



Antimicrobial resistance in the globalized food chain: a One Health perspective applied to the poultry industry

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Abstract

Antimicrobial resistance (AMR) remains a major global public health crisis. The food animal industry will face escalating challenges to increase productivity while minimizing AMR, since the global demand for animal protein has been continuously increasing and food animals play a key role in the global food supply, particularly broiler chickens. As chicken products are sources of low-cost, high-quality protein, poultry production is an important economic driver for livelihood and survival in developed and developing regions. The globalization of the food supply, markedly in the poultry industry, is aligned to the globalization of the whole modern society, with an unprecedented exchange of goods and services, and transit of human populations among regions and countries. Considering the increasing threat posed by AMR, human civilization is faced with a complex, multifaceted problem compromising its future. Actions to mitigate antimicrobial resistance are needed in all sectors of the society at the human, animal, and environmental levels. This review discusses the problems associated with antimicrobial resistance in the globalized food chain, using the poultry sector as a model. We cover critical aspects of the emergence and dissemination of antimicrobial resistance in the poultry industry and their implications to public health in a global perspective. Finally, we provide current insights using the multidisciplinary One Health approach to mitigate AMR at the human-animal-environment interface.

Keywords Aminoglycosides · β -Lactams · Broilers · Fluoroquinolones · Gram-negative · Gram-positive · Tetracyclines

Poultry industry and its role in the global food supply

The consumption of animal protein is the basis of human food. Poultry meat production, which accounts for approximately one-third of the overall meat production worldwide [1, 2], has increased rapidly over the last 50 years. The poultry industry will drive meat production growth

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in the coming decade, and it is projected to account for half of all additional meat produced in that time period [3]. Poultry meat output was forecasted to expand and reach 137 million tonnes in 2020, a 2.6% rise compared to 2019, despite the dampened food services sales during the COVID-19 pandemic, intentional production curbs, and avian influenza outbreaks [4]. The USA, Brazil, the European Union, and China continue to be the leading producers of poultry meat globally [3]. Out of the total meat trade globally (37.6 million tonnes), 14.1 million tonnes (37.5%) relate to poultry meat exported by leading producing countries, making chicken the most exported type of meat. This highlights the great economic importance of the poultry industry as a global commodity, leveraging worldwide economic activities in the industrial, commercial, and service sectors.

The relative affordability is an important driver for the increasing consumer willingness to replace beef and pork by poultry meat. Advantageous economies of scale and cost-efficiencies lead to cascading effects through poultry value chains from production to foreign trade [3]. Therefore, the industrial poultry production has a tradition to be in the vanguard in terms of technological advances in the areas of nutrition, breeding, management, and health.

Moreover, the increasing demand for poultry meat is related to the growing demand for meat protein globally as the human population is expected to reach 9.7 billion in 2050 and 10.9 billion in 2100 [5]. Human population in less developed countries will at least double by 2050, whereas much lower growth rates will occur in developed areas [5]. For example, there is a dramatic increase (725 %) in the projected demand for poultry meat in South Asia by 2030, especially in countries such as India, where poultry meat consumption is estimated to grow from 1.05 to 9.92 million tonnes annually over the next three decades [6].

Given the increasing demand for poultry meat in highly populated urban areas, the poultry industry will move towards more intensified trade activities over the next decades. Technological development in the poultry industry will consolidate as a key driver for the success of the business and for reaching the society needs. These technologies must also improve the capacity of the industry to properly handle the increasing amount of animal waste, such as poultry litter, as it can be a vehicle for the spread of antibiotics, biocides, metabolites, pathogens, and antimicrobial resistance genes (ARG) in the environment [7]. The contamination of soil and water sources, such as rivers and ground- and coastal water, can have a profound and direct impact on the environmental resistome, especially in countries under poor sanitation conditions [7, 8]. In this scope, the technological development of the poultry industry should be embedded in Environmental, Social, and Governance (ESG) integrated initiatives for increasing performance and sustainability.

The concept of antimicrobial resistance and its burden to public health

Antimicrobial resistance is a natural, ancient, and common phenomenon in bacteria inhabiting any biological system [9]. However, some factors can favor the emergence and spread of antimicrobial resistance among bacteria. Anthropogenic factors seem to play an important role in this aspect, such as the overuse and misuse of antimicrobial agents, which generates selection pressure favoring the multiplication of antimicrobial-resistant organisms [10]. Antimicrobial agents can eliminate or inhibit the growth of susceptible bacteria, allowing resistant strains to colonize and multiply in a given environment. Several experimental, epidemiological, and ecological studies have confirmed this hypothesis [11]. Therefore, antimicrobial-resistant pathogens have the potential to cause hard-to-treat diseases, increased mortality rates, and economic burden. Infections caused by antimicrobial-resistant pathogens are considered major global public health crisis by the World Health Organization as the discovery of effective antimicrobial drugs does not keep pace with the increasing antimicrobial resistance rates in bacteria.

The World Bank estimated that the loss of the annual global gross domestic product (GDP) by 2050 would fall 1.1% in the best case scenario of low AMR impact to 3.8% in the worst case scenario, which translates to an annual shortfall of \$3.4 trillion by 2030 [12]. The O'Neill report on Review on Antimicrobial Resistance estimated the burden to be 10 million lives per year and cumulative 100 trillion USD of economic output by 2050 [13]. An estimation of global infections caused by drug-resistant WHO priority pathogens placed third-generation cephalosporin-resistant *Escherichia coli* and *Klebsiella pneumoniae* to be responsible for approximately 4.6 to 6.4 million bloodstream infections in 2014 and carbapenem-resistant ones for approximately 0.4 to 0.5 million bloodstream infections [14].

Important considerations on antimicrobials used in the poultry industry

In quantitative terms, the amount of antimicrobial drugs used in livestock globally was estimated in 63,151 tonnes in 2010, and it is expected to increase 67% by 2030, reaching approximately 105,500 tonnes [15] following the rising demand for livestock products by the human population in middle-income countries. The amount of antimicrobials used in livestock is estimated to nearly double in countries such as the BRICS countries (Brazil, Russia,

India, China, and South Africa). In spite of all efforts and competence of researchers working in this field, accurate numbers on the amount of antimicrobials used in food animals are extremely difficult to be reached because of lack of information and complexity of available data. There are important initiatives addressing data acquisition and availability, especially those led by the World Organization for Animal Health (OIE) database that serves as a global reference to national systems for monitoring the use of antimicrobials in animals. Nevertheless, these initiatives are at an early stage of development and face innumerable challenges related to the inaccessibility of data or calculation errors from many countries that are unable to provide reasonable accurate information [16]. Therefore, current calculations in terms of the amount of antimicrobials used in animals seem to be underestimated.

According to the 2016 OIE report [16], data collected from 93 countries globally indicated that Americas, Asia, and East Oceania used 87.13% from 97,784 tonnes of antimicrobial agents intended for animal use that year, even though only 18 countries were able to distinguish which drugs were used in terrestrial food animals. Tetracyclines (33.8%) and penicillins (20.4%) were identified as the most frequently used drugs in these animals. According to this same report, increased estimates of antimicrobial use were observed after adjustments by animal biomass, as shown in Table 1 [16].

Antimicrobial drugs have played an important role in intensive animal production systems, and a wide variety of antimicrobials can be used in poultry production [17, 18]. Antimicrobial therapy of diseased animals is not only the most promising alternative to overcome the economic burden caused by bacterial infectious diseases in livestock, but also represents a welfare principle that must be used to avoid suffering of any food animal. Therefore, the

success of poultry farming and other animal production systems will continue to depend on the existence of effective antimicrobials.

