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(Article begins on next page)

1	Campylobacter spp. prevalence and mitigation strategies in the broiler production chain
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20 ABSTRACT

This study aims to discuss the microbial ecology of the broiler gut environment, 21 Campylobacter prevalence across the broiler production chain with a follow-up focus on a 22 possible mitigation strategy based on the use of bacteriophages. Scientific literature published 23 from the last two decades was reviewed and data were collected to establish the ranges of 24 25 *Campylobacter* loads from different samples. Results showed that the pathogen load in the sample is likely to increase from the different stages of the production chain. Contamination of 26 water and feed represents the most notable source of contamination during the primary 27 28 production, while cross-contamination of broiler carcasses, skin, and meat occurs during the slaughter, dressing, and processing via machinery, work surfaces, water, and air partially due to 29 the leaking of contaminated feces from visceral rupture. Knowledge gaps were identified and 30 included: a lack of studies detecting Campylobacter in broilers in most of the European countries 31 over the last decade and a low number of studies determining the bacterial load in crates used to 32 transport broilers to the slaughterhouse. Determining the prevalence of *Campylobacter* in the 33 broiler industry will enable us to set critical control points to produce broiler flocks and meat 34 products with a low risk of *Campylobacter* contamination. 35

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37 Keywords: Microbiota, chicken, bacteriophage, biocontrol, poultry, public health risk

1. INTRODUCTION

Zoonoses are defined as those infectious diseases naturally transmitted from vertebrate 39 animals to humans through direct or indirect contact (food or water contamination). Zoonotic 40 agents include a wide variety of bacteria, viruses, protozoa, insects, and helminths. According to 41 the World Health Organization (WHO), almost 600 million cases of food-borne zoonoses were 42 43 reported worldwide in 2015, of which 52% were caused by pathogenic bacteria (WHO, 2015). Campylobacter is the most common pathogen responsible for food-borne zoonotic diseases in 44 humans and it is considered a serious public health issue in both developing and developed 45 46 countries. The most recent summary report of the European Food Safety Authority (EFSA) stated that campylobacteriosis is the most frequently reported food-borne zoonoses in the 47 European Union (EU), with 220 682 confirmed cases in 2019, reported from 18 different 48 members states (EFSA and European Centre of Disease Prevention and Control (ECDC), 2019). 49 The most common source of infection in humans due to Campylobacter are broiler meat and 50 milk (EFSA and European Centre of Disease Prevention and Control (ECDC), 2019). 51 52 *Campylobacter* is a genus of microaerophilic and Gram-negative bacteria belonging to the Proteobacteria phylum. Bacterial cells generally appear as slender, spirally curved, or 53 54 comma-shaped rods characterized by inability to form endospores and ability to change to spherical or coccoid form under stress conditions (Lastovica et al., 2014). Campylobacter species 55 have an optimal growth temperature, O₂, and CO₂ concentration of 30- 42°C, 5-10%, and 3-5%, 56 57 respectively (Lastovica et al., 2014). In terms of biochemical characteristics of Campylobacter species, they are oxidase-positive, with the capacity to reduce fumarate to succinate but are not 58 59 able to metabolize lipids, starch, gelatin, and casein (Lastovica et al., 2014). Currently, 33 species have been taxonomically described from this genus, but this number is increasing due to 60

61 the identification of new species (bacterio.net). The main natural reservoir of

thermophilic/thermotolerant Campylobacter species has been extensively reported in warm-62 blood animals, including most mammals, birds, and food-producing animals (Silva et al., 2011). 63 The mechanism of colonization, adherence, and invasion of host by pathogenic Campylobacter 64 species such as C. jejuni, has been previously reviewed elsewhere (Elmi et al., 2021). In 65 66 summary, successful colonization of the bacteria depends on the ability to attach to the gut mucosa, motility, chemotaxis, spiral shape of the cell, the functionality of the flagella, production 67 of toxins and secreted proteins (cytolethal distending toxin -CDT) and other virulence factors 68 (Elmi et al., 2021). 69

Interestingly, *Campylobacter* does not multiply outside a warm-blooded host due to the 70 71 absence of microaerobic conditions. However, they can survive when protected from dryness. In 72 this regard, it has been shown that Campylobacter can survive up to 3 months in slurries and dirty water (Nicholson et al., 2005). The mechanism of survival of the Campylobacter species 73 74 when exposed to stress environments has been explained by its ability to form a biofilm on abiotic surfaces, this biofilm ensures a supply of nutrients and mechanical protection to survive 75 (Johnson et al., 2017). Although the bacteria cannot multiply outside the animal hosts or in food 76 77 during storage, it has been shown that they are able to develop complex mechanisms of virulence which remain poorly understood. 78

Recent studies have facilitated a greater appreciation of the complex mechanism of
virulence of the members of the *Campylobacter* genus. The full genome of *Campylobacter jejuni*, has elucidated strain-specific genetic diversity with high genome plasticity (Bacon et al.,
2000). The ability to survive and adapt to stress environments indicates that *C. jejuni* harbors
complex virulence and fitness factors (Tegtmeyer et al., 2021). Virulence factors and

84	pathogenicity islands in C. jejuni have been identified and are reported elsewhere (Ali et al.,
85	2012; Bacon et al., 2000; Sierra-Arguello et al., 2021; Tang et al., 2020; Zhang et al., 2017).
86	Virulence-associated genes such as <i>flaA</i> , <i>cadF</i> , <i>cdtA</i> , <i>cdtB</i> , <i>cdtC</i> , <i>cheY</i> , <i>iamA</i> , and <i>virB11</i> were
87	recently identified in C. jejuni and Campylobacter coli isolates originating from broiler of 31-day
88	or 37-day age at the rearing period to slaughtering process (Tang et al., 2020). Besides the
89	identification of virulence-associated genes, invasion-associated genes, plasmid genes and CDT-
90	associated genes were also recognized in C. jejuni strains isolated from cloacal, broiler carcasses,
91	and broiler slaughterhouses (Sierra-Arguello et al., 2021). The aforementioned genes are
92	involved in the adhesion, invasion, chemotaxis, motility, toxin-activity, and host immune system
93	evasion. Interestingly, there is a difference of virulence factors between C. jejuni and C. coli,
94	where a higher number of virulence genes were retrieved in C. jejuni if compared with C. coli,
95	this difference might contribute to the higher colonization of C. jejuni in the broilers' intestines
96	(Tang et al., 2020; Zhang et al., 2017).

Antibiotic resistance in *Campylobacter* is also considered a global trend. In this regard, *C. jejuni* and *C. coli* had shown a multi-drug resistance to several antibiotics such as
tetracyclines, macrolides, aminoglycosides, and β-lactams. However, a higher number of
antibiotic resistance genes were retrieved for *C. jejuni* if compared with *C. coli* (Tang et al.,
2020). On the increase of antibiotic resistance to more than one class of antibiotics, further
research is needed to understand the mechanism of antimicrobial resistance to improve not only
human but also animal health.

