

Influence of cell history on the subsequent inactivation of campylobacter jejuni during cold storage under modified atmosphere

Benjamin Duque, Nabila Haddad, Albert Rossero, Jeanne Marie Membre, Sandrine Guillou

▶ To cite this version:

Benjamin Duque, Nabila Haddad, Albert Rossero, Jeanne Marie Membre, Sandrine Guillou. Influence of cell history on the subsequent inactivation of campylobacter jejuni during cold storage under modified atmosphere. Food Microbiology, 2019, 84, pp.1-8. 10.1016/j.fm.2019.103263. hal-02617961

HAL Id: hal-02617961 https://hal.inrae.fr/hal-02617961v1

Submitted on 26 Oct 2021

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



Distributed under a Creative Commons Attribution - NonCommercial 4.0 International License

Version of Record: https://www.sciencedirect.com/science/article/pii/S0740002019301431 Manuscript_00f4bd8465f1c045b50cdffcc542a0e4

1 Influence of cell history on the subsequent inactivation of Campylobacter jejuni during

2 cold storage under modified atmosphere

- 3
- 4 Benjamin Duqué, Nabila Haddad, Albert Rossero, Jeanne-Marie Membré and Sandrine
- 5 Guillou*
- 6 SECALIM, INRA, Oniris, Université Bretagne Loire, 44307, Nantes, France
- 7 *Corresponding author. SECALIM, INRA, Oniris, Université Bretagne Loire, 44307, Nantes,
- 8 France. Tel.: +33 240 687 763
- 9 *E-mail address*: sandrine.guillou@oniris-nantes.fr (S. Guillou)
- 10

11 Abstract

12 Worldwide, *Campylobacter* infections are the main cause of human bacterial enteritis 13 and broiler meat is considered as the most important source of human campylobacteriosis. 14 Some mitigation strategies have been focused on reduction of *Campylobacter* at the 15 slaughtering steps. This study aimed to determine the influence of consecutive stresses 16 inspired by slaughtering steps on the subsequent inactivation of Campylobacter jejuni during 17 cold storage under different modified atmospheres. Using a full experimental design, three 18 strains of *C. jejuni* of poultry origin were submitted to consecutive heat (46°, 50° or 54°C for 4 19 min) and cold (-4° or 3°C for 2 h) stresses by plunging cultures into baths at appropriate 20 temperatures. Cultures were then stored at 6°C during seven days under modified 21 atmospheres (70% O_2 / 30% CO_2 or 50% CO_2 / 50% N_2). Inactivation of *C. jejuni* induced by 22 cold storage was shown to depend significantly (P<0.0001) upon the heat stress previously 23 applied. It was shown to be the highest under the atmosphere enriched in oxygen, after 24 application of 54°C. Strain inactivation variability was also quantified. These results show that 25 consecutive stresses influence further inactivation of C. jejuni during storage and 26 consequently the contamination level at consumer's plate. 27 28 **Keywords:** foodborne pathogen; slaughter process; stresses; food safety; strain 29 variability

31 1. Introduction

Since 2005, *Campylobacter* has been the main cause of bacterial enteritis worldwide in humans (EFSA and EDC 2018). Despite being largely under-estimated, the number of reported confirmed cases of *Campylobacteriosis* was 246,158 in 2017 with an EU notification rate of 64.8 per 100,000 population (EFSA and EDC 2018). Infection with *C. jejuni* can lead to chronic sequelae, such as Guillain-Barré syndrome (GBS), characterized by the damage of the peripheral nervous system leading to a reversible neuromuscular paralysis (Nyati and Nyati 2013).

The main reservoirs for *C. jejuni* are avian species and farmed poultry (Young et al. 2007). In food, the occurrence of *Campylobacter* remains high in broiler meat (EFSA and EDC 2018) and it is considered to be the most important single source of human *Campylobacteriosis*. In 2017, 37.4% of the 13,445 samples of fresh broiler meat (single or batch, aggregated data from all sampling stages) were found to be *Campylobacter* positive (EFSA and EDC 2018).

44 Several microbiological risk assessment studies have been performed to assess the 45 potential effects of control measures on prevention/reduction of Campylobacter concentration 46 in broiler meat production. Some studies focused more on pre-slaughter stages associated 47 with broiler rearing (Hermans et al. 2011, Lin 2009) including the use of probiotics (Saint-Cyr 48 et al. 2017) or vaccine (Meunier et al. 2017). Another effective strategy to reduce the number 49 of campylobacteriosis cases may reside in limiting the entry of chicken carcasses highly 50 contaminated by Campylobacter into the market. In this context, the European Commission 51 has just set a microbiological process hygiene criterion for Campylobacter in broiler carcasses (European Commision 2017). A limit of 1000 CFU.g⁻¹ on carcasses after chilling 52 53 has been defined. The EFSA (European Food Safety Authority) determined that more than 54 50% human risk reduction could be achieved if broiler carcasses complied with this new microbiological criteria of 1000 CFU.g⁻¹ (European Commision 2017). For food producers, 55 56 the compliance with this hygienic criterion requires improvements in slaughter hygiene to 57 limit cross-contamination but also investigation of new control measures which may favor

Campylobacter inactivation following slaughter. Focusing on the susceptibility of the bacteria
to the processing steps may represent one way to tackle *Campylobacter* risk.

60 C. jejuni is able to survive to slaughterhouse environments and during poultry 61 processing. In the poultry farm-to-fork chain, several steps may be stressful for bacteria 62 regarding its growth requirements (microaerophily and thermotolerance), such as scalding, 63 chilling and storage. Scalding consists in immersing chicken carcasses into a hot water bath, 64 which may induce heat stress to C. jejuni. At the end of the slaughter process, the chilling 65 step enables quick refrigeration of chicken carcasses, which is necessary to lower the growth 66 rate of pathogenic and spoilage microorganisms (James et al. 2006), is supposed to 67 generate cold stress to C. jejuni. At last, chicken cuts are generally conditioned under 68 modified atmosphere and stored under chilled temperature until consumption. Depending on 69 the producer, the gas mix of the package may contain high concentrations of oxygen or not. 70 This latter step induces both cold and possibly oxidative stresses. However, if the influence 71 of single stresses on *Campylobacter* behavior is well-described, the effect of stresses 72 occurring sequentially still needs to be more deeply understood in order to point out 73 conditions of slaughter steps that favor *Campylobacter* inactivation during poultry products 74 storage.