Besides therapy, antimicrobial drugs have been also used in poultry production for other purposes, such as metaphylaxis, i.e., the prophylactic treatment of the entire flock including animals that show no clinical sign of the disease in order to prevent the spread of a given disease, or as growth promoters, which are antimicrobials added to animal feed at low dosages to enhance animal performance. Although the use of antimicrobial drugs represents a driver for antimicrobial resistance in bacterial populations of every microenvironment using increased selective pressure, this phenomenon is significantly increased when antimicrobials are misused. Therefore, how antimicrobials are used in the food animal industry in terms of antimicrobial classes, the doses, and the purpose could have a great impact on the emergence and dissemination of antimicrobial resistance. While specific approved antimicrobials in veterinary medicine have been used therapeutically following veterinary recommendations, misuse of antimicrobials in food animals has been reported, such as the indiscriminate use of unprescribed drugs added to animal feed [19] or even added to vaccines [20] aiming to prevent disease and to improve animal performance. Thus, the use of antimicrobials in food animals should be also assessed in a qualitative perspective, given the fact that the non-therapeutic administration of antimicrobial drugs to livestock at sub-inhibitory dosages are supposed to have a greater impact on the emergence and dissemination of antimicrobial resistance. The problem is critical when these drugs relate to antimicrobial classes that are considered critically important to human health [21], which makes the non-therapeutic use of these drugs a global health concern. There are indications that antimicrobials in animal food production are directly related to the increase of antimicrobial resistance [22–25].

Importantly, there is a lack of new active principles or candidate interventions capable of overcoming resistance mechanisms developed by bacteria. Therefore, the rational use of these drugs seems the most adequate alternative to guarantee the effectiveness of already existing antimicrobials for treatment of diseases in animals and humans.

As also occurring in other sectors, certain animal production practices might contribute to the spread of antimicrobial-resistant bacteria. These practices include overprescription of broad-spectrum antimicrobials to animals [26], feeding of certain antibiotics at low doses for growth promotion purposes [27, 28], and the use of non-approved or off-label use of drugs [29].

However, this drug overuse might be deposited as antibiotic residues in foodstuffs such as meat, eggs, and milk, to name a few [30–33]. Several side effects have been related to antibiotic residues in foodstuffs, such as toxicity (bone

Table 1 Amounts of antimicrobial drugs used in food animals by animal biomass before and after adjustments, by country estimates of data coverage. Table was adapted from OIE report of the year 2020 [16].

OIE region	Countries*	Antimicrobial amounts used by animal biomass	
		Not adjusted (mg/kg)	After adjustment (mg/kg)
Africa	21	39.17	45.25
Americas	11	114.54	138.07
Asia, Far East, and Oceania	19	237.72	240.57
Europe	40	66.91	68.55

*Number of countries providing data for food animals in all rounds of data collection for 2016, from 4 OIE regions

marrow and hepatotoxicity), nephropathy, immunopathological effects (allergy), mutagenicity, and, especially, transfer of antibiotic resistance between MDR and commensal bacteria that may contribute to the imbalance of gut microbiota posing hazards to human health [30–33].

To address this concern effectively, some actions delivered by all sectors are needed to avoid the spread of these antimicrobial residues in foodstuffs. First, farmers must have the drug withdrawal period as a priority to safeguard human health. Second, the poultry industries must establish measurement strategies to detect antimicrobials in poultry products and related sources, including quantitative (liquid chromatography, capillary electrophoresis, enzyme linked immunosorbent assay), semi-quantitative (microbiology assay), and qualitative (x-ray diffraction) techniques [33]. Lastly, food authorities must be aware and monitor the previous steps made by the whole food chain sector.

Emergence and spread of antimicrobial resistance in the poultry production: a two-faceted problem

As in other sectors, the use of drugs in poultry production eliminates susceptible bacteria favoring the selection of resistant isolates. These persistent isolates become

predominant and can transfer their resistance genes to both clonal descendants and also to other isolates from the same bacterial species or even from other species [34]. In this context, the problem of antimicrobial resistance in poultry production needs to be understood from two different perspectives: (i) the emergence and dissemination of antimicrobial-resistant zoonotic pathogens that represent a direct hazard in public health and (ii) the emergence and dissemination of commensal bacteria carrying ARGs in mobile genetic elements that can be further contributed as source of important resistance genes potentially transferred to pathogenic bacteria within a given host or in the environment [7, 8]. This explains the high complexity of antimicrobial resistance in the poultry industry, as illustrated in Fig. 1.

Another critical issue in the acquisition of resistance mechanisms is the identification of transmission routes of resistant bacteria along the food chain. Although it remains difficult to identify when the contamination begins, several insights can be denoted in One Health framework, including risk factors associated with the farm environment, food industry, and household settings as summarized in Fig. 1. Among these factors, transmission via vectors, water, soil, plants, animal feed, poultry, and anthropological actions play a crucial role to the development and spread of antimicrobial resistance in a global scale [36].

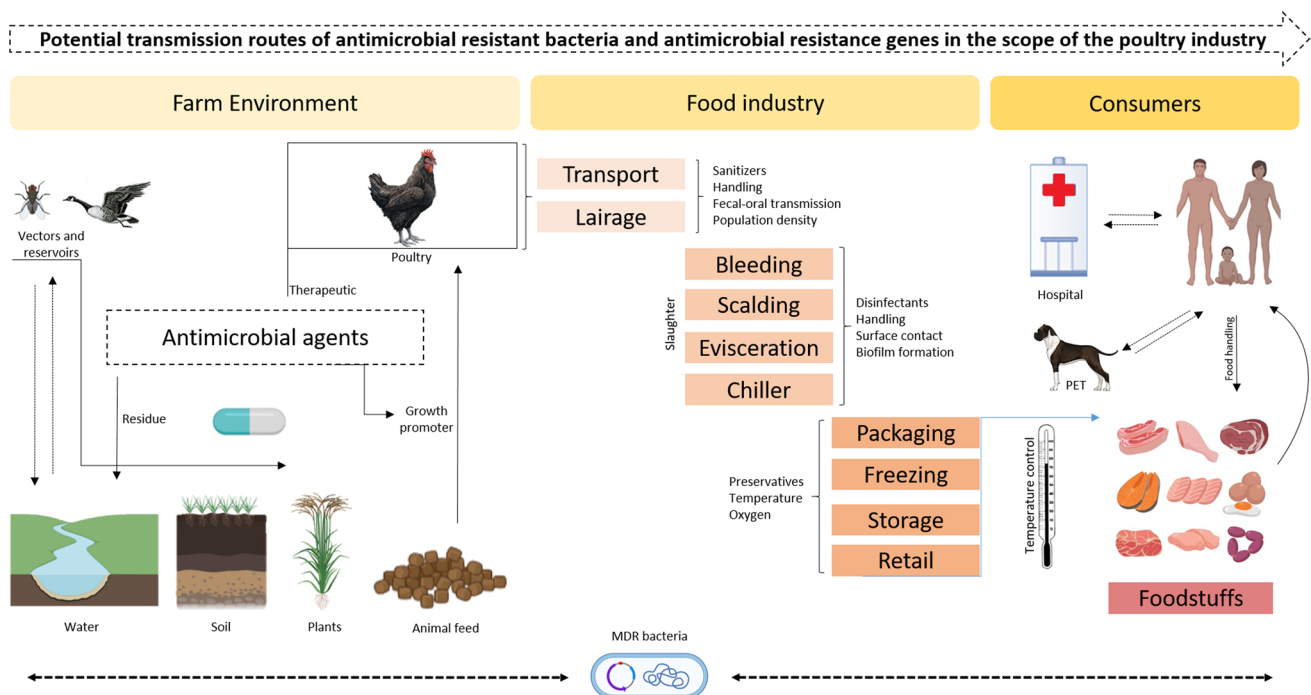


Fig. 1 Potential transmission routes of antimicrobial-resistant bacteria and antimicrobial resistance genes in the scope of the poultry industry (adapted from Monte et al. [35]). This figure was created using BioRender.com.