Besides the genetic makeup, the main factors that influence the occurrence of *Campylobacter* in broilers are related to the host gut environment, production chain, or farm practices (Barker et al., 2020; Djennad et al., 2017; McKenna et al., 2020; Perez-Arnedo and Gonzalez-Fandos, 2019; Sibanda et al., 2018; Tang et al., 2020). A conceptual framework of the factors increasing
the occurrence of *Campylobacter* and a prevention guideline to stipulate the best conditions and
food processing management to reduce the risk of *Campylobacter* contamination in the broiler
production chain has been developed (EFSA, Panel on Biological Hazards, 2011; Lyngstad,
Jonsson, Hofshagen, & Heier, 2008).

112 Several intervention methods have been developed in recent years, such as the combination of strict biosecurity measures, good manufacturing practice (GMP), hazard analysis and critical 113 control points (HACCP), Campylobacter vaccines, antibiotic alternatives to control 114 115 Campylobacter, probiotics, and phytochemicals (Deng et al., 2020; European Food Safety Authority Panel on Biological Hazards, 2011; Umar et al., 2016; Ushanov et al., 2020). 116 However, the problem has not been completely eradicated and the prevalence of this pathogen is 117 118 still high. Ante- and post-mortem veterinary inspections of broilers are routinely used at the slaughterhouse level as a strategy to ensure that meat does not bear fecal or other contaminants. 119 However, the presence of Campylobacter in broiler carcasses cannot be detected visually. As an 120 121 attempt to mitigate this issue, the application of *Campylobacter*-specific bacteriophages has emerged as one of the most promising approaches to be applied within the farm-to-fork poultry 122 123 process (Atterbury et al., 2003; Fischer et al., 2013; Hammerl et al., 2014; Kittler et al., 2013; Richards et al., 2019). In this context, this review focuses on discussing the most updated 124 scientific achievements made on the microbial ecology of the gastrointestinal (GI) tract of 125 126 broilers and the interaction between chickens' gut microbiota and Campylobacter, *Campylobacter* prevalence across the broiler production chain with a follow up of the application 127 128 of bacteriophage along the farm-to-fork process.

2. MICROBIAL ECOLOGY IN CHICKEN' ECOSYSTEMS

Chicken ecosystems harbor complex, diverse, and dynamic microbial communities 131 composed of bacteria, protozoa, fungi, yeasts, bacteriophages, and other viruses. The integrity of 132 the GI tract and the gut microbiota composition has a direct influence on chicken' health, 133 affecting the development of their digestive and immune systems (Clavijo and Flórez, 2018; 134 135 Khan et al., 2020). The avian gut microorganisms are mainly responsible for the continuous generation of energy and nutrients, such as vitamins (K and B groups), amino acids, short-chain 136 fatty acids (SCFA like, butyric, propionic, and lactic acids), ammonia, antimicrobial compounds 137 138 (bacteriocins) and the decrease of triglyceride concentrations in the avian gut (Vispo and Karasov, 1997). The positive effect of the production of SCFA on the host includes the inhibition 139 140 of pathogens, reduction of the pH levels in the colon, and stimulation of the cell proliferation in 141 the gut epithelium (Christl et al., 1997; Dibner and Richards, 2005; Ricke, 2003). In contrast, chickens' growth can be depressed when competition for energy and protein occurs between the 142 143 commensal microbiota. As a result of this competition, toxic metabolites (amino acid catabolites) can be produced and impact the growth of the animal (Shang et al., 2018). 144

As reported in recent years, the composition and function of chickens' microbiota vary depending on biological changes within and between hosts (age, sex, maternal factors, and breed), and environmental factors (biosecurity levels, housing, litter, feed access and antibiotic administration, hygiene, location, and climate) (Kers et al., 2018). Regarding the biological variation within and between hosts, it has been demonstrated that the chickens' microbiota richness increases during the first week of life (Ballou et al., 2016; Crhanova et al., 2011), while the number of different microbial taxa decreases with chicken age (Lu et al., 2003). The

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microbial composition of chickens does not only change with chicken age but it is also influenced by the location in the digestive tract and diet (Shang et al., 2018).

154 Significant progress has been made in understanding the microbial taxonomic 155 composition of the different sections of the chickens' GI tract. Overall, Firmicutes is the phylum most abundant in both ilea and ceca of chickens (Kumar et al., 2018; Lu et al., 2003; Oakley et 156 157 al., 2014). Interestingly, in the ceca, the relative abundance of members of the *Clostridium* genus increased by 10 fold between weeks 1 and 6 post-hatch (Oakley and Kogut, 2016). Concerning 158 159 the bacterial community within the small intestine, in this section lactobacilli mainly dominated 160 the microbial ecosystem (Gong et al., 2007; Lu et al., 2003). According to Lu et al., the microbial composition of the ceca and ileum of three days of age broilers (chickens for meat 161 production) fed with a vegetarian corn-soy diet was mainly composed of *Lactobacillus* 162 delbrueckii (13 and 42%, respectively), Clostridium spp. (31 and 1%, respectively) and 163 *Clostridium perfringens* (13 and 16%, respectively), however differences in the microbial 164 165 composition between these sections (ceca and ileum) were reported (Lu et al., 2003). Regarding seven to 49 days old chickens, Clostridium spp. remained the most abundant bacteria in the ceca, 166 followed by *Ruminococcus*, while differences in the relative abundance of bacterial species 167 168 overtime were reported in the ileum (Lu et al., 2003). In detail, Lactobacillus acidophilus (50-59%) was the most abundant bacteria in chickens of seven to 21 days of age, while a unique 169 community was reported in the ileum of three, 28, and 49 days of age broilers. Interestingly, 170 171 regardless of the absence of *Clostridium* spp. and *Ruminococcus* spp. in the ileum section at an early age (between three to 14 days), significant differences of the microbiota between the 172 different sections of the GI tract (ileum and ceca) were only found after 14 days of age (Lu et al., 173 2003). 174

Overall, it is still not clear whether *Proteobacteria* and *Bacteroidetes* are also abundant 175 phyla during the first six weeks in the chicken's ceca (Shang et al., 2018). These contradictory 176 results may be related to the different chicken breeds studied. According to Kers and colleagues 177 (2018) a variation at the phylum level from the ceca samples of broiler breeds (meat production) 178 and laying-type chickens (egg production) were observed at different time points using 16S 179 180 rRNA gene amplicon sequencing. Specifically, at zero hr, Firmicutes was the most abundant phylum reported in meat-type chicks (Pedroso et al., 2016; Danzeisen et al., 2011), while 181 182 Proteobacteria was identified in laying-type chicks (Ballou et al., 2016). This variability may be 183 due to sample types (fecal vs cecal), feed intervention, and/or the technical aspects of the microbial identification as reported elsewhere (Shang et al., 2018). However, from one to 42 184 days of age, *Firmicutes* was the most abundant phylum regardless of the type of broiler breeds. 185 Furthermore, *Candidatus arthromitus*, a desirable bacterium commonly associated with healthy 186 GI tracts in animals, has been also identified in the jejunum and ileum of chickens (Gong et al., 187 2007, 2002). 188

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2.1 Chicken diet and intestinal microbiota interaction