75 Indeed, the exposure to a stress may condition the bacterial behavior following the exposure 76 to another stress. Increased resistance has been for example already reported in Bacillus 77 (den Besten et al. 2010, Desriac et al. 2013). For instance, after being subjected to mild acid 78 or oxidative stress, C. jejuni appeared to be more resistant to a subsequent lethal stress 79 (Murphy et al. 2003a, Murphy et al. 2003b, Oh et al. 2015). Such cross-protection response 80 from one stress to another has also been reported, like genes involved in heat shock 81 response in C. jejuni NCTC 11168 which were upregulated in response to acid stress (Reid 82 et al. 2008b). Additionally, starved *C. jejuni* cells were able to better withstand heat stress 83 (Klančnik et al. 2009).

84 Besides, it is well established that strains have different behavior when subjected to 85 stress (Chan et al. 2001, Cools et al. 2003, Newell et al. 2001) and this is highly relevant for

- 86 poultry product in the sense that poultry meat is naturally contaminated by different strains of
- 87 *C. jejuni* and often more than one species (Colles et al. 2003, Johnsen et al. 2006, Rivoal et

88 al. 2005)

- 89 In the current study, several strains of *C. jejuni* were submitted to stresses inspired from
- 90 those encountered during slaughter, and subsequently incubated under modified
- 91 atmosphere. The effect of cell history *i.e.* the effect of hot and cold temperature applied
- 92 following successive heat and cold stresses, on the subsequent inactivation of *C. jejuni*
- 93 strains during chilled storage under various modified atmospheres, was then quantitatively
- 94 analyzed.
- 95

96 2. Materials and methods

97

2.1 Bacterial strains and culture conditions

Experiments were performed on three C. jejuni strains isolated from poultry products. 98 99 C. jejuni C09MJLT205 was isolated from chicken legs distributed at the market level 100 (Guyard-Nicodeme et al. 2013). C. jejuni RM1221 was isolated by Fouts et al. (2005). The 101 strain C. jejuni C97Anses640 was isolated from a poultry product (Guyard-Nicodeme et al. 102 2016). All strains were stored at -80°C in brain heart infusion (BHI, Biomérieux, Marcy 103 l'Etoile, France) supplemented with 20% (v/v) glycerol. C. jejuni strains were routinely 104 cultured on Karmali (Oxoid, Dardilly, France) agar plates at 42°C for 48-72 h under 105 microaerophilic conditions (5% O₂, 10% CO₂, and 85% N₂). Then, one or two colonies were 106 transferred into 20 mL of Mueller-Hinton broth (MH, Oxoid, Dardilly, France) and incubated 107 for 20 h under the same conditions under agitation (90 rpm). A 1/100th dilution was 108 transferred into a fresh MH broth and incubated under the same conditions for 18 h, so that 109 the culture reached stationary phase. Finally, a 1/10th dilution was carried out to obtain what 110 we called the 'starter culture' further used for challenge-test experiments.

To enumerate viable *C. jejuni* cells, suspensions were serially tenfold diluted and
surface plated on Columbia sheep blood agar plates (Biomérieux, Marcy l'Etoile, France) and
incubated for 48 h at 42°C under microaerophilic conditions using a SPIRAL plater
(EasySpiral Interscience, Saint Nom, France).

115

116

2.2 Experimental procedure

C. jejuni strains were submitted to stresses inspired from conditions encountered
during slaughter chicken process. Slaughter is associated with several steps considered as
stressful for *C. jejuni*. These steps are scalding, chilling, and are followed by cold storage
under modified atmosphere. Scalding consists in immersing chicken carcasses into hot water
bath. Then, chicken carcasses are submitted to a guick refrigeration during the chilling step.

At last, after cutting carcasses in pieces, chicken cuts are conditioned under modifiedatmosphere and stored under chilled temperature.

124 The three previously mentioned stresses were reproduced as closely as possible in 125 the laboratory, and named as heat stress, cold stress and storage stress, respectively. The 126 first two stresses, *i.e.* heat and cold stresses were assumed to influence the future behavior 127 of *C. jejuni* during chilled storage under modified atmosphere. As such, they were considered 128 as participating to *C. jejuni* adaptive history (Fig. 1).

129 Ten mL of starter culture with an initial concentration of 8 log₁₀ CFU.mL⁻¹ from each strain

130 were transferred into a glass tube and submitted to consecutive heat, cold temperature and

131 storage at 6°C under modified atmospheres according to a full experimental design.

Heat stress consisted in immersing *C. jejuni* cultures in hot water baths at 46°, 50° or 54°C for 4 min. 54°C was chosen as the maximal observed temperature of scalding baths in French chicken slaughterhouses, 46 °C, the maximum growth temperature (Hazeleger et al. 1998) of *C. jejuni* and 50°C, an intermediate temperature.

After exposure to heat stress, tubes were immediately cooled for 5 min in a water bath
at 22°C, corresponding to the conditions of ambient temperature occurring during the five
minutes between scalding and chilling steps in slaughterhouse.

Cold stress was applied immediately after cooling by plunging the previous cultures into cold water with ethylene glycol at -4° or 3°C for 2 h. The temperature of 3°C was chosen because it represents the targeted core temperature of poultry carcasses during the chilling step and -4°C is the minimal surface temperature measured during different visits performed in several slaughterhouses. Strains were exposed to cold stress for 2 h similarly as chicken carcasses during the chilling step.

At last, cultures were stored at 6°C for seven days under the modified atmospheres
(70% O₂ / 30% CO₂ or 50% N₂ / 50% CO₂) commonly used for the packaging of poultry cuts.
These two atmospheres were generated by filling jars with the *ad hoc* gas mixture (Air
Liquide, Carquefou, France). Cultures were placed in bottles with porous silicone caps
(Hirschmann Laborgeräte GmbH & Co. KG, Eberstadt, Allemagne) enabling gas exchange.

The temperature of 6°C was chosen according to the average temperature of consumer
refrigerators (Laguerre et al. 2001).

During experiments, the temperature applied during heat and cold stress was
recorded using a temperature probe Kistock KTT220 (KIMO, Montpon, France) placed in a
control tube. The gas composition (% O₂ and % CO₂) of the jar atmosphere was measured
using the Oxybaby M+ device (WITT-GASETECHNIK GmbH & Co KG, Witten, Germany). It
was performed just after filling with gas mixture and regularly over time to check the stability
of the gas composition.
After each step, viable counts of *C. jejuni* were enumerated to determine the viability loss

resulting from the application of each step, *i.e.* $\Delta \log_{10}$. A total of 152 (3 strains x 4 repetitions

160 x 12 experimental conditions + 8 controls) experiments were performed.

161

162 *2.3 Statistical analysis*

Four independent replicates were carried out and statistical analysis was performed on $\Delta \log_{10}$ transformed values. An ANOVA (α =5%) was performed using the software XLSTAT (version 19.03.45028), an Addin of the Microsoft Excel software.