Antimicrobial resistance in zoonotic pathogens associated with poultry products

A wide variety of antimicrobial-resistant bacteria of public health significance has been detected in broilers and laying hens. In this topic, we describe antimicrobial resistance levels among the most relevant zoonotic pathogens associated with chickens and poultry products: *Salmonella* spp. [37–39], *Campylobacter* spp. [37, 40, 41], *Enterococcus* spp. [42–44], *Escherichia coli* [43, 45], and *Staphylococcus aureus* [42, 46].

Salmonella spp.

Salmonellosis affects hundreds of people around the world, causing high fever and diarrhea that can progress to death [47]. Because of its global importance, many countries have implemented surveillance programs for *Salmonella* in livestock, allowing the acquisition of important information about antimicrobial resistance. More than 2,650 serovars of *Salmonella enterica* have been already identified [48], and several are found in foods such as chicken meat and eggs, considered key sources of infection and concern for public health [49–51]. In chickens, resistant *Salmonella* spp. not only can cause economic losses, but they can potentially cause occupational salmonellosis in farmers and keepers [52–54].

Salmonella diseases in birds can be divided into three distinct conditions: fowl typhoid, caused by *Salmonella Gallinarum* biovar Gallinarum (SG); pullorum disease, caused by *Salmonella Gallinarum* biovar Pullorum (SP); and avian paratyphoid, resulting from infection by any other serovar except those previously mentioned [55]. SG and SP only affect chickens and cause severe clinical signs and mortality, while SG affects birds of any age, and SP-associated infections occur in young birds. Chickens of light lineages are more resistant to fowl disease compared to semi-heavy and heavy lineages [56]. In contrast, paratyphoid *Salmonella* infections do not lead to clinical signs, but there are exceptions that can cause disease and mortality in chicks [55].

The epidemiology of salmonellosis is complex. The high density in the poultry houses along with the persistence of the microorganism in water, feed ingredients, litter, and fomites as well as in biological vectors on the farm environment facilitate its dissemination [57]. Transmission occurs through horizontal and vertical routes in the case of pullorum disease and avian paratyphoid [55, 58], and only by horizontal route in fowl typhoid [59].

A previous study on semi-heavy laying hens showed that treatment with antimicrobials was able to reduce

mortality caused by fowl typhoid only temporarily; death and compromised egg production were observed after treatment was suspended [59]. Thus, the efficacy of antimicrobials in controlling disease is limited and can promote the emergence of resistant *Salmonella* strains [55, 59]. In the case of paratyphoid *Salmonella*, multi-resistance is of concern as it promotes failures in the treatment of infection in humans who have ingested chicken meat or eggs contaminated by zoonotic serotypes such as *S. Enteritidis*, *S. Typhimurium*, *S. Heidelberg*, *S. Hadar*, *S. Infantis*, *S. Mbandaka*, *S. Saintpaul*, and *S. Newport*, among others [60–62]. A high and diverse level of antimicrobial resistance was identified in approximately 4,000 isolates from 14 different studies (Table 2). Nalidixic acid (95.2%), amoxicillin (91.6%), ampicillin (100%), erythromycin (100%), penicillin G (100%), sulfamethoxazole (100%), and tetracycline (93%) showed the highest resistances, while higher sensitivities were obtained for aminoglycosides (spectinomycin - 96% and gentamicin - 99.8%).

The emergence and spread of antimicrobial-resistant, in particular multidrug-resistant *Salmonella* strains (MDR strains), is a major public health concern [63]. While *Salmonella* continues to be a major cause of foodborne disease in most countries, the reduced number of antimicrobials available for therapeutic purposes in humans and animals exacerbates the issue [64]. The proportion of *Salmonella* isolates resistant to ceftriaxone and ciprofloxacin in the USA has increased year on year [65]. MDR *Salmonella* recovered from chicken samples during veterinary inspection service in the USA has increased, remarkably for *Salmonella* Infantis [65]. Nontyphoidal MDR *Salmonella* is estimated to be responsible for 20,800 infections per year in the USA [66]. The most common antimicrobial resistance patterns of multidrug-resistant *Salmonella* strains related to important therapeutic antimicrobial classes are used in humans, including penicillins, tetracyclines, cephalosporins, and fluoroquinolones [28].

The frequency and extent of resistance in *Salmonella* varies according to the antimicrobial use and the ecological differences in its epidemiology [67]. Some multidrug-resistant *Salmonella* strains with extensive resistance profiles pose greater threats to public health and have been associated with higher mortality and morbidity rates [41]. Multidrug resistance in *Salmonella* spp. to ampicillin, chloramphenicol, streptomycin, sulfonamides, and tetracycline (ACSSuT) has been associated with invasive disease in humans [64]. Another multidrug-resistant pattern with additional resistance to amoxicillin-clavulanic acid and ceftriaxone (ACSSuTAuCx) had been associated with severe disease in humans [68], while the monophasic ST35 variant is associated to colistin resistance [69].

Table 2 Summary of antimicrobial resistance results in *Salmonella* spp. isolated from poultry published in the last 5 years, within the five continents.

