I₀ corresponds to egg incubation at 37.6°C, 56% RH. Cold incubation I₁ 699 700 corresponds to egg exposure for 30 min at 15°C at d 18 and 19 of incubation, eggs being incubated in the same conditions than for I₀ the remaining time. Chickens were reared in floor 701 pens either under control temperature T₀ from 32°C at d 0 to 21°C at d 21 or under cold 702 temperature T₁ from 29°C at d 0 to 21°C at d 21. From d 22 to d 40, the ambient temperature 703 in T₀ and T₁ was maintained at 21°C. Different letters (a, b, c) correspond to significant 704 705 differences (P < 0.05) between groups.

706

707 Figure 3. Effects of sex, incubation condition and postnatal rearing condition on parameters related to welfare troubles observed on carcasses after slaughter at d 40. A- Occurrence of 708 hock burns. B- Occurrence of pododermatitis. Control incubation I₀ corresponds to egg 709 incubation at 37.6°C, 56% RH. Cold incubation I₁ corresponds to egg exposure for 30 min at 710 15° C at d 18 and 19 of incubation, eggs being incubated in the same conditions than for I₀ the 711 remaining time. Chickens were reared in floor pens either under control temperature T₀ from 712 32°C at d 0 to 21°C at d 21 or under cold temperature T₁ from 29°C at d 0 to 21°C at d 21. 713 From d 22 to d 40, the ambient temperature in T₀ and T₁ was maintained at 21°C. Different 714 letters (a, b, c) correspond to significant differences (P < 0.05) between groups. 715

| Incubation groups | Control Incubation (I ₀) 37,6°C and 56% Relative Humidity (RH) | Cold Incubation (I_1) 37,6°C and 56% RH and at d 18 and 19 of incubation, 15°C, 81% RH 30 min/d | | | | | |
|----------------------|---|--|--|--|--|--|--|
| Postnatal | Control Incubation Control temperature $(I_0 T_0)$ $32^{\circ}C d 0 \rightarrow 21^{\circ}C d 21$ | Cold incubation Control temperature $(\mathbf{I}_1 \mathbf{T}_0)$ $32^{\circ}C d 0 \rightarrow 21^{\circ}C d 21$ | | | | | |
| groups | Control Incubation Cold temperature $(I_0 T_1)$ 29°C d 0 \rightarrow 21°C d 21 | Cold incubation Cold temperature $(I_1 T_1)$ 29°C d 0 \rightarrow 21°C d 21 | | | | | |

Figure 1

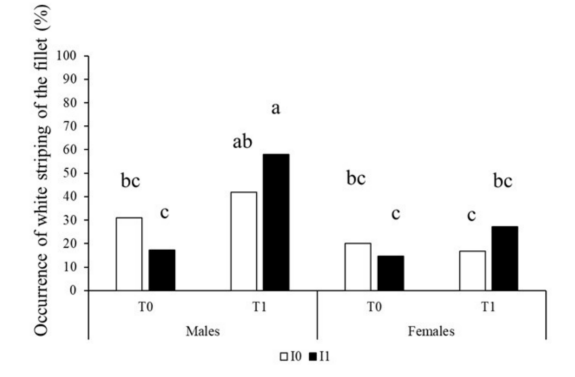


Figure 2

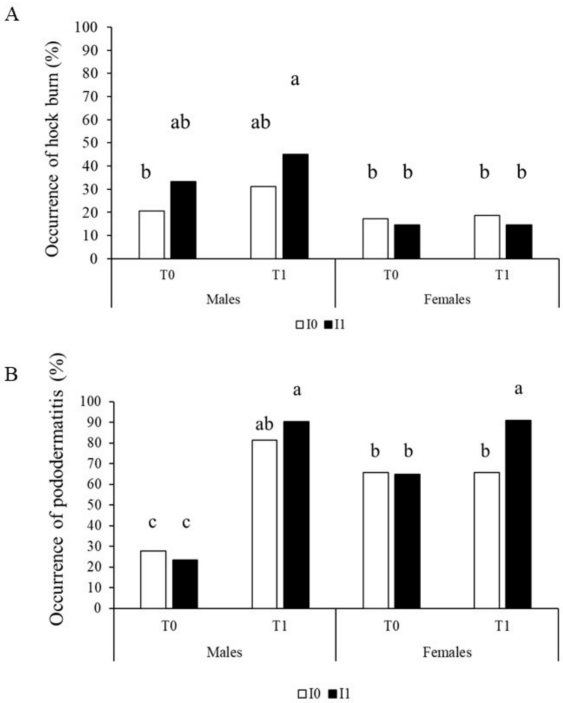


Figure 3