167 **3. Results**

168 169 3.1 Cumulative effect of subsequent heat, cold and storage stresses on inactivation of C. jejuni (Δlog_{total})

From the starter culture with an initial concentration of 8 log₁₀ CFU.mL⁻¹, the mean
cumulative *C. jejuni* inactivation, due to the consecutive heat, cold and storage stresses,
varied from 0.02 to 4.9 log₁₀ CFU.mL⁻¹ as function of strains and atmosphere conditions.
Detailed results are available in supplementary material Table S1.

174 During experiments, controls were performed and submitted only to the cold storage 175 under modified atmosphere. The inactivation induced by the only storage step varied from 176 0.9 to $1.69 \log_{10} CFU.mL^{-1}$ with the $70\%O_2 / 30\% CO_2$ atmosphere and from 0.1 to $0.3 \log_{10}$ 177 $CFU.mL^{-1}$ with the $50\% N_2 / 50\% CO_2$ atmosphere.

178 For the bacterial samples undergoing all steps, the storage step with the $70\%O_2$ / 179 30% CO₂ atmosphere, had the main impact on *C. jejuni* inactivation, the bacterial population 180 decrease varied from 1.0 to 3.9 log₁₀ CFU.mL⁻¹ (Fig. 2a). In addition, the highest inactivation 181 during storage was reached after the application of the highest temperature during heat 182 stress. Heat only produced an inactivation following application of 54°C with a mean inactivation between 0.8 and 0.9 log₁₀ CFU.mL⁻¹, whereas the inactivation was negligible 183 184 when 50° and 46°C were applied. In our conditions, cold temperature had also a negligible 185 impact on the *C. jejuni* inactivation, indeed a maximum of 0.3 log₁₀ CFU.mL⁻¹ reduction was 186 achieved. The highest cumulative inactivation was obtained for the C97anses640 strain. This 187 strain appeared to be less resistant than C09MJLT205 and RM1221 strains, especially 188 during storage. For instance, the inactivation of strain C97anses640 at 50°C / -4°C was 3.3 log₁₀ CFU.mL⁻¹ (mean value) compared to 1.2 log₁₀ CFU.mL⁻¹ for both C09MJLT205 and 189 190 RM1221 strains (Fig. 2b).

191 Using the 50% N_2 / 50% CO₂ atmosphere, the cumulative inactivation of *C. jejuni* was 192 mainly attributable to the storage step and varied from 0.1 to 0.9 log₁₀ CFU.mL⁻¹ (Fig. 2b). 193 However, especially at 54°C, the heat stress contributed equally with the cold storage to the 194 cumulative inactivation, except for the strain RM1221. Indeed, for the latter, the inactivation
195 in the hot water bath was higher than during the storage step, *i.e.* mean values of 1.5 and 0.9
196 log₁₀ CFU.mL⁻¹, respectively.

197

3.2 Effect of heat/cold temperature, modified atmosphere, strains and interactions *on subsequent inactivation during storage affected by cell history (Δlog_{storage})*When subjected to mild stress, bacteria may become more resistant to another stress
applied subsequently. In this study, we tested if the application of successive heat and cold
temperature enhanced the resistance of *C. jejuni* during storage. In other words, we studied
the effect of cell adaptive story on the subsequent behavior of *C. jejuni*.

204 The influence of the different factors and their interaction on inactivation of C. jejuni 205 during cold storage was analysed by ANOVA. The ANOVA highlighted that the temperature 206 of the cold bath did not influence significantly the inactivation during storage. Therefore, a 207 new ANOVA was performed without considering cold stress. The model was significant (p < 208 0.0001) as indicated in Table 1. The ANOVA showed that there were significant effects of 209 strain, temperature applied during heat stress, and packaging atmosphere. During the cold storage, strains C09MJLT205 and RM1221 exhibited similar inactivation around 1.1 log10 210 211 CFU.mL⁻¹. In contrast, the strain C97anses640 appeared to be less resistant to storage than 212 the two other strains with a mean inactivation of 1.8 log10 CFU.mL⁻¹ (Fig. 3a).

213 Regarding heat stress, the highest temperature of 54°C induced the highest inactivation

associated with storage. Indeed, the mortality reached a mean of 1.9 log₁₀ CFU.mL⁻¹

compared to that obtained at 46° and 50°C (1 \log_{10} CFU.mL⁻¹) (Fig. 3b).

216 Inactivation of *C. jejuni* during storage was more than five times higher under the 217 modified atmosphere 70% O_2 / 30% CO_2 than under the atmosphere 50% N_2 / 50% CO_2 (p < 218 0.0001), with a mean inactivation of 2.2 log₁₀ CFU.mL⁻¹ and 0.4 log₁₀ CFU.mL⁻¹, respectively 219 (Fig. 3c). 220

The ANOVA showed significant interactions between strains and atmosphere,

temperature applied during heat stress and atmosphere.

222 While the mean inactivation resulting from storage under the atmosphere 50% N₂ / 50% CO₂ 223 was low and similar no matter the strain used, (between 0.3 and 0.5 \log_{10} CFU.mL⁻¹), the 224 inactivation was largely higher under 70% O₂ / 30% CO₂ and different between strains (p< 225 0.0001). Indeed, strain C97anses640 appeared to be less resistant than the strains 226 C09MJLT205 and RM1221. The mean inactivation of strain C97anses640 was 3.2 log₁₀ 227 CFU.mL⁻¹ compared to that of C09MJLT205 and RM1221 strains, *i.e.* 1.9 and 1.7 log₁₀ 228 CFU.mL⁻¹, respectively (Fig. 3d). Besides, this study pointed out that similar inactivation 229 during heat treatment does not necessarily lead to similar inactivation during storage. Indeed, 230 under 70% O_2 / 30% CO_2 atmosphere, the inactivation during heat stress was similar for 231 each strain but higher for the strain C97anses640 during the storage. Thus, this result 232 highlights the importance of the cell history, and the difficulty to predict future behavior while 233 considering only a previous phenotypic response to a specific stress.

A significant interaction between atmosphere and heat stress was characterized by a similar mean inactivation at 46° and 50°C under the atmosphere 70% O_2 / 30% CO_2 , (1.7 and 2.1 log₁₀ CFU.mL⁻¹, respectively), compared to that obtained at 54°C, *i.e.* 3.0 log₁₀ CFU.mL⁻¹ (Fig. 3e). Under the atmosphere 50% N₂ / 50% CO₂, the inactivation following application of 54°C was also the highest and was 0.8 log₁₀ CFU.mL⁻¹. In contrast, the inactivation resulting from application of 46°C was slightly higher than that obtained following application of 50°C, *i.e.* 0.4 and 0.1 log₁₀ CFU.mL⁻¹ respectively.