Study	Region	Reference	Publishing year	No. of isolates	Resistance profile (%)
1	Africa	Abd-Elghany et al.	2015	166	AMP (74,7), AMX (91,6), C (47), CIP (19,3), ERY (100), GM (1,2), KA (9,6), NAL (95,2), NEO (61,4), NOR (30,1), OT (89,2), P (92,8), S (67,5), SUL (83,1)
2	Africa	Zishiri et al.	2016	102	AMP (47), AMX (31), C (31), ERY (18), GM (48), KA (74), S (12), SXT (84,3), T (93), TMP (78,4)
3	Africa	Egualé	2018	26	AMC (42,3), AMP (42,3), C (42,3), CFN (46,2), CIP (7,7), FOX (0), GM (7,7), KA (42,3), NAL (19), NEO (11,5), NIT (26,7), S (92,3), SUL (92,3), SXT (3,9), T (30,8), TMP (3,9)
4	North America	Ladely et al.	2016	S. Kentucky 600	AK (0), AMC (15,3), AMP (16,7), C (2), CEF (14,8), CIP (0), CRO (15,2), FOX (14,3), GM (2), KA (1,3), NAL (0,3), S (46,5), SUL (3,5), SXT (0,2), T (50)
5	North America	Nisar et al.	2017	19	AMC (15,8), AMP (10,5), AZI (0), C (5,2), CEF (5,2), CIP (0), FOX (10,5), GM (10,5), NAL (0), S (21,1), SUL (21,1), SXT (0), T (21,1)
6	South America	Donado-Godoy et al.	2014	39	AMP (82), CEF (74), CIP (69), CTX (69)
7	South America	Vinueza-Burgos et al.	2016	62	AMP (67,7), C (64,5), CAZ (6,5), CIP (83,9), COL (16,1), CTX (71), FFC (67,7), GM (66,1), KA (51,6), NAL (85,5), S (79), SUL (88,7), T (80,6), TMP (80,6)
2	South America	Zishiri et al.	2016	24	AMP (100), AMX (83), C (4,2), ERY (62,5), GM (12,5), KA (16,7), S (12,5), SXT (50), T (83), TMP (66,7)
8	South America	Cunha-Neto et al.	2018	31	AMP (25), AZI (21,8), C (3,1), CEF (6,3), CFN (25), CIP (0), ENR (0), FFC (0), FOX (18,8), GM (3,1), NAL (0), NIT (0), S (0), SUL (100), SXT (75), T (9,4), TMP (87,5)
9	Asia	Thung et al.	2016	11	AMC (0), AMP (72,7), AMX (27,3), CAZ (0), CFZ (27,3), CIP (27,3), ERY (100), GM (0), KA (0), NAL (9,1), P (100), S (9,1), T (0), TMP (0), VAN (100)
10	Asia	Trongjit et al.	2017	375	AMP (70,7), C (10,7), CAZ (4,3), CIP (0,6), CPD (4,9), CTX (3,8), GM (2,9), S (28), SUL (69,5), T (26), TMP (29,6)
11	Asia	Uddin et al.	2018	150	AMC (30), AMP (75), AZI (70), BAC (56), C (41), CE (0), CIP (4), CRO (0), ERY (40), FEP (0), KA (8), NAL (50), NEO (80), NV (74), S (31), SPT (4), T (89), TZP (0)
12	Europe	de Jong et al.	2014	42	AMC (2,9), AMP (35,4), C (3,3), CIP (0), COL (10,4), CTX (13,1), GM (0,2), NAL (22,8), S (24,3), SXT (26,8), T (15)
13	Europe	Franco et al.	2015	42	AMP (78,6), C (4,8), CIP (83,3), CTX (71,4), GM (2,4), KA (40,5), NAL (83,3), S (19), SUL (85,7), T (85,7), TMP (83,3)
14	Europe	EFSA	2016	2210	AMP (19,1), AZI (1,9), C (4), CAZ (2,6), CIP (53,5), COL (8,3), CTX (2,3), GM (6,6), NAL (48,7), SUL (45,1), T (40,4), TGC (9,3), TMP (16,9)

AK amikacin, AMC amoxicillin/clavulanic acid, AMP ampicillin, AMX amoxicillin, AZI azithromycin, BAC bacitracin, C chloramphenicol, CAZ ceftazidime, CE cefradine, CEF ceftiofur, CFN cephalothin, CFZ cefazoline, CIP ciprofloxacin, CLX cefalexin, COL colistin, CPD cefpodoxime, CRO ceftriaxone, CTX cefotaxime, ENR enrofloxacin, ERY erythromycin, FEP cefepime, FFC florfenicol, FOX cefoxitin, GM gentamicin, KA kanamycin, NAL nalidixic acid, NEO neomycin, NIT nitrofurantoin, NOR norfloxacin, NV novobiocin, OT oxytetracycline, P penicillin, S streptomycin, SP spermidine, SPT spectinomycin, SUL sulfamethoxazole, SXT sulfamethoxazole/trimethoprim, T tetracycline, TGC tigecycline, TMP trimethoprim, TZP piperacillin/tazobactam, VAN vancomycin

Campylobacter spp.

Campylobacter species can cause disease in animals and humans [70] and are a major cause of foodborne diseases worldwide. Both *Campylobacter jejuni* (*C. jejuni*) and *C. coli* can colonize the gut of broilers. Human infections, of

particular concern when involving children [71], are usually caused by consumption of contaminated poultry products, even though occupational transmission has been observed.

Due to the high occurrence of *Campylobacter* spp. in the environment, including soil and water, and the high diversity of susceptible vector hosts [72, 73], the epidemiology is

very complex. Wild animals and pests, as well as humans and vehicles, can introduce *Campylobacter* spp. onto the farm [38]. Even though the real role of the feed as source of *Campylobacter* is unclear [41], the bacteria can survive and multiply in foods under different storage temperatures [74]. High morbidity can be observed in domestic birds, and transmission occurs through both horizontal and vertical routes [41, 72, 75].

When infecting the bird, *Campylobacter* spp. effectively colonizes the intestine, mainly the mucosa of the cecal crypts. It can invade the intestinal epithelium and multiply rapidly in the intestinal mucus to avoid clearance [74]. In addition, it is able to persist under commensal conditions due to the weak immune response of the animal, making the bird a reservoir of human campylobacteriosis [41, 72].

Antimicrobial therapy, besides being ineffective in eliminating *Campylobacter* due to its high occurrence and commensal relationship in birds [41], can promote the emergence of resistant strains. Therefore, biosecurity practices are the most appropriate way to reduce the infection at the flock level [72]. Other approaches such as the use of organic acids to reduce the pH of the gastrointestinal tract and competitive exclusion can reduce intestinal colonization by the microorganism [41, 75].

In nine recent studies, antimicrobial resistance profiles of more than 2,000 isolates of *Campylobacter* spp. were evaluated in broilers [76–79], laying hens [77], chicken carcasses [77, 80–82], and chicken meat or offal [81, 83, 84]. High frequencies of resistance to nalidixic acid (100–38.5%), ampicillin (100–8.8%), cephalixin (100–85.7%), ciprofloxacin

Table 3 Summary of antimicrobial resistance results in *Campylobacter* spp. isolated from poultry published in the last 5 years, within the five continents.

Study	Region	Reference	Publishing year	No. of isolates	Resistance profile (%)
1	Africa	Karikari et al.	2017	<i>C. jejuni</i> 39	AMP (100), C (84,6), CIP (69,2), CLX (100), CTX (87,2), ERY (100), GM (23,1), KA (28,2), NAL (38,5), NOR (48,7), SXT (92,3), T (100)
				<i>C. coli</i> 14	AMP (100), C (64,3), CIP (64,3), CLX (85,7), CTX (85,7), ERY (100), GM (0), KA (28,6), NAL (7,4), NOR (0), SXT (92,9), T (92,9)
				<i>C. lari</i> 9	AMP (100), C (55,6), CIP (100), CLX (100), CTX (100), ERY (100), GM (33,3), KA (33,3), NAL (100), NOR (77,8), SXT (88,9), T (100)
2	North America	Kassem et al.	2017	<i>C. jejuni</i> 45	CIP (15,6), ERY (20), S (11,1), T (64,4), TYL (33,3)
				<i>C. coli</i> 155	CIP (7,1), ERY (26,5), S (10,3), T (66,5), TYL (28,4)
3	South America	Donado-Godoy et al.	2014	29	CIP (97), ERY (21), T (83)
4	South America	Panzenhagen et al.	2016	<i>C. jejuni</i> 44 <i>C. coli</i> 38	CIP (100), ENR (100) CIP (100), ENR (100)
5	South America	Vinueza-Burgos et al.	2017	<i>C. jejuni</i> 48	CIP (97,9), ERY (4,2), GM (2,1), NAL (100), S (8,4), T (83,4)
				<i>C. coli</i> 170	CIP (100), ERY (25,8), GM (1,2), NAL (99,3), S (11,2), T (67,6)
6	Asia	Han et al.	2016	<i>C. jejuni</i> 160	CIP (88,1), CLD (68,1), ERY (14,3), FFN (16,3), GM (47,5), LEV (78,1), S (43,8), T (79,4)
				<i>C. coli</i> 130	CIP (100), CLD (98,5), ERY (93,5), FFN (8,5), GM (90,8), LEV (98,5), S (17,8), T (98,5)
7	Asia	Kottawatta et al.	2017	<i>C. jejuni</i> 20	AMP (45), C (0), CIP (80), CLA (0), ERY (5), GM (10), NAL (80), NEO (10), S (5), SUL (0), T (85), TUL (5)
				<i>C. coli</i> 45	AMP (8,8), C (2,2), CIP (84,4), CLA (6,6), ERY (11,1), GM (2,2), NAL (84,4), NEO (8,8), S (4,4), SUL (2,2), T (24,4), TUL (6,6)
8	Europe	Wieczorek e Osek	2015	<i>C. jejuni</i> 576	CIP (74,8), ERY (0,9), T (46,5)
				<i>C. coli</i> 575	CIP (88,3), ERY (4), T (65,7)
9	Europe	García-Sánchez et al.	2018	<i>C. jejuni</i> 55	AZI (96), CIP (100), ERY (100), GM (100), NAL (100), T (98)
				<i>C. coli</i> 19	AZI (63), CIP (79), ERY (63), GM (95), NAL (100), T (100)
				<i>C. lari</i> 2	AZI (0), CIP (100), ERY (0), GM (50), NAL (100), T (100)