The characteristics of the chickens, feed management, the use of medications or vaccines, 190 the environmental conditions of the poultry house, and the housing systems are known factors 191 that have a short- or long-term effect on the intestinal microbiota composition and immune 192 193 system development of chickens (Kers et al., 2018). The effect of feed management on the intestinal microbiota has been extensively studied (Apajalahti, 2005; McKenna et al., 2020; 194 Singh et al., 2014; Stanley et al., 2012; Takeshita et al., 2021). Differences in the relative 195 196 abundance of bacterial species in fecal and cecal samples of broilers following high and low growth diets were reported elsewhere (Singh et al., 2014; Stanley et al., 2012). Knarreborg and 197

colleagues (2002) demonstrated that the divergence in feed can also increase or decrease the 198 relative abundance of a specific bacterial group: the aforementioned study shows how pellet feed 199 increases the number of Enterococcus spp. and coliforms and decreases lactobacilli species and 200 C. perfringens in the ileum of broilers when compared with mash feed (Knarreborg et al., 2002). 201 In contrast, when chickens consumed corn, this diet favors a decrease in the number of clostridia, 202 203 enterococci, and lactobacilli, while when chickens consumed wheat, it favors the increase of bifidobacteria (Apajalahti, 2005). Interestingly, the amount of protein in the chicken feed also 204 changed microbial composition, where high amount of protein showed a lower relative 205 206 abundance of lactobacilli species compared with chicken feed with a low amount of proteins diet (Takeshita et al., 2021). The difference in microbial community structure between production 207 systems together with different management paraments such as stocking density has also shown 208 209 to alter the microbiota of broilers (McKenna et al., 2020). To date, broilers' microbiota studies have focused on identifying bacterial composition while the identification of other components, 210 211 such as fungi, phages, or viruses, remain unclear. A better understanding of the role and interactions between mycobiota, phagobiota, and virobiota with the broiler microbial ecosystem 212 may help to improve chicken productivity, health, and welfare and develop novel strategies for 213 controlling the prevalence of *Campylobacter* spp. in broilers (Silva et al., 2011). 214

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2.2 Campylobacter and its interactions with gut microbiota in chickens

Campylobacter typically occurs within two weeks onwards and increases during broilers
life cycle (Ijaz et al., 2018; Kalupahana et al., 2013; McKenna et al., 2020; Neill et al., 1984;
Thibodeau et al., 2015). It usually grows in the ceca and cloaca and can also colonize the spleen,
blood, and liver (Lin, 2009). *C. jejuni* and *C. coli* are the most commonly isolated *Campylobacter* species in broiler samples at different production stages. Interestingly, recent

studies show contradictory results on how microbial ecology influences Campylobacter 221 colonization. According to Sofka and colleagues the presence of Campylobacter decreased the 222 diversity of intestinal microbiota when compared with non-colonized broilers (Sofka et al., 223 2015). In alignment with the aforementioned study, a significant difference in the relative 224 abundance of the microbial operational taxonomic units detected in the ceca of chickens fed with 225 226 different diets, at different ages from three different commercial broiler farms were reported between Campylobacter-positive and -negative chickens (Takeshita et al., 2021). Interestingly, 227 the decrease in lactobacilli abundance in chicken ceca was associated with high levels of 228 229 Campylobacter, while the growth of Campylobacter increased the levels of Enterobacteriaceae (Sakaridis et al., 2018). However, according to McKenna and colleagues the presence of 230 *Campylobacter* in broilers reared under standard industrial growing systems increased the cecal 231 232 microbial community structure (McKenna et al., 2020). Whereas the transplantation of cecal microbial in chickens shows no significant difference in the ceca microbial communities of 233 different inbred chickens (Chintoan-Uta et al., 2020). The factors affecting host-pathogen 234 ecology in terms of the microbiome and the microbial dynamics and *Campylobacter* presence 235 remain poorly studied at an industrial or small-scale farm level. 236

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238 3. CAMPYLOBACTER PREVALENCE IN THE BROILER PRODUCTION CHAIN

In terms of *Campylobacter* diversity in the broiler chain production, a recent study has demonstrated that *C. jejuni* predominates during the rearing broiler period while more isolates of *C. coli* were detected during the slaughtering process (Chen et al., 2010; Tang et al., 2020; Zhang et al., 2018).

Campylobacter from chicken reservoirs may reach humans through the environment or by 243 direct contact and mainly through the consumption of raw and undercooked contaminated broiler 244 meat and meat products. The Scientific Opinion of the Panel on Biological Hazards (BIOHAZ), 245 EFSA in 2010 estimated that the majority of human campylobacteriosis is attributed to the 246 chicken reservoir as a whole (50 - 80%), while the handling, preparation, and consumption of 247 248 broiler meat may account only for 20 to 30% (EFSA, 2010). Other food products such as unpasteurized dairy products and contaminated water are also vehicles of transmission of this 249 pathogen, but Campylobacter infections are less common from these sources compared with 250 251 meat products (Josefsen et al., 2015). To provide an indication of the possible control points for *Campylobacter* at the broiler meat production chain, this review describes the prevalence of 252 *Campylobacter* along the process and discusses the risk factors that influence the level of 253 254 contamination (Figure 1).

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3.1 Primary production

The increases of *Campylobacter* loads during the first weeks of age of commercial flocks 256 (Umar et al., 2016) indicate that vertical transmission of the pathogen does not commonly occur; 257 however, it still represents an important risk factor for the *Campylobacter* colonization in 258 259 broilers (Bull et al., 2008; European Food Safety Authority Panel on Biological Hazards, 2011; Humphrey, 2006; Tang et al., 2020). The protection of young broilers' GI tract against 260 colonization of Campylobacter has been associated with Campylobacter-specific maternal 261 262 antibodies (MAB) (Sahin et al., 2001; Vandeplas et al., 2010). MAB, predominantly immunoglobulin G (IgG) class, are transported from the egg yolk across the yolk sac membrane 263 into the embryonic circulation to protect young broiler chickens from C. jejuni infections 264 (Linden and Roth, 1978). However, MAB levels change with the increase in the broiler age. In 265

detail, the highest level of MAB is reported during the first two weeks after hatching, followed 266 by a decrease, reaching minimal values at the third and fourth weeks of age (Sahin et al., 2001; 267 Vandeplas et al., 2010). The increased risk of *Campylobacter* colonization in broilers from week 268 three to week six was demonstrated elsewhere, as shown in Table 1 (Ingresa-Capaccioni et al., 269 2016; Perez-Arnedo and Gonzalez-Fandos, 2019; Tang et al., 2020; Tangkham et al., 2016). 270 271 Interestingly, the prevalence of *Campylobacter* spp. and *C. jejuni* at the end of the rearing period is higher in open housing compared with environmentally controlled housing (Tangkham et al., 272 2016). Noteworthy, a higher prevalence of C. jejuni isolates compared with C. coli was observed 273 274 from the broiler rearing period at the farm level in China (Tang et al., 2020), while in Spain, C. *coli* has not been detected throughout chickens progeny (Ingresa-Capaccioni et al., 2016; Perez-275 Arnedo and Gonzalez-Fandos, 2019). 276

277 Natural colonization of broilers by single or multiple *Campylobacter* species rapidly occurs through horizontal transmission from vectors such as domestic pets, insects, rodents and 278 migratory birds, farm equipment, transport vehicles, farmworkers, drinking water, feed, litter, 279 air, or thinning (Bang et al., 2002; Carvalho et al., 2010; Gharib Naseri et al., 2012; Koolman et 280 al., 2014; Robyn et al., 2013; Schroeder et al., 2014; Stern et al., 2001). Based on the most recent 281 282 studies considered in this review, the highest prevalence of *Campylobacter* presence has been detected on drinking water, feed, workers boots, and broilers' fecal and cecal samples after 283 thinning, while the lowest prevalence of this bacteria was found on samples from the litter, air 284 285 and the infrastructure of the farm (Table 1).