241

242 4. Discussion

The objective of this study was to assess the effect of the cell history on the subsequent survival of three *C. jejuni* strains. Stressful steps from slaughterhouses were identified and then reproduced in laboratory.

The first step was to immerse the bacteria in hot water bath at temperature ranging from 46° to 54°C. The inactivation during this step varied between 0 and 1.5 log₁₀CFU.mL⁻¹ 248 according to strain and temperature. From heat inactivation, it was possible to calculate the 249 decimal reduction time or D-value. These D-values corresponds to the time necessary to kill 250 90% of an initial population or decrease the initial population by one log. The D-value is a 251 common metric and enables to compare more easily results from different studies. In our 252 experimentations, heat stress was applied during 4 min. We measured that one minute was 253 the time necessary to reach the temperature desired. Thus, once reached, the set 254 temperature was maintained during 3 min. For a selected temperature, D-values were 255 calculated by dividing the 3-min constant temperature application by $\Delta \log$ lost following 256 application of the heat stress.

257 All collected D-values from different studies in the literature were gathered in Table 2. We 258 can see an important variability between results of decimal reduction times for C. jejuni, 259 mainly at 50°C. It appears that variability was higher at 50°C than at 55°C, which was also 260 mentioned by McCarthy et al. (2018). In our study, C. jejuni was submitted to heat stress in 261 suspension rather than on chicken muscle like in the real scalding step. However, the high 262 variability of collected D-values could not enable to visualize a difference of heat resistance 263 between meat matrices and laboratory media, as it could have been expected. Indeed, it has 264 been shown that the heat resistance was higher for Salmonella when attached to muscle as 265 opposed to free cells (Humphrey et al. 2007). C. jejuni cells could also be better recovered 266 from 5-min boiling if they were previously attached to chicken muscle as opposed to carrots (de Jong et al. 2012). D_{50°C}-values varied from 3.5 to 39 min which corresponds to an 267 inactivation from 0.08 to 0.9 log₁₀ in 3 min. D_{55°C}-values varied between 1.5 and 2.8 min and 268 269 were found in the range of what was observed in previous studies (mean $D_{55^{\circ}C}$ of 1.5 ± 1.3 270 min). In light of the variation illustrated in Table 2, heat strain sensitivity, experimental 271 method, and variables such as medium may greatly influence the inactivation of C. jejuni. In 272 our study, for 46°C and 50°C, no inactivation has been observed. Indeed, we considered that inactivation was significant if $\Delta \log$ was greater than 0.5 \log_{10} CFU.mL⁻¹ *i.e.* the experimental 273 274 error due to microbial enumeration (Pujol et al. 2012). These results are in accordance with 275 the very high D-values found by Nguyen et al. (2006).

276 To our knowledge the effect of cold temperature applied for a short period (*i.e.* two or 277 several hours) on C. jejuni has not yet been reported. In this study, application of a cold 278 temperature during two hours had no significant effect on the survival of *C. jejuni*. However, 279 these results have to be interpreted with caution, especially regarding possible extrapolation 280 with the chilling step of the slaughter process. Unlike our experimental conditions, the cold 281 temperature applied during the chilling step is generally associated with desiccation due to 282 ventilation. In laboratory, such conditions could not be mimicked. Nevertheless, the potential 283 influence of desiccation might induce microbial inactivation. For instance, Zoz et al. (2016) 284 have highlighted the impact of relative humidity on survival of *Listeria monocytogenes*. 285 Likewise, a study led by (Rivoal et al. 2016), investigated the relation between temperature, 286 chilling time and air velocity on the inactivation of eight strains of C. jejuni by the use of a 287 miniaturized chilling room prototype. This work highlighted the significant effect of the 288 duration of the chilling step on the inactivation of *C. jejuni*. A significant interaction between 289 the temperature and the air flow during the process was also pointed out by the authors. 290 However, it is important to reinforce here that carcasses contaminated with more than 3 log 291 CFU.g⁻¹ of Campylobacter could not be significantly decontaminated during the chilling step 292 in slaughtering house.

293 After the application of hot and cold temperature at the slaughterhouse, C. jejuni was 294 submitted to storage for several days under the same conditions as those found at retailers' 295 and then at consumers' home. During storage, C. jejuni undergoes a combination of cold 296 temperature and modified atmosphere. It was found in the current study that the inactivation 297 varied from -1.0 to -3.9 log₁₀ CFU.mL⁻¹ in the atmosphere 70% O₂ / 30% CO₂ and from 0 to -298 0.9 \log_{10} CFU.mL⁻¹ in the atmosphere 50% N₂ / 50% CO₂. The inactivation of *C. jejuni* was 299 significantly influenced by the atmosphere, with a higher effect of the atmosphere 70% O_2 / 300 30% CO₂. Indeed, *Campylobacter* are microaerophilic organisms. The presence of O₂ in high 301 concentrations is toxic for the pathogen. For many years, studies have investigated the effect 302 of storage during several days under aerobiosis condition or modified atmosphere. On 303 various meat matrices, the reduction of *Campylobacter* count in aerobiosis at 4°C varied from 304 0.3 to 2 log₁₀ CFU.g⁻¹ (Bhaduri and Cottrell 2004, Blankenship and Craven 1982, Eideh and 305 Al-Qadiri 2011, Koidis and Doyle 1983, Phebus et al. 1991, Vashin and Stoyanchev 2011). In 306 broth, similar results have also been found (Chaisowwong et al. 2012, Chan et al. 2001, 307 Garenaux et al. 2008). However, the inactivation of *C. jejuni* was higher in our work 308 compared to what is commonly observed in the literature. Indeed, before the storage step, 309 bacteria were subjected to heat and cold stress which may induce a hurdle effect. Besides, 310 one of the atmospheres used during our experiments had a concentration in O₂ much higher 311 than in air. In other studies, it has been found that the inactivation increased with 312 atmospheres enriched in O₂ compared to atmospheres containing only CO₂ or N₂. Boysen et 313 al. (2007) showed that in broth, the inactivation under atmosphere $70\% O_2 / 30\% CO_2$ was 314 higher than in atmosphere 70% N_2 / 30% CO_2 and 100% $N_2,$ with a reduction of 0.5 to 2 log_{10} 315 CFU.mL⁻¹ and 0.3 to 0.8 log₁₀ CFU.mL⁻¹ respectively.