AMP ampicillin, AZI azithromycin, C chloramphenicol, CIP ciprofloxacin, CLA clarithromycin, CLD clindamycin, CLX cefalexin, CTX cefotaxime, ENR enrofloxacin, ERY erythromycin, FFN florfenicol, GM gentamicin, KA kanamycin, LEV levofloxacin, NAL nalidixic acid, NEO neomycin, NOR norfloxacin, S streptomycin, SUL sulfamethoxazole, SXT sulfamethoxazole/trimethoprim, T tetracycline, TUL tulathromycin

(100-7.1), erythromycin (100-0.9%), gentamicin (100-0%), and tetracycline (100-24.4%) were detected (Table 3).

Enterococcus spp.

Enterococcus species are part of the natural gut microbiota of both humans and animals and frequently contaminate the environment and foods, including chicken meat [85]. Although some strains have been used in probiotics, others have been reported as pathogenic strains for chickens [86]. In this case, there might be mortality without clinical signs during the acute phase, ranging from 1 to 3 weeks after infection [86]. In the case of clinical manifestation, septicemia is the most common observed condition. In general, transmission occurs via both horizontal and vertical routes. Among the pathogenic specimens for domestic birds, *E. cecorum* and *E. faecalis* can be highlighted [86].

E. cecorum has grown in importance from the past two decades after the epidemic of pathogenic strains in broilers, causing the so-called free thoracic vertebra (FTV) osteomyelitis [87]. The infection causes abscesses in the thoracic vertebrae that compress the spinal cord, leading the bird to remain supported under the tibiotarso-tarsometatarsal joint

resulting from symmetrical paralysis of the posterior limbs [88, 89]. In addition, septicemia may occur with the appearance of fibrinoid pericarditis, hepatitis, and splenomegaly, which can progress to death [90]. Importantly, pathogenic strains are able to colonize the gut of birds in earlier development states compared with commensal strains, which has been attributed as an important driver for the occurrence of the disease [88].

On the other hand, *E. faecalis* has been associated with omphalitis and yolk sacculitis followed by sepsis and death of chicks in the first week of life, which may or may not be associated with *Escherichia coli*. Surviving animals may develop valve endocarditis and die in the chronic phase of the infection [86].

Enterococcus spp. can easily acquire resistance to antimicrobials and, therefore, play a central role in its dissemination [86]. Pathogenic *E. cecorum* strains are more resistant to antimicrobials than commensals [90]. Vancomycin-resistant *Enterococcus* spp. strains have been associated with economic losses in animal production and also with nosocomial diseases in humans [91–93].

Studies performed between 2014 and 2017 identified the presence of Enterococci species in broilers, laying hens [74,

Table 4 Summary of antimicrobial resistance results in *Enterococcus* spp. isolated from poultry published in the last 5 years, within the five continents.

Study	Region	Reference	Publishing year	No. of isolates	Resistance profile (%)
1	Africa	Ngbede et al.	2016	53	AMP (43,4), C (9,4), CIP (11,3), ERY (73,5), GM (47,1), QD (9,4), RIF (15,1), S (47,1), T (81,1), VAN (0)
2	North America	Kilonzo-Nthenge et al.	2015	132	AK (19,7), AMC (10,6), AMP (14,4), AZI (6,8), C (3), CEC (3), ERY (62,9), FOX (7,6), KA (47), P (52,3), S (81,8), T (44,7)
3	North America	Tyson et al.	2017	5455	C (0,3), CIP (17,8), DAP (1,9), ERY (28,1), GM (6,6), KA (19,8), LZD (0,1), NIT (24,7), P (13,3), QD (20,5), S (16,6), T (60), TGC (0,1), TYL (27,2), VAN (0)
4	South America	Donado-Godoy et al.	2014	<i>E. faecalis</i> 121 <i>E. faecium</i> 60	AMP (0), CIP (32), ERY (86), VAN (2) AMP (0), CIP (33), ERY (62), QD (60), VAN (0)
5	South America	Cavaco et al.	2017	<i>E. faecalis</i> 40	C (2,5), CIP (2,5), ERY (7,5) FFC (2,5), GM (2,5), KA (2,5), LZD (7,5), S (5), T (7,5), TZD (7,5)
6	South America	Braga et al.	2017	<i>E. faecalis</i> 12	AMP (0), AMX (0), BAC (0), CEF (8,3), GM (33,3), NEO (25), PNV (0), VAN (8,3)
7	Asia	Usui et al.	2014	<i>E. faecalis</i> 117	AMP (0), C (27,4), ENR (17,9), ERY (70,9), GM (11,1), KA (37,6), MY (73,5), OT (69,2), S (51,3), VAN (0)
8	Europe	Maajost et al.	2015	<i>E. faecalis</i> 126 <i>E. faecium</i> 18	AMC (0), AMP (0), C (8), CIP (5), ERY (44), GM (44), MY (99), P (0), T (82), TYL (44), VAN (0) AMC (17), AMP (28), C (6), CIP (56), ERY (61), GM (72), MY (72), P (44), T (67), TYL (56), VAN (0)
9	Europe	Stepien-Pysnick	2016	911	AMC (4), AMX (4,4), DOX (67,3), ENR (69,4), FFC (15,7), MY (56,1), SXT (88), TYL (71,4), VAN (0,1)

AK amikacin, AMC amoxicillin/clavulanic acid, AMP ampicillin, AMX amoxicillin, AZI azithromycin, BAC bacitracin, C chloramphenicol, CEC cefaclor, CEF ceftiofur, CIP ciprofloxacin, DAP daptomycin, DOX doxycycline, ENR enrofloxacin, ERY erythromycin, FFC florfenicol, FOX cefoxitin, GM gentamicin, KA kanamycin, LZD linezolid, MY lincomycin, NEO neomycin, NIT nitrofurantoin, OT oxytetracycline, P penicillin, PNV penicillin/novobiocin, QD quinupristin/dalfopristin, RIF rifampicin, S streptomycin, SXT sulfamethoxazole/trimethoprim, T tetracycline, TGC tigecycline, TYL tyrosine, TZD tedizolid, VAN vancomycin

94–97], and chicken meat [98–100], with frequencies ranging from 30.7 to 92%. High levels of resistance (Table 4) were found against drugs belonging to the class of aminoglycosides (streptomycin, 81.8–5%), tetracyclines (doxycycline, 67.3%; tetracycline, 82–7.5%), and quinolones (ciprofloxacin 97–2.5%; enrofloxacin, 69–17.9%). Vancomycin resistance was low (2–0.1%), with higher levels observed in isolates from chickens affected with vertebral osteomyelitis [101].