The colonization of the flocks with *Campylobacter* can also be introduced from external visitors to the farm, maintenance personnel, bird catching crew, close livestock, when operators visit more than one farm or are negligent regarding hygienic practices (Berndtson et al., 1996;

Cardinale et al., 2004; Hald et al., 2000; Johnsen et al., 2007; Kapperud et al., 1993; A. M. 289 Ridley et al., 2011). Besides human traffic, the proximity of the fresh litter, larger thinning 290 crews, increasing the number of broiler houses on-site, and the presence of dead broilers in the 291 house enhances the survival of Campylobacter and thus the risk of positive flocks is increased 292 (Cardinale et al., 2004; Koolman et al., 2014; Lyngstad et al., 2008). Inadequate biosecurity such 293 294 as broiler houses older than 15 years, absence of anterooms and barriers in each house, the use of shared tools between houses, long downtime, and drinking systems with bells or cups represent a 295 common risk factor for Campylobacter colonization of broiler flocks (Sommer et al., 2016). 296 297 Partial thinning of broiler flocks has also been considered as a potential risk factor for Campylobacter colonization of the remaining animals, due to the difficulty of keeping 298 299 biosecurity measures during the process (Hermans et al., 2011). 300 The dispersion of *Campylobacter* originating from broilers into the environment represents an important factor leading to increased contamination. In broiler, C. jejuni is the most 301 predominant species colonizing the flocks, followed by C. coli and occasionally by other species 302 (Rossler et al., 2019; Umar et al., 2016). Once the first bird is infected, Campylobacter is 303 horizontally transmitted to most of the birds in a flock within only a few days, reaching between 304 10^6 and 10^8 CFU/g in their intestinal tract, and they remain colonized until slaughter (Marotta et 305 al., 2015). 306

The increased water consumption by chickens during summer months increases the risk of drinking water contaminated with *Campylobacter* (Herman et al., 2002; Lyngstad et al., 2008). This association has been observed in northern European countries, such as Sweden, Denmark, Norway, and the Netherlands (Jore et al., 2010). The changes in the temperature throughout the year can explain the increase in water consumption during summer. However, other factors such as the increment in the abundance of transmission vectors of *Campylobacter* (flies) and the need
for ventilation in the poultry house of broilers can also increase the probabilities of the
introduction of *Campylobacter* into the environment (Hald et al., 2008; Jore et al., 2010). Also, it
must be underlined that, as expected, climate change is an important factor that might increase
the prevalence of *Campylobacter* in the future, placing public health at risk. It is worth noting
that colonized birds are predominantly asymptomatic, with no negative effect on their health
(Pielsticker et al., 2012).

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3.2 Transportation before slaughter

The transportation step consists of moving the broilers from farm facilities to the slaughterhouse. During this process, the animals are confined in crowded crates or cages, deprived of water and feed, and undergo continuous stress that affect not only the meat quality but also the *Campylobacter* prevalence, which could increase due to cross-contamination. If a single broiler is colonized, *Campylobacter* will be spread to the environment and will be transmitted to the rest through fecal matter or feathers contact (Stern et al., 2001; Whyte et al., 2001).

In comparison with the primary production, the pre- and post-transportation steps have 327 been less studied over the last two decades (Table 2). Overall, from the literature available we 328 observed that during transportation, the floor and transportation cage/crates account for the 329 highest risk of cross-contamination in broilers (Stern et al., 2001; Willis et al., 2002), while no 330 clear contamination level in fecal samples between pre-and post-transportation steps is observed 331 (Whyte et al., 2001). One likely explanation for this phenomenon is that catching and placing the 332 333 chickens in cages/crates to transport them to the processing plant increases the risk of contamination, mostly due to cross-contamination during transportation (Slader et al., 2002). 334

However, recent literature has demonstrated the ineffectiveness of cleaning naturally
contaminated crates before using them to transport broilers and reduce *Campylobacter* infections
(A. Ridley et al., 2011). Further, transportation stress alters excretion rates of *Campylobacter* in
the fecal material of broilers (Whyte et al., 2001).

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3.3 Slaughter, dressing, and processing

340 Colonization of *Campylobacter* in broilers during rearing and transportation steps contributes to the contamination of this bacteria in the slaughterhouse causing cross-341 contamination. During the slaughter process, the transmission of Campylobacter-infected flocks 342 343 to non-infected (Shange et al., 2019; Umar et al., 2016) is mainly due to the leaking of contaminated feces from visceral rupture to the chickens (García-Sánchez et al., 2019; Hermans 344 et al., 2011). In the European Union (EU), it was observed that batches of broilers whose 345 intestines were colonized with Campylobacter yielded carcasses with high numbers of 346 Campylobacter (EFSA, Panel on Biological Hazards, 2011). However, Campylobacter counts on 347 carcasses varied among slaughterhouses. These differences found on the levels of 348 *Campylobacter* loads among slaughterhouses might be related to the different hygiene practices 349 used between countries. Interestingly, *Campylobacter* strains in chickens are not necessarily the 350 351 same as those isolated from processed carcasses, which suggests that cross-contamination occurs during processing (Slader et al., 2002). 352

This cross-contamination can occur during the entire slaughter, dressing, and processing including the chilling room, plucking, evisceration and portioning areas, or via machinery, work surfaces, process water and air (Allen et al., 2003; Arnold and Silvers, 2000; Corry and Atabay, 2003; Haas et al., 2005; Hue et al., 2010; Johnsen et al., 2007). Overall, based on the most recent studies considered in this review, the highest prevalence of *Campylobacter* spp. was detected on

the defeathering, evisceration, operation tables, worker's gloves, shackles, and conveyor belt 358 equipment, while the lowest prevalence of Campylobacter spp. was detected on the sink, floor 359 and chopping boards and knife swabs during slaughter, dressing, and processing as shown in 360 Table 3 (García-Sánchez et al., 2017; Khan et al., 2018; Tang et al., 2020; Zhang et al., 2018). In 361 agreement with data from the primary production, where the surfaces and equipment in the 362 363 facility are the main sources for *Campylobacter*, a high prevalence of this bacteria in broiler samples and carcasses has been also reported during the slaughter, dressing and processing 364 (Table 3) (Carrillo et al., 2014; Casagrande Proietti et al., 2018; García-Sánchez, Melero, Diez, 365 366 Jaime, & Rovira, 2018; Ingresa-Capaccioni et al., 2016; Khan et al., 2018; Korsak, Maćkiw, Rożynek, & Żyłowska, 2015; Perez-Arnedo & Gonzalez-Fandos, 2019; Williams & Oyarzabal, 367 2012; Zhang et al., 2018). 368