Such results have also been found when *C. jejuni* was submitted to paraquat, a chemical
oxidizing agent (Garenaux et al. 2008). After 7 days of exposition, the population was
reduced by 3 log₁₀ CFU.mL⁻¹.

319 In this work, thermal steps were applied successively in order to assess the impact of 320 cell history on the subsequent inactivation of C. jejuni during storage. Some studies have 321 already investigated the potential effect of cell history or adaptation of one stress to another 322 stress. Indeed, microorganisms enduring stressful environments can protect or crossprotect 323 themselves to survive subsequent homogeneous or heterogeneous stresses, which is called 324 adaptive tolerance response (ATR). It has been shown that C. jejuni can induce an ATR after 325 aerobic or acid stress (Gaynor et al. 2005, Jones et al. 1993, Ma et al. 2008, Martinez-326 Rodriguez and Mackey 2005, Murphy et al. 2003a, Murphy et al. 2003b, Reid et al. 2008a). 327 On the other hand, in our study it has been shown that after successive application of heat 328 and cold temperature, if an ATR was induced, it was insufficient to produce an adaptation 329 effect since the inactivation was higher for stressed bacteria compared to unstressed ones. 330 This study was conducted on three strains isolated from poultry and more particularly

331 on product from retailed market. Tackling strain variability is not easy because the features of

332 the strains which are associated with highest or lowest sensitivity to processing steps are not 333 known. The choice was done in this study to select strains (i) representative of strains 334 commonly found in poultry, and (ii) exhibiting a priori different behaviors regarding selected 335 stresses (data not shown). Firstly, the representativeness of strains in poultry was sought by 336 determining the strain genotype by MultiLocus Sequence Typing (MLST). The strain 337 C97anses640 belonged to the Sequence Type-45 complex (ST-45), the strain C09MJLT205, 338 to the ST-21 complex and the strain RM121, to the ST-354 complex. The ST-21 and ST-45 339 are the two main clonal complexes encountered in chicken (and poultry meat) worldwide. 340 Interestingly, they are also commonly found among human isolates (Colles et al. 2003, de 341 Haan et al. 2010, Dingle et al. 2002, Guyard-Nicodeme et al. 2015, Habib et al. 2009, 342 Levesque et al. 2008, Ragimbeau et al. 2008, Sheppard et al. 2009). Secondly, the behavior 343 variability between strains was determined regarding their inactivation during cold storage 344 under modified atmosphere. High response variability between C. jejuni strains has also 345 been reported in various studies (Habib et al. 2010, Ligowska et al. 2011, Vashin and 346 Stoyanchev 2011). These findings highlighted the difficulty and the importance to take into 347 account strain variability in microbial inactivation studies, and, more generally in exposure 348 assessment.

The experimental laboratory conditions applied during the present study differ from what *C. jejuni* cells really experience during the slaughtering steps because they are not attached on muscle but free in liquid suspension, and because the chilling step does not induce desiccation. Here reside some limitations of laboratory experiments. They offer the possibility of replicating standardized procedures and measures, which is necessary to obtain the proof of concept. Passed this step, a validation step in laboratory on chicken muscle will be necessary and is currently being processed.

In conclusion, to better predict *Campylobacter* contamination in poultry processing,
it is essential to assess its behavior after applying not only one stress but the whole
sequence of stresses encountered during the process. Indeed, cell history plays an important
role and may induce some physiological responses such as ATR which influences *C. jejuni*

360 further behavior regarding subsequent stresses. In some cases, the bacteria is able to better 361 resist to another stress whereas sometimes, the application of successive stresses induces a 362 hurdle effect, and in contrast, enhances bacterial inactivation. Finally, this study pointed out 363 that there is no clear correlation between the amount of log inactivation observed in 364 preliminary processing steps and the amount of log inactivation occurring at the latest 365 processing steps such as storage, rendering it very difficult to predict the microbial behavior 366 during the storage at retail and consumer's house from phenotypic information resulting from 367 application of previous slaughtering steps.

A new generation of predictive models, including cell history information, will be necessary to progress along this route, and omics methods might help. These new tools consider not only the bacterial behavioral response, but also the adaptation response depending on the bacterial gene expression and regulation. The investigation of predictive models including gene expression would enable to fine-tune existing exposure assessment models and consider strain-dependent physiological response.

375 Acknowledgements

- We are grateful to the French Agency for Food, Environmental and Occupational Health &
- 377 Safety (ANSES) of Ploufragan who realized the MLST analysis.
- We also especially thank the Regions Pays de La Loire and Bretagne through the "Pôle
- 379 Agronomique Ouest" for their financial support in the framework of the Biomics project.

381 References

- Berrang, M.E. and Dickens, J.A., 2000. Presence and level of *Campylobacter spp*. on broiler
 carcasses throughout the processing plant. Applied Poultry Science 9, 43-47.
- Bhaduri, S. and Cottrell, B., 2004. Survival of cold-stressed *Campylobacter jejuni* on ground
 chicken and chicken skin during frozen storage. Applied and Environmental
 Microbiology 70, 7103-7109.
- Blankenship, L.C. and Craven, S.E., 1982. *Campylobacter jejuni* survival in chicken meat as
 a function of temperature. Applied and Environmental Microbiology 44, 88-92.
- Boysen, L., Knochel, S. and Rosenquist, H., 2007. Survival of *Campylobacter jejuni* in
 different gas mixtures. FEMS Microbiology Letters 266, 152-157.
- Chaisowwong, W., Kusumoto, A., Hashimoto, M., Harada, T., Maklon, K. and Kawamoto, K.,
 2012. Physiological Characterization of *Campylobacter jejuni* under Cold Stresses
 Conditions: Its Potential for Public Threat. Journal of Veterinary Medical Science 74,
 43-50.
- Chan, K.F., Le Tran, H., Kanenaka, R.Y. and Kathariou, S., 2001. Survival of Clinical and
 Poultry-Derived Isolates of *Campylobacter jejuni* at a Low Temperature (4 C). Applied
 and Environmental Microbiology 67, 4186-4191.
- Colles, F.M., Jones, K., Harding, R.M. and Maiden, M.C.J., 2003. Genetic Diversity of
 Campylobacter jejuni isolates from Farm Animals and the Farm Environment. Applied
 and Environmental Microbiology 69, 7409-7413.
- 401 Cools, I., Uyttendaele, M., Caro, C., D'Haese, E., Nelis, H.J. and Debevere, J., 2003.
 402 Survival of *Campylobacter jejuni* strains of different origin in drinking water. Journal of
 403 Applied Microbiology 94, 886-892.
- de Haan, C.P., Kivisto, R., Hakkinen, M., Rautelin, H. and Hanninen, M.L., 2010. Decreasing
 trend of overlapping multilocus sequence types between human and chicken *Campylobacter jejuni* isolates over a decade in Finland. Applied and Environmental
 Microbiology 76, 5228-5236.

de Jong, A.E., van Asselt, E.D., Zwietering, M.H., Nauta, M.J. and de Jonge, R., 2012.
Extreme Heat Resistance of Food Borne Pathogens *Campylobacter jejuni*, *Escherichia coli*, and *Salmonella typhimurium* on Chicken Breast Fillet during
Cooking. Int J Microbiol 2012, 196841.

den Besten, H.M., Arvind, A., Gaballo, H.M., Moezelaar, R., Zwietering, M.H. and Abee, T.,
2010. Short- and long-term biomarkers for bacterial robustness: a framework for
quantifying correlations between cellular indicators and adaptive behavior. PLoS One
5, e13746.