Escherichia coli

Escherichia coli (*E. coli*), a Gram-negative species belonging to the Enterobacteriaceae family, is of particular importance in the human-animal-environment triad. This species is often studied as a marker for antimicrobial resistance, mainly because of its wide distribution and its capacity to harbor several resistance genes in mobile genetic elements, therefore serving as a source of antimicrobial resistance genes to pathogenic strains [25, 102, 103]. In domestic birds, avian pathogenic *E. coli* (APEC) are causative agents of colibacillosis [104–106]. The number of serotypes causing the disease has been increasing [107], although O1 and O78 remain as the most commonly detected serovars [104].

Colibacillosis is a complex local or systemic syndrome, in which the pathogen is easily transmitted orally, by inhalation or vertical route [108, 109]. *E. coli* containing several virulence genes are shown to lead to persistent and severe clinical diseases in birds [107]. In general, it is considered an opportunistic infection, occurring after the immune response becoming reduced due to stressful situations to the animal, immunosuppression, co-infections, and environmental problems (e.g., excess of ammonia), among others. However, more recently it has also been considered a primary pathogen that causes high economic loss [104, 109]. As for other bacteria species, its epidemiology is complex. Although pathogenic *E. coli* can be introduced onto the farm by different routes, the contact with wild birds, insects, and rodents [108] is of special importance.

The incubation period is short, and birds can be asymptomatic and remain as such until death; in case of coli septicemia, birds become apathetic or moribund [108]. Clinical signs vary according to the syndrome, but the different conditions can result in localized anatomopathological changes, such as coliform omphalitis, coliform cellulitis, swollen head syndrome, and salpingoperitonitis in adults, or systemic such as colisepticemia, respiratory colisepticemia (aerosaculitis, chronic respiratory and complicated chronic respiratory diseases), osteoarthritis, synovitis, and coligranuloma (Hjarre's disease). They can lead to productive losses, ranging from rejects and carcass condemnation to death [108, 109]. Young birds are more susceptible to the pathogen, and high mortality rates are observed in embryos [108].

The success of vaccines has been challenged by the extensive number of serotypes. Therefore, the control of colibacillosis is very dependent of biosecurity measures [108]. The use of bacteriophages has been shown to help prevent colibacillosis with initial development in the respiratory tract and air sacs [110], and *in ovo* administration of bacteriophages showed similar effect to treatment with enrofloxacin [111]. However, the use of bacteriophages is limited due to the type, bird age, administration route, dose used, and the fact that the elimination of the bacteria in the host can result in the release of endotoxins that lead to fever and toxic shock [112]. Control of colibacillosis with antimicrobials is a common practice, but there are problems resulting from the acquisition of resistance by microorganisms [112], since resistant strains colonizing the gastrointestinal tract can be transmitted to humans and other animals [95, 113].

Table 5 shows the resistance levels of *E. coli* isolates from chickens and chicken meat [76, 114–126]. Some studies reported resistance to aminoglycosides, β -lactam groups (penicillins, cephalosporins), and fluoroquinolones in all tested isolates.

Staphylococcus aureus

Staphylococcus aureus (*S. aureus*) is an opportunistic agent that causes nosocomial diseases, which can lead from mild skin lesions to fatal endocarditis [127]. The greatest concern related to the worldwide spread of *S. aureus* [128] is the emergence of strains of methicillin-resistant *S. aureus* (MRSA), often found in humans and animals [129]. They are commonly isolated from chickens and can be transmitted to humans by direct contact or consumption of meat [130, 131]. Between 1998 and 2012, eight human outbreaks attributed to *S. aureus* of poultry origin were reported, resulting in 133 sick people and 19 hospitalizations in the USA [132].

Avian staphylococcosis can result in injuries that lead to economic losses due to decreased production and carcass condemnation. The epidemiology is complex since it is widespread in the environment and in the skin and mucosa of animals and humans [133]. MRSA strains have already been detected in rodents [134], which can introduce multi-resistant pathogens in poultry farms [133].

Table 6 summarizes the results of antimicrobial resistance in *S. aureus* obtained from studies between 2015 and 2018. The greatest resistances were found for amoxicillin (100%), amoxicillin-clavulanic acid (80–5%), ampicillin (100–5.9%), cefoxitin (100–10.5%), kanamycin (79.3–5, 2%), penicillin (100–2.1%), and tetracycline (91.2–10%). ST398 was the most detected strain among all isolates described in these studies that originated from chickens [123, 135–137] and chicken meat [138–142].

Table 5 Summary of antimicrobial resistance results in *Escherichia coli* isolated from poultry published in the last 5 years, within the five continents.

Study	Region	Reference	Publishing year	No. of isolates	Resistance profile (%)
1	Africa	Chishimba et al.	2016	77	AMP (100), C (57,1), CAZ (100), CIP (48,1), GM (37,7), NAL (48,1), NOR (54,5), S (20,8), SXT (41,6), T (59,7)
2	Africa	Awogbemi et al.	2018	10	AMC (100), AMP (90), AMX (100), C (90), ERY (80) GM (50), OB (80), P (80), S (70), T (90)
3	Africa	Osmam et al.	2018	30	AMP (90), AMX (90), C (53,3), CE (0), CEF (56,7), CIP (30), CLD (96,7), COL (0), DOX (100), ENR (60), ERY (96,7), FOS (0), GM (10), NEO (73,3), NOR (93,3), OT (100), PEF (80), RIF (86,7), S (93,3), SP (86,7), SPT (100), SXT (76,7), UB (90)
4	North America	Johnson et al.	2017	75	AMC (21), AMP (41), C (7), CEF (12), CIP (1,3), CRO (13), FOX (12), GM (55), KA (13), S (47), SUL (63), SXT (12), T (51)
5	South America	Donado-Godoy et al.	2014	164	AMP (55), CIP (23), CTX (24)
6	South America	Koga et al.	2015	39	AMC (28,2), AMP (100), C (17,9), CAZ (5,1) CIP (43,6), CTX (94,9), ENR (41), GM (43,6), KZ (100), NAL (74,4), NIT (10,3), NOR (30,8), SXT (33,3), T (97,4)
7	South America	Monte et al.	2017	8	AMX (50), CEF (87,5), COL (62,5), CRO (100), CTX (100), FOX (37,5), GM (62,5), PB (87,5), SXT (12,5), T (50)
8	South America	Ferreira et al.	2018	36	AMC (100), AZI (44,4), C (47,2), CAZ (66,7), CIP (100), CTX (100), FEP (13,9), FOX (72,2), GM (30,6), LEV (94,4), NAL (100), SXT (30,6), T (88,9)
9	Asia	Nguyen et al.	2015	895	AK (5,4), AMC (47,9), AMP (86), C (68,1), CAZ (2), CIP (32,5), CRO (2,5), GM (19,9), SXT (69,7), T (93,4)
10	Asia	Brower et al.	2017	1556	AMP (43,8), C (7,1), CFX (4), CIP (39,4), COT (42,2), GM (12,9), IMP (0), NAL (86,1), NIT (18,4), T (47)
11	Asia	Amir et al.	2017	450	AMX (55,1), AZI (39,8), C (56,2), CE (64,7), CIP (72,7), CRO (9,1), GM (59,6), NAL (67,6), OT (99,3), P (80,7)
12	Asia	Hussain et al.	2017	168	C (8), CAZ (37), CIP (70), COT (45), FOS (4), GM (32), T (84)
13	Europe	Hasman et al.	2015	5	AK (0), ATM (100), CAZ (100), CIP (20), CTX (100), COL (100), DOR (0), DOX (40), ETP (0), FEP (0), GM (0), IMP (0), LEV (20), MEM (0), MH (20), PB (100), SXT (60), T (0), TGC (0), TIM (40), TZP (0)
14	Europe	Lambrecht et al.	2017	35	AMP (100), AZI (2,9), C (42,9), CAZ (62,9), CIP (91,4), COL (2,9), CTX (65,7), GM (2,9), MEM (0), NAL (54,3), SUL (80), T (62,9), TGC (0), TMP (74,3)