369 The variation of the prevalence of *Campylobacter* spp. during the different processes' steps can also be explained due to technical aspects of *Campylobacter* detection (sampling 370 procedures, storage, DNA extraction, selection of targeting region, and PCR primers and the 371 sequencing platforms used). Culture-based isolation approaches is considered as a standard 372 method for the detection and enumeration of the different *Campylobacter* spp. of products 373 374 intended for human consumption, animal feeding, environmental samples in the area of food and feed production, and samples from the primary production stage (ISO, 2017). However, the 375 limitations of this technique rely on the difficulties to isolate *Campylobacter* from samples with 376 377 heavy contamination and the rapid loss in cultivability of isolates. To overcome the challenges in traditional phenotype-based methods for the identification of Campylobacter, different DNA-378 based approaches have become widely used due to the speed, and reproducibility to confirm 379 Campylobacter identification (Johannessen et al., 2020). Nevertheless, besides the great 380

advances made in establishing a less time-consuming sampling protocol and more amenable to
 couple with DNA-base methods, currently, there is no standard procedure for a fast screening of
 Campylobacter at the retail level. One also notes that pathogenesis or virulence factors that
 certain *Campylobacter* sequence types may have is an important feature to consider for
 predicting future *Campylobacter* outbreaks and accurate identification in the context of risk
 assessment.

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8 4. CONTROL STRATEGIES

A direct relationship between the reduction of *Campylobacter*'s load at the different broiler 389 production stages and the reduction of public health risk has been linked to an effective control 390 391 strategy. Reducing the numbers of *Campylobacter* on the carcasses by one log₁₀-unit, would reduce the public health risk by between 50 and 90%, and, reducing counts by more than two 392 log₁₀-units would reduce the public health risk by more than 90% (European Food Safety 393 394 Authority Panel on Biological Hazards, 2011). To reduce *Campylobacter* loads, the EU has recently developed the Commission Regulation (EU) 2017/1495, which sets microbiological 395 limits regarding Campylobacter spp. in carcasses of broilers (European Commission, 2017). In 396 recent years several *Campylobacter* control strategies have been developed; most of them 397 focused on the reduction of *Campylobacter* colonization at the farm level which consequently 398 399 decreased *Campylobacter* loads into the slaughterhouse, resulting in a low concentration or absence of the pathogen on the final product (Wagenaar et al., 2006). 400

401 The main strategies to control *Campylobacter* spp. colonization at the farm level is based 402 on the reduction of environmental exposure (biosecurity and hygienic measures), the increase of

broiler resistance to colonization (competitive exclusion, vaccination, application of pre-and pro-403 biotics, organic acids, or phytocompounds etc.), the use of alternative antimicrobials 404 (bacteriophage therapy and bacteriocin treatment), and/or selection of specific breeding to 405 increase the resistance of broiler chickens to colonization (European Food Safety Authority 406 Panel on Biological Hazards, 2011; Umar et al., 2016). Besides, the BIOHAZ sets sanitation 407 408 practices during thinning to prevent *Campylobacter* from entering broiler houses at primary production, and the application and monitor system of the decontamination of carcasses (using 409 chemical or physical treatments) are recommended (EFSA, Panel on Biological Hazards, 2011). 410 At the transportation stage the improvement of hygienic measures by removing feed and 411 litter, cleaning and disinfecting transport crates, are the main strategies studied (Meunier et al., 412 413 2016). However, besides the importance of the transportation step, at the moment the BIOHAZ 414 has not published any recommendation to prevent and/or reduce the contamination of *Campylobacter* during this processing step. At slaughter, dressing, and processing the most 415 common and effective strategies used to reduce Campylobacter loads is the application of 416 417 specific food safety protocols and strict hygienic practices (HACCP), separating Campylobacterinfected flocks from non-infected, physical treatments (scalding, chilling) and chemical 418 419 decontamination of carcasses using chlorine compounds or chlorine-based antimicrobials (Osimani et al., 2017; Silva et al., 2011). One of the disadvantages of using physical treatments 420 421 is that it contributes to the change of organoleptic properties of the food products, which would 422 make them less desirable to the consumers. In addition, physical decontamination is allowed in the United States but not in the EU. 423

In the EU, bacteriophages or bacteriocins in the feed are used to reduce the load of *Campylobacter* in the GI tract of broilers before slaughtering, a reduction of the slaughter age of

broilers, implementation and improvement of the sanitation practices during slaughter (including
the design of adequate equipment with the prevention of fecal leakage), and training food
handlers with better hygienic practices to prevent or reduce the *Campylobacter* colonization in
the slaughter, dressing and processing steps are recommendations made by the EFSA to promote
good processing practices (EFSA, Panel on Biological Hazards, 2011).

431

432 **5. THE APPLICATION OF BACTERIOPHAGES TO COMBAT**

433 *CAMPYLOBACTER* IN BROILER PRODUCTION

434 Despite the extensive efforts from the broiler industry, food safety authorities, and academia, there is no effective, reliable, and practical intervention control strategy able to reduce 435 436 the prevalence of Campylobacter from the farm-to-fork process. As the incidence of antibioticresistant Campylobacter strains is increasing, the development of novel non-antibiotic anti-437 *Campylobacter* treatments is becoming critical (Johnson et al., 2017). Treatment strategies that 438 439 have shown highly promising results for *Campylobacter* control in broiler chickens are currently under development. Among them, the use of specific bacteriophages (phages) as biocontrol 440 agents is considered one of the most promising strategies to reduce the prevalence of 441 *Campylobacter* in the broiler production chain (Atterbury et al., 2003; Carvalho et al., 2010; El-442 443 Shibiny et al., 2009; Fischer et al., 2013; Hammerl et al., 2014). Bacteriophages are viruses that specifically infect and kill bacteria, widely distributed in the environment from the human GI 444 tract to the deep ocean, and often naturally present in animals such as, broilers (Dion et al., 2020; 445 Nafarrate et al., 2021). The interest in using phages as a safety strategy in food production relies 446 447 on its selectivity towards the pathogen of concern, it is harmless to humans, animals, and plants, and does not affect the existing commensal microbiota or alter food properties. 448