- Desriac, N., Broussolle, V., Postollec, F., Mathot, A.G., Sohier, D., Coroller, L. and
 Leguerinel, I., 2013. *Bacillus cereus* cell response upon exposure to acid
 environment: toward the identification of potential biomarkers. Frontiers in
 Microbiology 4, 284.
- Dingle, K.E., Colles, F.M., Ure, R., Wagenaar, J.A., Duim, B., Bolton, F.J., Fox, A., Wareing,
 D.R.A. and Maiden, M.C.J., 2002. Molecular Characterization of *Campylobacter jejuni*Clones A Basis for Epidemiologic Investigation. Emerging Infectious Diseases 8, 949955.

424 Doyle, M.P. and Roman, D.J., 1981. Growth and survival of *Campylobacter fetus* subsp.
425 *jejuni* as a Function of Temperature and pH. Journal of Food Protection 44, 596-601.

Duffy, L.L., Blackall, P.J., Cobbold, R.N. and Fegan, N., 2014. Quantitative effects of in-line
 operations on *Campylobacter* and *Escherichia coli* through two Australian broiler
 processing plants. International Journal of Food Microbiology 188, 128-134.

429 EFSA and EDC, 2018. The European Union summary report on trends and sources of 430 zoonoses, zoonotic agents and food-borne outbreaks in 2017. EFSA Journal 16, 262.

431 Eideh, A.M.F. and Al-Qadiri, H.M., 2011. Effect of Refrigerated and Frozen Storage on the
432 Survival of *Campylobacter jejuni* in Cooked Chicken Meat Breast. Journal of Food
433 Science 76, 17-21.

- European Commision, 2017. Commission regulation (EU) 2017/1495 of 23 August 2017
 amending regulation (EC) no 2073/2005 as regards *Campylobacter* in broiler
 carcases. Official Journal of the European Union L 218, 1-6.
- Fouts, D.E., Mongodin, E.F., Mandrell, R.E., Miller, W.G., Rasko, D.A., Ravel, J., Brinkac,
 L.M., DeBoy, R.T., Parker, C.T., Daugherty, S.C., Dodson, R.J., Durkin, A.S.,
 Madupu, R., Sullivan, S.A., Shetty, J.U., Ayodeji, M.A., Shvartsbeyn, A., Schatz,
 M.C., Badger, J.H., Fraser, C.M. and Nelson, K.E., 2005. Major structural differences
 and novel potential virulence mechanisms from the genomes of multiple
- 442 *Campylobacter* species. PLoS Biology 3, e15.
- Garenaux, A., Jugiau, F., Rama, F., de Jonge, R., Denis, M., Federighi, M. and Ritz, M.,
 2008. Survival of Campylobacter jejuni strains from different origins under oxidative
 stress conditions: effect of temperature. Curr Microbiol 56, 293-297.
- Gaynor, E.C., Wells, D.H., MacKichan, J.K. and Falkow, S., 2005. The *Campylobacter jejuni*stringent response controls specific stress survival and virulence-associated
 phenotypes. Molecular Microbiology 56, 8-27.
- Guyard-Nicodeme, M., Keita, A., Quesne, S., Amelot, M., Poezevara, T., Le Berre, B.,
 Sanchez, J., Vesseur, P., Martin, A., Medel, P. and Chemaly, M., 2016. Efficacy of
 feed additives against *Campylobacter* in live broilers during the entire rearing period.
 Poultry Science 95, 298-305.
- Guyard-Nicodeme, M., Rivoal, K., Houard, E., Rose, V., Quesne, S., Mourand, G., Rouxel,
 S., Kempf, I., Guillier, L., Gauchard, F. and Chemaly, M., 2015. Prevalence and
 characterization of *Campylobacter jejuni* from chicken meat sold in French retail
 outlets. International Journal of Food Microbiology 203, 8-14.
- Guyard-Nicodeme, M., Tresse, O., Houard, E., Jugiau, F., Courtillon, C., El Manaa, K.,
 Laisney, M.J. and Chemaly, M., 2013. Characterization of *Campylobacter* spp.
 transferred from naturally contaminated chicken legs to cooked chicken slices via a
 cutting board. International Journal of Food Microbiology 164, 7-14.

Habib, I., Miller, W.G., Uyttendaele, M., Houf, K. and De Zutter, L., 2009. Clonal population
structure and antimicrobial resistance of *Campylobacter jejuni* in chicken meat from
Belgium. Applied and Environmental Microbiology 75, 4264-4272.

- Habib, I., Uyttendaele, M. and De Zutter, L., 2010. Survival of poultry-derived *Campylobacter jejuni* of multilocus sequence type clonal complexes 21 and 45 under freeze, chill,
 oxidative, acid and heat stresses. Food Microbiology 27, 829-834.
- Hazeleger, W.C., Wouters, J.A., Rombouts, F.M. and Abee, T., 1998. Physiological Activity
 of *Campylobacter jejuni* Far below the Minimal Growth Temperature. Applied and
 Environmental Microbiology 64, 3917-3922.
- Hermans, D., Van Deun, K., Messens, W., Martel, A., Van Immerseel, F., Haesebrouck, F.,
 Rasschaert, G., Heyndrickx, M. and Pasmans, F., 2011. *Campylobacter* control in
 poultry by current intervention measures ineffective: urgent need for intensified
 fundamental research. Veterinary Microbiology 152, 219-228.
- Humphrey, T., O'Brien, S. and Madsen, M., 2007. Campylobacters as zoonotic pathogens: A
 food production perspective. International Journal of Food Microbiology 117, 237-257.
- James, C., Vincent, C., de Andrade Lima, T. and James, S., 2006. The primary chilling of
 poultry carcasses a review. International Journal of Refrigeration 29, 847-862.
- Johnsen, G., Kruse, H. and Hofshagen, M., 2006. Genotyping of *Campylobacter jejuni* from
 Broiler Carcasses and slaughterhouse environment by amplified fragment length
 polymorphism. Poultry Science 85, 2278-2284.
- Jones, D.M., Sutcliffe, E.M., Rios, R., Fox, A.J. and Curry, A., 1993. *Campylobacter jejuni*adapts to aerobic metabolism in the environment. Journal of Medical Microbiology 38,
 145-150.
- Klančnik, A., Guzej, B., Jamnik, P., Vuckovic, D., Abram, M. and Mozina, S.S., 2009. Stress
 response and pathogenic potential of *Campylobacter jejuni* cells exposed to
 starvation. Research in Microbiology 160, 345-352.
- Koidis, P. and Doyle, M.P., 1983. Survival of *Campylobacter jejuni* in fresh and heated red
 meat. Journal Food of Protection 46, 771-774.