AK amikacin, AMC amoxicillin/clavulanic acid, AMP ampicillin, AMX amoxicillin, AZI azithromycin, C chloramphenicol, CAZ ceftazidime, CE cefradine, CEF ceftiofur, CFX cefuroxime, CIP ciprofloxacin, CLD clindamycin, COL colistin, COT co-trimoxazole, CRO ceftriaxone, CTX cefotaxime, DOR doripenem, DOX doxycycline, ENR enrofloxacin, ERY erythromycin, ETP ertapenem, FEP cefepime, FLM flumequine, FOS fosfomicin, FOX cefoxitin, GM gentamicin, IMP imipenem, KA kanamycin, KZ cefazoline, LEV levofloxacin, MEM meropenem, MH minocycline, NAL nalidixic acid, NEO neomycin, NIT nitrofurantoin, NOR norfloxacin, OB cloxacillin, OT oxytetracycline, P penicillin, PB polymyxin B, PEF pefloxacin, RIF rifampicin, S streptomycin, SP spermidine, SPT spectinomycin, SUL sulfamethoxazole, SXT sulfamethoxazole/trimethoprim, T tetracycline, TGC tigecycline, TIM ticarcillin/clavulanic acid, TMP trimethoprim, TZP piperacillin/tazobactam

Poultry-derived products as source of antimicrobial resistance genes to the human microbiota

Bacteria can acquire resistance against antimicrobial drugs by either spontaneous mutation (in both pre-existing or newly acquired genes) or horizontal gene transfer [143]. Considering the possibility of commensal bacteria to carry and spread ARGs through horizontal gene transfer mechanisms, foodstuffs are potential sources of ARGs to human

bacteria. Horizontal gene transfer refers to the transfer of ARGs [144] from resistant to susceptible strains, allowing bacteria to successfully adapt and colonize new environments [11]. Genes coding for antimicrobial resistance can be found in various mobile DNA elements, e.g., plasmids, transposons, genomic islands, and integrons [145]. Horizontal gene transfer can occur through bacterial transformation, transduction, or conjugation. Bacterial transformation is the process by which a recipient bacterium takes up and assimilates free DNA, which carries the resistance genes

Table 6 Summary of antimicrobial resistance results in *Staphylococcus aureus* isolated from poultry published in the last 5 years, within the five continents.

Study	Region	Reference	Publishing year	No. of isolates	Resistance profile (%)
1	Africa	Chairat et al.	2015	19	CIP (5,2), CLD (0), ERY (21), FOX (10,5), KA (5,2), OXA (10,5), P (94,7), S (0), T (36,8), TOB (0)
2	Africa	Sallam et al.	2015	288	AK (53,1), AMP (5,9), AMX (8,7), C (52,1), CIP (71,2), ERY (11,1), GM (64,2), NET (70,5), OB (1), P (2,1), RIF (50), S (28,8), SXT (77,1), T (18,8), VAN (87,5)
3	Africa	Mkize et al.	2017	29	AMP (27,6), C (69), ERY (62,1), FOX (76), GM (55,2), KA (79,3), S (62,1), T (69), TMP (58,6), VAN (14)
4	Africa	Awogbemi et al.	2018	10	AMC (80), AMP (100), AMX (100), C (70), GM (30), OB (90), P (80), S (50), T (10)
5	North America	Abdalahman et al.	2015	113	AMP (92), AZI (39,8), C (2,7), CIP (23), CLD (31,9), DOX (38,1), ERY (39,8), FOX (23,9), GM (28,3), KA (32,7), OXA (30,1), P (57,5), RIF (8,8), SXT (16,8), T (47,8), VAN (23)
6	Asia	Ali et al.	2017	56	ERY (80,4), GM (28,6), NEO (25), P (94,6), T (82,1)
7	Asia	Ruban et al.	2017	MRSA 80	FOX (67), MET (45), OXA (65)
8	Asia	Kim et al.	2018	121	AMC (5), AMP (51,2), C (2,5), CAZ (2,5), CFN (1,7), CFX (1,7), CFZ (1,7), CIP (33,9), CLD (19), CTX (1,7), DOX (20,7), ERY (20,7), FEP (1,7), GM (19), KA (24,7), P (51,2), RIF (2,5), SXT (0,8), T (38,8)
9	Europe	Kraushaar et al.	2016	68	C (2,9), CIP (39,7), CLD (88,2), ERY (85,3), FD (2,9), FOX (100), GM (2,9), KA (22,1), P (100), QD (80,9), S (27,9), SUL (1,5), T (91,2), TIA (63,2), TMP (73,5)

AK amikacin, AMC amoxicillin/clavulanic acid, AMP ampicillin, AMX amoxicillin, AZI azithromycin, C chloramphenicol, CAZ ceftazidime, CFN cephalothin, CFX cefuroxime, CFZ cefazoline, CIP ciprofloxacin, CLD clindamycin, CTX cefotaxime, DOX doxycycline, ERY erythromycin, FEP cefepime, FD fusidic acid, FOX ceftoxitin, GM gentamicin, KA kanamycin, MET methicillin, NEO neomycin, NET netilmicin, OB cloxacillin, OXA oxacillin, P penicillin, QD quinupristin/dalfopristin, RIF rifampicin, S streptomycin, SUL sulfamethoxazole, SXT sulfamethoxazole/trimethoprim, T tetracycline, TIA tiamulin, TMP trimethoprim, TOB tobramycin, VAN vancomycin

from the environment [146]. Bacterial transduction involves the transfer of DNA from bacteriophages to bacteria [146]. Plasmid transfer is probably the most frequent mechanisms of horizontal transfer of resistance genes. It refers to the self-transfer of a conjugative plasmid or transposon to a recipient cell [146]. Conjugative plasmids are the most common vectors for the transmission of antimicrobial resistance genes [143].

Plasmids are autonomously replicating extra-chromosomal DNA molecules that range from <2 and >100 kb in size [146]. They can confer resistance to one or several classes of antimicrobials, including beta-lactams, aminoglycosides, tetracyclines, phenicols, sulfonamides, trimethoprim, macrolides, and quinolones [147].