449	Campylobacter-specific phage cocktail (phiCcolBB12, phiCcolBB35, and phiCcolBB37) has
450	been applied at pre-slaughter and post-slaughter stages to reduce bacterial loads (Carvalho et al.,
451	2012). In detail, the application of Campylobacter-specific bacteriophages in the broiler
452	production chain has been tested in several studies, focusing on the administration of
453	bacteriophages into the drinking water during the rearing cycle of broilers (Loc Carrillo et al.,
454	2005; El-Shibiny et al., 2009; Fischer et al., 2013; Hammerl et al., 2014; Kittler et al., 2013;
455	Richards, Connerton, & Connerton, 2019) or using phages on raw and processed meat or raw
456	liver (Atterbury et al., 2003; Firlieyanti et al., 2016; Goode et al., 2003).
457	During the rearing cycle of broilers, the reduction rates in the cecal content achieved by the
458	addition of single bacteriophages or bacteriophage cocktails (phage NCTC 12673, 12674, and
459	12678) showed promising potential reducing bacterial loads between 2.5 to 3.2 log CFU/g
460	(Fischer et al., 2013; Kittler et al., 2013). In addition, it was demonstrated that no adverse effects
461	on the broiler microbiota occur after administering a phage cocktail (CP20 and CP30A) in
462	contrast to administering broad-spectrum antibiotic treatments, which can yield dysbiosis in the
463	gut microbiota (Richards et al., 2019). Overall, the results from most of the studies in broilers
464	conclude that bacteriophages (NCTC 12672, 12673, 12674, 12678, 12669, 12671, 12684, CP8,
465	CP34, CP81, Cj6, phiCcolIBB35, phiCcolIBB37, and phiCcolIBB12) can effectively reduce
466	Campylobacter levels if they are administered 24-48 h prior to slaughter (Ushanov et al., 2020).
467	Besides the use of bacteriophages in the primary production, a mean reduction of
468	approximately one log CFU/g of Campylobacter on broiler products was obtained using single
469	bacteriophages (NCTC 12674 and 12673) during meat processing (Atterbury et al., 2003; Goode
470	et al., 2003). In this case, different authors suggest the application of bacteriophages
471	(phiCcolIBB35, phiCcolIBB37, and phiCcolIBB12) at high titers to achieve successful

reductions in Campylobacter counts (Carvalho et al., 2012). In addition, bacteriophages (CP8 472 and CP30) have also shown successful dispersal of Campylobacter biofilms and a reduction of 473 viable cells ranging from one to three log CFU/cm² (Siringan et al., 2011), indicating an 474 additional potential field for phage application to target biofilms in meat processing facilities. 475 From an epidemiological point of view, the bacteriophage treatment in the production chain 476 477 can contribute to a drastic reduction of the infection risk for the consumer. According to the 478 model prediction, phage therapy could lead to a reduction of *Campylobacter* in the fecal matter and on the surface of broiler chicken meat (one log each), resulting on a 90% risk reduction for 479 480 consumer infection (Havelaar et al., 2007). Clinical data suggest the tolerability and/or effectiveness of phage therapy to reduce antibiotic-resistant infections in humans, but also phage 481 482 resistance (El Haddad et al., 2019; Zhvania et al., 2017). Bacteria can promote phage attack 483 mainly through spontaneous chromosomal mutations governed by Darwinian dynamics, leading to the emergence of phage resistance and consequently treatment failure (Luong et al., 2020). 484 485 Identifying new phages with different binding sites to improve efficacy may aid in the prevention of problems related to phage resistance (Wright et al., 2019). 486 More research is needed to find routes of administration, phage selection, the order of phage 487 exposure frequency of administration, dosage, phage resistance, pharmacokinetic and 488 pharmacodynamic properties of the phages, and improve bacteriophage efficacy against 489 *Campylobacter* and broiler meat safety. It is worth noting that bacteriophages should not be 490 considered as a substitute for the control strategies developed so far, but rather seen as a 491 complementary strategy. Successful control of *Campylobacter* could probably be achieved by 492 493 implementing strict biosecurity and hygiene measures in combination with bacteriophage

treatments.

496 **6. CONCLUSION**

The high occurrence of *Campylobacter* along the broiler production chain is a serious threat 497 to public health. This review revealed that abiotic factors have the potential to contribute to 498 cross-contamination of *Campylobacter*. Furthermore, the transfer of contaminated content of the 499 500 bird intestine, or persisting biofilm on equipment/surfaces represents likely the source of crosscontamination during the broiler production. Effective *Campylobacter* control measures along 501 the whole broiler production chain are, therefore, needed to improve broiler meat safety, 502 resulting in a reduction of the incidence of human campylobacteriosis. The use of 503 Campylobacter-specific bacteriophages at different points from farm-to-fork (in livestock, 504 505 slaughter, and/or processing facilities) has been proposed as an additional strategy of a multistage bio-security measure to assure safer chicken products for the consumer. The use of 506 507 multi-omics approaches can help us to increase our understanding of the ability of this foodborne 508 pathogen to persist through the water and the food chain, its environmental niche, and how it interacts with bacteriophages. Progress in this field will help us to better understand how to 509 assess the environmental conditions and nutritional requirements to reduce the risk of 510 *Campylobacter* contamination in the broiler production chain. 511

512

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961 Table legend

Table 1. Prevalence of *Campylobacter* spp, *Campylobacter jejuni* and *Campylobacter coli*from different sample types at different sampling points during the broiler primary production
using culture-based methods for the detection of *Campylobacter* expresses as percentage or log
CFU/g

Table 2. Prevalence of *Campylobacter* spp, *Campylobacter jejuni* and *Campylobacter coli* from fecal samples and different equipment used to transport broilers during the pre-and post transportation of broilers to the slaughterhouse using culture-based methods for the detection of
 Campylobacter expresses as percentage

Table 3. Prevalence of *Campylobacter* spp, *Campylobacter jejuni* and *Campylobacter coli*from different sample types during slaughter, dressing and processing using different types of
polymerase chain reaction for the detection of *Campylobacter* expresses as percentage

974 **Figure legend**

975 Figure 1. Key steps identified and used to assess the prevalence of *Campylobacter* in the

- broiler production chain. The location of studies investigating the effect of bacteriophage is
- 977 indicated with a bacteriophage icon. Color in blue is the prevalence of *Campylobacter* spp. and
- 978 color in yellow is the prevalence of *Campylobacter jejuni*, both expressed as percentage.

Table 1

FACTORS	DESCRIPTION	SAMPLE	CAMPYLOBACTER	C. JEJUNI	C. COLI	REFERENCE
Broiler age						
	14 d	Cloacal	5.00 %	100.00 %		(Ingresa-Capaccioni et al., 2016)
	42 d	Cloacal	62.00 %	67.00 %		(Ingresa-Capaccioni et al., 2016)
	14	Cloacal	0.00 %			(Perez-Arnedo and Gonzalez-Fandos, 2019)
	42 d	Cloacal	0.00 - 100.00 %			(Perez-Arnedo and Gonzalez-Fandos, 2019)
	31 d	Cloacal	39.30 %	46.00 %	26.00 %	(Tang et al., 2020)
	37 d	Cloacal	60.00 %	74.00 %	38.00 %	(Tang et al., 2020)
	7 d (control housing)	Fecal	5.30 %	5.30 %		(Tangkham et al., 2016)
	42 d (control housing)	Fecal	26.00 %	26.00 %		(Tangkham et al., 2016)
	21 d (control housing)	Fecal	83.30 %	18.70 %		(Tangkham et al., 2016)
	7 d (open housing)	Fecal	0.00 %	0.00 %		(Tangkham et al., 2016)
	21 d (open housing)	Fecal	93.30 %	70.70 %		(Tangkham et al., 2016)
	42 d (open housing)	Fecal	37.30 %	00 - 37.30 %		(Tangkham et al., 2016)
Water						
	Environmentally control		0.00 - 66.70 %	0.00 - 16.70 %		(Tangkham et al., 2016)
	Open		0.00 - 83.30 %	0.00 - 83.30 %		(Tangkham et al., 2016)
	Water of 31 d age		0.00 %			(Tang et al., 2020)
	Water of 37 d age		0.00 %			(Tang et al., 2020)
Feed						(Tangkham et al., 2016)
	Environmentally control		0.00 - 83.30 %	0.00 - 33.30 %		(Tangkham et al., 2016)
	Open		0.00 - 33.30 %	0.00 - 33.30 %		(Tangkham et al., 2016)