- Laguerre, O., Derens, E. and Palagos, B., 2001. Study of domestic refrigerator temperature
 and analysis of factors affecting temperature: a French survey. International Journal
 of Refrigeration 25, 653-659.
- Levesque, S., Frost, E., Arbeit, R.D. and Michaud, S., 2008. Multilocus sequence typing of *Campylobacter jejuni* isolates from humans, chickens, raw milk, and environmental
 water in Quebec, Canada. Journal of Clinical Microbiology 46, 3404-3411.
- Ligowska, M., Cohn, M.T., Stabler, R.A., Wren, B.W. and Brøndsted, L., 2011. Effect of chicken meat environment on gene expression of *Campylobacter jejuni* and its relevance to survival in food. International Journal of Food Microbiology 145, 111-115.
- Lin, J., 2009. Novel approaches for *Campylobacter* control in poultry. Foodborne pathogens
 and disease 6, 755-765.
- 501 Ma, Y., Hanning, I. and Slavik, M., 2008. Stress-induced adaptive tolerance response and 502 virulence gene expression in *Campylobacter jejuni*. Journal of Food Safety 29, 126-503 143.
- Martinez-Rodriguez, A. and Mackey, B.M., 2005. Physiological changes in *Campylobacter jejuni* on entry into stationary phase. International Journal of Food Microbiology 101,
 1-8.
- McCarthy, Z., Smith, B., Fazil, A., Wu, J., Ryan, S.D. and Munther, D., 2018. pH dependent
 C. jejuni thermal inactivation models and application to poultry scalding. Journal of
 Food Engineering 223, 1-9.
- 510 Meunier, M., Guyard-Nicodeme, M., Vigouroux, E., Poezevara, T., Beven, V., Quesne, S.,
- 511 Bigault, L., Amelot, M., Dory, D. and Chemaly, M., 2017. Promising new vaccine 512 candidates against *Campylobacter* in broilers. PLoS One 12, e0188472.
- 513 Murphy, C., Carroll, C. and Jordan, K.N., 2003a. Identification of a novel stress resistance 514 mechanism in *Campylobacter jejuni*. Journal of Applied Microbiology 95, 704-708.

- Murphy, C., Carroll, C. and Jordan, K.N., 2003b. Induction of an adaptive tolerance response
 in the foodborne pathogen, *Campylobacter jejuni*. FEMS Microbiology Letters 223,
 89-93.
- Newell, D.G., Shreeve, J.E., Toszeghy, M., Domingue, G., Bull, S., Humphrey, T. and Mead,
 G., 2001. Changes in the carriage of *Campylobacter* strains by poultry carcasses
 during processing in abattoirs. Applied and Environmental Microbiology 67, 26362640.
- Nguyen, H.T., Corry, J.E. and Miles, C.A., 2006. Heat resistance and mechanism of heat
 inactivation in thermophilic *Campylobacters*. Applied and Environmental Microbiology
 72, 908-913.
- 525 Nyati, K.K. and Nyati, R., 2013. Role of *Campylobacter jejuni* infection in the pathogenesis of 526 Guillain-Barre syndrome: an update. Biomed Research International 2013, 852195.
- 527 Oh, E., McMullen, L. and Jeon, B., 2015. Impact of oxidative stress defense on bacterial
 528 survival and morphological change in *Campylobacter jejuni* under aerobic conditions.
 529 Frontiers in Microbiology 6, 295.
- Osiriphun, S., Tuitemwong, P., Koetsinchai, W., Tuitemwong, K. and Erickson, L.E., 2012.
 Model of inactivation of *Campylobacter jejuni* in poultry scalding. Journal of Food
 Engineering 110, 38-43.
- 533 Phebus, R.K., Draughon, F.A. and Mount, J.R., 1991. Survival of *Campylobacter jejuni* in 534 modified atmosphere packaged turkey roll. Journal of Food Protection 54, 194-199.
- Pujol, L., Kan-King-Yu, D., Le Marc, Y., Johnston, M.D., Rama-Heuzard, F., Guillou, S.,
 McClure, P. and Membre, J.M., 2012. Establishing equivalence for microbial-growthinhibitory effects ("iso-hurdle rules") by analyzing disparate *Listeria monocytogenes*data with a gamma-type predictive model. Applied and Environmental Microbiology
 78, 1069-1080.
- 540 Ragimbeau, C., Schneider, F., Losch, S., Even, J. and Mossong, J., 2008. Multilocus 541 sequence typing, pulsed-field gel electrophoresis, and fla short variable region typing

- 542 of clonal complexes of *Campylobacter jejuni* strains of human, bovine, and poultry 543 origins in Luxembourg. Applied and Environmental Microbiology 74, 7715-7722.
- Reid, A.N., Pandey, R., Palyada, K., Naikare, H. and Stintzi, A., 2008a. Identification of *Campylobacter jejuni* genes involved in the response to acidic pH and stomach
 transit. Applied and Environmental Microbiology 74, 1583-1597.
- Reid, A.N., Pandey, R., Palyada, K., Whitworth, L., Doukhanine, E. and Stintzi, A., 2008b.
 Identification of *Campylobacter jejuni* genes contributing to acid adaptation by
 transcriptional profiling and genome-wide mutagenesis. Applied and Environmental
 Microbiology 74, 1598-1612.
- Rivoal, K., Poezevara, T., Quesne, S. and Chemaly, M., 2016. Optimisation des conditions
 de refroidissement des volailles à l'abattoir pour contrôler la contamination en *Campylobacter*. In: Agence nationale de sécurité sanitaire de l'alimentation, d. l. e. e.
 d. t. (ed.), Journée d'information et d'échanges sur *Campylobacter*, Ploufragan.
- Rivoal, K., Ragimbeau, C., Salvat, G., Colin, P. and Ermel, G., 2005. Genomic diversity of *Campylobacter coli* and *Campylobacter jejuni* isolates recovered from free-range
 broiler farms and comparison with isolates of various origins. Applied and
 Environmental Microbiology 71, 6216-6227.
- Saint-Cyr, M.J., Haddad, N., Taminiau, B., Poezevara, T., Quesne, S., Amelot, M., Daube,
 G., Chemaly, M., Dousset, X. and Guyard-Nicodeme, M., 2017. Use of the potential
 probiotic strain *Lactobacillus salivarius* SMXD51 to control *Campylobacter jejuni* in
 broilers. International Journal of Food Microbiology 247, 9-17.
- Sheppard, S.K., Dallas, J.F., MacRae, M., McCarthy, N.D., Sproston, E.L., Gormley, F.J.,
 Strachan, N.J., Ogden, I.D., Maiden, M.C. and Forbes, K.J., 2009. *Campylobacter*genotypes from food animals, environmental sources and clinical disease in Scotland
 2005/6. International Journal of Food Microbiology 134, 96-103.
- 567 Sörqvist, S., 1989. Heat resistance of *Campylobacter* and *Yersinia* strains by three methods.
 568 Journal of Applied Bacteriology 67, 543-549.

569	Vashin, I.T. and Stoyanchev, T.T., 2011. Influence of temperature on Campylobacter jejuni
570	survival rate in pork meat. Bulgarian Journal of Veterinary Medicine 14, 25-30.
571	Waterman, S.C., 1982. The heat-sensitivity of Campylobacter jejuni in milk. Journal of
572	Hygiene 88, 529-533.
573	Young, K.T., Davis, L.M. and Dirita, V.J., 2007. Campylobacter jejuni: molecular biology and
574	pathogenesis. Nature Reviews Microbiology 5, 665-679.
575	Zoz, F., Iaconelli, C., Lang, E., Iddir, H., Guyot, S., Grandvalet, C., Gervais, P. and Beney, L.,
576	2016. Control of Relative Air Humidity as a Potential Means to Improve Hygiene on
577	Surfaces: A Preliminary Approach with Listeria monocytogenes. PLoS One 11, e0148418.

581 Table 1: Results from ANOVA of the influence of the different factors and their interaction on

Source	DF	Sum Sq	Mean Sq	F value	Р
Model	9	160.941	17.882	35.661	< 0.0001
Error	124	62.181	0.501		
Total	133	223.122			
Heat	2	19.254	9.627	19.198	< 0.0001
Atmosphere	1	110.907	110.907	221.169	< 0.0001
Strain	2	16.725	8.363	16.677	< 0.0001
Heat*Atmosphere	2	6.419	3.210	6.401	0.002
Atmosphere*Strain	2	12.806	6.403	12.768	< 0.0001

582 inactivation of *C. jejuni* during cold storage

	рН	Medium	Strain	D₅₀∘c (min)	D₅₅∘c (min)	Reference	
			C97anses640	No inactivation	2.5*		
_	7.2 7.4	Mueller-hinton broth	C09MJLT205	No inactivation	2.8*	This study	
m			RM1221	No inactivation	1.5*		
edi		Heart infusion broth	AR6	36	5.3	(Nguyen et al. 2006)	
E P			L51	39	4.6		
La			H-840	ND	0.6		
	ND	1% peptone solution	composite of 5			(Blankenship and Craven 1982)	
			strains	ND	1.1		
	ND	Physiological saline	1503	ND	1.14	(Sörqvist 1989)	
			FRI-CF3	5.4	0.7		
			FRI-CF6	4.7	1		
	6.8 7.2	Skim milk Milk	FRI-CF8	3.5	0.9	(Doyle and Roman 1981)	
			FRI-CF12	4.4	1		
			FRI-CF16	5.1	0.9		
			24791	5.7	ND		
×			16000	ND	ND		
atri			21033	7.2	ND		
Ĕ			172589	ND	ND	(Waterman 1982)	
000			16509	7.3	1.1		
ŭ			5388	36	ND		
	ND	Chicken carcasses	H-840	ND	2.1		
			composite of 5			(Blankenship and Craven 1982	
			strains	ND	2.3		
	8	Chicken carcasses	Endogeneous flora	ND	0.8*	(Osiriphun et al. 2012)	
	6	Lamb meat	FRI-CF8	5.9	1	(Kaidis and Dayla 1982)	
			FRI-CF31P	11.2	1.2	(Koldis and Doyle 1983)	

Table 2: D-values (min) of *C. jejuni* collected from different studies

			FRI-CF401S	13.2	ND	
			FRI-CF402S	11.1	ND	
			FRI-CF403S	8.96	ND	
			FRI-CF404S	13.3	1.23	
1	ND	Chicken carcasses	Endogeneous flora	ND	0.9	(Berrang and Dickens 2000)
1	ND	Chicken carcasses	Endogeneous flora	ND	1.4	(Duffy et al. 2014)

584

585 ND: Not Determined

586 *D_{55°C}-values were calculated using the formula log D $_{55°C}$ = log D $_{54°C}$ - (T $_{55°C}$ - T $_{54°C}$) / Z

587 Z = 4.99°C was estimated by calculating the mean Z reported from different studies (Blankenship and Craven 1982, Doyle and Roman 1981, Sörqvist 1989, Waterman 1982).

588 Figures legends:

- 589 Figure 1. Global scheme of experimental procedure. The different factors are represented with their associated levels. After each step, the
- 590 inactivation (viable bacterial population N lost) was calculated (Δlog).
- 591 **Figure 2.** Contribution of factors (temperature applied during hot and cold bath, strain and gas composition of the atmosphere) on the
- 592 inactivation of *C. jejuni* induced by each step of the experimental procedure (hot bath in white, cold bath in black, storage at 6°C in grey) under
- atmosphere 70% O_2 / 30% CO_2 (a) and atmosphere 50% N_2 / 50% CO_2 (b).
- 594 **Figure 3.** Effect of factors and interactions on inactivation during storage at 6°C under modified atmosphere: strain effect (a), mean effect of
- 595 temperature applied during heat stress all strains combined (b), mean effect of the gas atmosphere composition all strains combined (c),
- 596 interaction between strains and atmosphere (d) and mean interaction between temperature of heat stress and atmosphere all strains combined
- 597 (e).
- 598 A,B,C Different capital letters indicate significant difference according the Least Significant test (P < 0.05) in inactivation.
- 599
- 600

Figure 1





Figure 2

Figure 3