	Feed of 31 d age		0.00 %		(Tang et al., 2020)
	Feed of 37 d age		0.00 %		(Tang et al., 2020)
itter					
	Covering shoe		20.00 %		(Schroeder et al., 2014)
\ir	Gelatin sample		15.00 %		(Schroeder et al., 2014)
	Air filter samples		0.00 - 10.00 %		(Johannessen et al., 2020)
Workers	Workers' boots swabs		0.00 - 60.00	9%	(Johannessen et al., 2020)
nfrastructure of the	e farm				
			0.00 - 12.50 %		(Bang et al., 2002)
	Floor 31 d age		0.00 %		(Tang et al., 2020)
	Floor 37 d age		0.00 %		(Tang et al., 2020)
	Bedding 31 d age		10.00 %		(Tang et al., 2020)
	Bedding 37 d age		10.00 %	1.00 %	(Tang et al., 2020)
	Sole 31 d age		4.70 %	1.00 %	(Tang et al., 2020)
	Sole 37 d age		10.50 %	2.00 %	(Tang et al., 2020)
	Net 31 d age		0.00 %		(Tang et al., 2020)
	Net 37 d age		0.00 %		(Tang et al., 2020)
	Stool 31 d age		0.00 %		(Tang et al., 2020)
	Stool 37 d age		6.70 %	1.00 %	(Tang et al., 2020)
Management					
	First thinning	Cecal	27.00 - 100.00 %		(Koolman et al., 2014)
	Second thinning	Cecal	90.00 - 100.00 %		(Koolman et al., 2014)
	Drinking water + allicin	Cecal		5.38 log CFU/g	(Robyn et al., 2013)
	Feed				

	Cecal	4.2 - 7.5 log CFU/g		(Gharib Naseri et al., 2012)
	Fecal	6.3 - 7.2 log CFU/g		(Gharib Naseri et al., 2012)
Drink water + probiotics				
	Cecal	4.1 - 6.6 log CFU/g		(Gharib Naseri et al., 2012)
	Fecal	5.4 - 6.4 log CFU/g		(Gharib Naseri et al., 2012)
Feed + plant derivate				
	Cecal	4.2 - 6.3 log CFU/g		(Gharib Naseri et al., 2012)
	Fecal	5.5 - 6.5 log CFU/g		(Gharib Naseri et al., 2012)
Feed + organic acids				
	Cecal	4.0 - 6.2 log CFU/g		(Gharib Naseri et al., 2012)
	Fecal	4.1 - 5.6 log CFU/g		(Gharib Naseri et al., 2012)
Feed + bacteriocin	Fecal	ND		(Stern et al., 2006)
Feed + bacteriophages	Fecal		5.00 log CFU/g	(Carvalho et al., 2010)

980 Abbreviations: C. jejuni; Campylobacter jejuni, C. coli; Campylobacter coli, ND; Not determined

Table 2

		PREVALENCE (%)	PREVALENCE (%)			
FACTORS	DESCRIPTION	CAMPYLOBACTER	C. JEJUNI	C. COLI	REFERENCE	
Equipment						
	Cage		36.80	ND	(Willis et al., 2002)	
	Floor		65.40	ND	(Willis et al., 2002)	
	Pre-transportation cage	6.20 - 30.00		ND	(Stern et al., 2001)	
	Post-transportation cage	42.50 - 85.00		ND	(Stern et al., 2001)	
Fecal						
	Pre-transportation	57.10 - 80.00		ND	(Whyte et al., 2001)	
	Post-transportation	60.00 - 80.00		ND	(Whyte et al., 2001)	

983 Abbreviations: C. jejuni; Campylobacter jejuni, C. coli; Campylobacter coli, ND; Not determined

PRODUCTION	FACTOR	DESCRIPTION	PREVALENCE (%)				REFERENCE
PRODUCTION STAGE	FACTORS	DESCRIPTION	CAMPYLOBACTER	C. JEJUNI	C. COLI	AMPLIFICATION GENE	
Slaughter	Environment	Dirty defeathering swabs	60.00	80.00	20.00	hipO-F, hipoO-R, hipO-P, ceuE-F, ceuE- R, ceuE-R, ceuE-P ⁴	(García-Sánchez e al., 2017)
		Clean defeathering swabs	54.50	90.00	10.00	hipO-F, hipoO-R, hipO-P, ceuE-F, ceuE- R, ceuE-R, ceuE-P ⁴	(García-Sánchez al., 2017)
		Dehairing swabs	31.80	3.00	4.00	16S rRNA, mapA, ceu E^{C}	(Tang et al., 2020
		Dirty evisceration swabs	78.00	100.00	0.00	hipO-F, hipoO-R, hipO-P, ceuE-F, ceuE- R, ceuE-R, ceuE-P ⁴	(García-Sánchez al., 2017)
		Clean evisceration swabs	56.40	100.00	0.00	hipO-F, hipoO-R, hipO-P, ceuE-F, ceuE- R, ceuE-R, ceuE-P ⁴	(García-Sánchez al., 2017)
		Evisceration	31.80	1.00	7.00	16S rRNA, mapA, ceu E^{C}	(Tang et al., 2020
		Dirty sink swabs	100.00	100.00	0.00	hipO-F, hipoO-R, hipO-P, ceuE-F, ceuE- R, ceuE-R, ceuE-P ⁴	(García-Sánchez al., 2017)
		Clean sink swabs	20.00	50.00	50.00	hipO-F, hipoO-R, hipO-P, ceuE-F, ceuE- R, ceuE-R, ceuE-P ⁴	(García-Sánchez al., 2017)
		Dirty floor swabs	22.70	100.00	0.00	hipO-F, hipoO-R, hipO-P, ceuE-F, ceuE- R, ceuE-R, ceuE-P ⁴	(García-Sánchez al., 2017)
		Clean floor swabs	30.00	8330	16.70	hipO-F, hipO-R, hipO-P, ceuE-F, ceuE- R, ceuE-R, ceuE-P ⁴	(García-Sánchez al., 2017)
		Operation table swabs		10.00 - 80.00	0.00 - 100.00	16S rRNA, mapA, ceu E^{C}	(Zhang et al., 20
		Partition	9.10	1.00	1.00	16S rRNA, <i>mapA</i> , $ceuE^{C}$	(Tang et al., 202
		Workers' gloves swabs		20.00 - 60.00	40.00 - 80.00	16S rRNA, <i>mapA</i> , $ceuE^{C}$	(Zhang et al., 20

Table 3

		Cooling	36.40	5.00	5.00	16S rRNA, mapA, $ceuE^{C}$	(Tang et al., 2020)
	Broiler slaughterhouse	Cloacal swabs		0.00 - 63.00	0.00 - 13.60	16S rRNA, mapA, ceu E^{C}	(Zhang et al., 2018)
		Carcass after plucking		0.00 - 73.30	0.00 - 85.70	16S rRNA, mapA, $ceuE^C$	(Zhang et al., 2018)
		Carcass after evisceration		0.00 - 80.00	0.00 - 80.00	16S rRNA, <i>mapA</i> , $ceuE^{C}$	(Zhang et al., 2018
		Carcass after washing		0.00 - 76.70	0.00 - 95.20	16S rRNA, <i>mapA</i> , $ceuE^{C}$	(Zhang et al., 2018
		Carcass after chilling		0.00 - 87.50	0.00 - 100.00	16S rRNA, <i>mapA</i> , $ceuE^C$	(Zhang et al., 2018
		Carcass entrance (41 - 44 days age)	41.40	38.00	22.00	16S rRNA, <i>mapA</i> , <i>ceuE</i> ^{C}	(Tang et al., 2020)
		Carcass after dehairing	12.90	3.00	23.00	16S rRNA, mapA, $ceuE^C$	(Tang et al., 2020)
		Carcass after evisceration	53.40	37.00	69.00	16S rRNA, <i>mapA</i> , ceuE ^C	(Tang et al., 2020)
		Carcass after cooling	14.80	12.00	16.00	16S rRNA, mapA, $ceuE^C$	(Tang et al., 2020)
		Carcass after partition	13.60	8.00	18.00	16S rRNA, mapA, ceu E^{C}	(Tang et al., 2020)
Dressing	Environment	Dirty shackles swabs	41.80	100.00	0.00	hipO-F, hipoO-R, hipO-P, ceuE-F, ceuE- R, ceuE-R, ceuE-P ⁴	(García-Sánchez e al., 2017)
		Clean shackles swabs	38.00	94.70	5.30	hipO-F, hipoO-R, hipO-P, ceuE-F, ceuE- R, ceuE-R, ceuE-P ⁴	(García-Sánchez e al., 2017)
		Dirty conveyor belt swabs	87.90	96.60	3.40	hipO-F, hipoO-R, hipO-P, ceuE-F, ceuE- R, ceuE-R, ceuE-P ⁴	(García-Sánchez e al., 2017)
		Clean conveyor belt swabs	3.30	100.00	0.00	hipO-F, hipoO-R, hipO-P, ceuE-F, ceuE- R, ceuE-R, ceuE-P ⁴	(García-Sánchez et al., 2017)
		Dirty sink swabs	0.00	0.00	0.00	hipO-F, hipoO-R, hipO-P, ceuE-F, ceuE- R, ceuE-R, ceuE-P ⁴	(García-Sánchez e al., 2017)
		Clean sink swabs	0.00	0.00	0.00	hipO-F, hipoO-R, hipO-P, ceuE-F, ceuE- R, ceuE-R, ceuE-P ⁴	(García-Sánchez e al., 2017)

		Dirty floor swabs	9.10	100.00	0.00	hipO-F, hipoO-R, hipO-P, ceuE-F, ceuE- R, ceuE-R, ceuE-P ⁴	(García-Sánchez et al., 2017)
		Clean floor swabs	10.00	100.00	0.00	hipO-F, hipO-R, hipO-P, ceuE-F, ceuE- R, ceuE-R, ceuE-P ⁴	(García-Sánchez e al., 2017)
	Broiler	Carcass processing plant	91.00			Not mentioned ^{<i>B</i>}	(Perez-Arnedo and Gonzalez-Fandos, 2019)
		Skin on thighs	43.50			glyA, hipO ^C	(Casagrande Proietti et al., 2018)
		Skin off breast	44.40			glyA, hip O^{C}	(Casagrande Proietti et al., 2018)
		Legs	54.30			Not mentioned ^B	(Perez-Arnedo and Gonzalez-Fandos, 2019)
		Breast	46.00			Not mentioned ^B	(Perez-Arnedo and Gonzalez-Fandos, 2019)
		Wings	87.00			Not mentioned ^{<i>B</i>}	(Perez-Arnedo and Gonzalez-Fandos, 2019)
Processing	Environment	Chopping board and knives swabs		14.00		hipO ^B	(Khan et al., 2018)
	Partition		13.60	8.00	18.00	16S rRNA, mapA, ceu E^{C}	(Tang et al., 2020)
	Broiler parts	Skin on thighs	51.20			glyA, hipO ^C	(Casagrande Proietti et al., 2018)
		Skin off breast	2.70			glyA, hipO ^C	(Casagrande Proietti et al., 2018)
		Unpacked thighs	51.60			23S rRNA, glyA, hipO, sapB2 ^C	(García-Sánchez e al., 2018)
		Unpacked breast	51.60			23S rRNA, glyA, hipO, sapB2 ^C	(García-Sánchez e al., 2018)
		Unpacked minced	21.90			23S rRNA, glyA, hipO, sapB2 ^C	(García-Sánchez e al., 2018)
		Unpacked marinated	56.30			23S rRNA, glyA, hipO, sapB2 ^C	(García-Sánchez e al., 2018)
		Mean unpacked	45.30			23S rRNA, glyA, hipO, sapB2 ^C	(García-Sánchez e al., 2018)
		Packed thighs	56.30			23S rRNA, glyA, hipO, sapB2 ^C	(García-Sánchez e al., 2018)
		Packed breast	45.30			23S rRNA, glyA, hipO, sapB2 ^C	(García-Sánchez e al., 2018)

Packed minced	14.00			23S rRNA, glyA, hipO, sapB2 ^C	(García-Sánchez e al., 2018)
Packed marinated	18.70			23S rRNA, glyA, hipO, sapB2 ^C	(García-Sánchez e al., 2018)
Mean packed	33.60			23S rRNA, glyA, hipO, sapB2 ^C	(García-Sánchez e al., 2018)
Raw broiler meat		36.00		hipO ^B	(Khan et al., 2018)
Broiler intestine		24.00		hipO ^B	(Khan et al., 2018
Feathers		8.00		hipO ^B	(Khan et al., 2018
Gizzard	59.75			23S rRNA, $mapA$, $ceuE$, $hipO^B$	(Korsak et al., 2015)
Heart	49.66			23S rRNA, mapA, ceuE, hip O^B	(Korsak et al., 2015)
Livers	44.08			23S rRNA, mapA, ceuE, hip O^{B}	(Korsak et al., 2015)
Fillet	61.00			23S rRNA, mapA, ceuE, hip O^{B}	(Korsak et al., 2015)
Breast	39.00			glyA, hipO, ask ^C	(Williams and Oyarzabal, 2012)
Tenderloins	26.00			glyA, hipO, ask ^C	(Williams and Oyarzabal, 2012)
Thighs	53.00 - 90.00	94.50	5.50	glyA, hipO, ask ^{AC}	(García-Sánchez al., 2017; Williams and Oyarzabal, 201

986 Abbreviations: C. jejuni; Campylobacter jejuni, C. coli; Campylobacter coli, ^A Real-Time Polymerase Chain Reaction (PCR), ^BPCR, ^CMultiplex PCR