The spread of antimicrobial resistance in Gram-negative bacteria has been largely attributed to DNA exchange via conjugation of plasmids carrying ARGs [148]. Multi-resistance plasmids can be transferred into a susceptible bacterial host independently of whether there is selection pressure for one or all the resistance genes [146]. Thus, the persistence of a gene conferring resistance to a given antimicrobial could occur without the selection pressure on the gene itself. Following the ban of avoparcin,

a glycopeptide antibiotic analogue of vancomycin, the prevalence of vancomycin-resistant *E. faecium* in broilers dropped from 72.7% in 1995 to 5.8% in 2000 [149]. However, the prevalence in pigs did not decrease until the use of tylosin was reduced. It was shown that tylosin provided the selective pressure for *E. faecium* to maintain the vancomycin-resistant vanA-erm(B) plasmid which also carried tylosin resistance genes [149]. This is also known as co-resistance, a mechanism of co-selection, when genes conferring a number of antimicrobial resistance phenotypes are located on the same genetic element such as a plasmid or integron [150]. Another mechanism of co-selection, known as cross-resistance, allows bacteria to survive the effects of more than one drug after acquisition of a given antimicrobial resistance gene or mutation [151].

The surveillance of commensal bacterial in foodstuff carrying antimicrobial resistance genes has been used in surveillance programs as an indicator of the level of antimicrobial resistance. Advances in high-throughput analytical methods can provide valuable information on qualitative assessment of chicken products regarding the levels of ARGs.

Alternatives to reduce the use of antimicrobials and multidrug-resistant foodborne pathogens in the poultry industry

As the major risks for human-associated diseases caused by multidrug-resistant pathogens have been linked to the consumption of contaminated food, or direct contact with animals, a thorough investigation of the source of antimicrobial-resistant foodborne pathogens upstream in the production cycles is needed to effectively tackle the problem.

For many pathogens, transmission pathways are often very complex and involve wildlife and the environment as reservoirs. In view of the amount of information available in the literature, this section will focus on the control of *Salmonella* organisms. A recent investigation of *Salmonella* recovered from captive wildlife host species and the environment in Ohio, USA, determined that indirect transmission of *Salmonella* could occur among animal hosts via environmental contamination [152]. The same study reported multidrug-penta-resistant *Salmonella* types (AmSTTeKmGm), suggesting that the role of wildlife and environment in the transmission of antimicrobial resistance to production animals and humans could be underestimated.

Interventions are typically most cost-effective and practical for the control of multidrug-resistant foodborne pathogens when applied upstream in the food supply chain at the farm level, reducing their occurrence and spread through the food chain.

Pre-harvest control of antimicrobial resistance

Strategies to prevent human MDR-foodborne pathogens, such as *Salmonella*, have broadened from pre-harvest control at the farm level to post-harvest control of food products [153]. The problem associated with the occurrence of these pathogens in the poultry industry is multifactorial and requires solutions addressing micro-ecosystems and multiple variables. This can be summarized into four key control strategies, namely, vaccination, testing, flock management and sanitation, and gastrointestinal colonization control [153].

Immune responses triggered by vaccinations can reduce the duration and severity of *Salmonella* infections in chickens, as well as prevent reinfections. However, it cannot completely prevent infections. Poor sanitation, inadequate vaccine management and environmental stress on the chickens can adversely impact vaccine performance. Vaccines also have limited efficacy against field strains or emerging *Salmonella* serotypes which are antigenically

different from vaccine strains. In response to the dramatic increase in human salmonellosis due to egg-transmitted *S. Enteritidis*, several testing and quality assurance programs for commercial laying flocks have been proposed by government agencies and the private sector. In the USA, a combination of environmental sampling, serologic testing, and bacteriological culturing of birds has been employed. The National Poultry Improvement Plan (NPIP) was designed to be a cooperative industry, state, and federal testing and monitoring program to eliminate some poultry diseases including *S. Pullorum* and *S. Enteritidis*. However, testing retains a degree of uncertainty and has limited value in predicting or responding to emerging reservoirs of *Salmonella* infection or serotypes. Effective biosecurity and hygiene practices as part of flock management and sanitation is crucial, since all other measures would fail if these were not executed appropriately. Effective biosecurity and hygiene practices could be further elaborated upon to include disinfection practices, pest controls, animal, person and vehicle movements, feed, water, and sources of poultry.

Gastrointestinal colonization control of foodborne pathogens

There are various products promoted as feed additives or treatments that not only improve animal health but can also protect birds from *Salmonella* infections. Some acidic compounds or organic acids have been shown to directly reduce colonization and shedding of *Salmonella* [154]. Prebiotics are non-digestible feed ingredients that stimulate the growth of one or a limited number of resident beneficial bacterial species in chickens [155]. Competitive exclusion products are gastrointestinal contents of adult poultry given to chicks to establish an adult microflora that is resistant to *Salmonella* colonization [156]. Probiotics are living microbial feed ingredients that have anti-inflammatory effects and the potential to modify the microbiological ecosystem within the poultry gut to control *Salmonella* [157]. Bacteriophages, live viruses that attack bacteria, have shown promise in reducing colonization in experimental trials [158], but field trial studies are limited [154]. Phytobiotics may modulate the chicken microbiome similar to antimicrobials [159], prevent the biofilm-production [160], and prevent gastrointestinal diseases in broilers [161], without causing neither collateral effects to birds [162] nor selection pressure to antimicrobial resistance in bacteria [163]. Finally, potential feed additives, such as flavophospholipol, could have the beneficial effect of reducing both *Salmonella* shedding and the number of animals carrying *Salmonella* at slaughter [164].

Surveillance programs to control antimicrobial resistance

Programs addressing the surveillance of antimicrobial resistance in the food chain have been implemented in some countries and are currently under implementation in many others. They are normally focused on the primary production level (pre-harvest) but can also include the surveillance in animal-derived products (post-harvest). Although targeting mainly foodborne zoonotic bacteria, these programs can also include classic zoonotic agents and animal pathogens as well.

In Europe, the basis for the monitoring of antimicrobial resistance in the food chain was established in 2003 by the Directive on the monitoring of zoonoses and zoonotic agents (Decision 2003/99/EC, 2003). Since then, monitoring has been implemented by means of the legislative authority of the European Commission under the scientific advice of the European Food Safety Agency (EFSA). Detailed information on the microbiological criteria applied to foodstuffs can be found in the Commission Regulation 2005/2073/EC (<https://eur-lex.europa.eu/legal-content/EN/ALL/?uri=CELEX%3A32005R2073>). Efforts towards the standardization of targeted bacteria, samplings procedures, antimicrobials to be used, and their respective cut-off values have been implemented through the Commission Decision 2007/516/EC and Decision 2013/652/EU and were guided by important reports published by EFSA [165–168] providing valuable information for the establishment of the targeted animal populations or food categories and the bacterial hazards. *Salmonella* surveillance in laying hens, broiler chickens, and turkeys are mandatory in the National Control Programs of all European member states, enabling retrospective analysis of antimicrobial resistance in these isolates. Among other foodstuff, fresh poultry has been monitored for ESBL-, AmpC-, or cabapemase-producing *E. coli* (Decision 2013/652/EU, 2013).

National monitoring systems: the US NARMS and the Brazilian PAN-BR

In USA, the National Antimicrobial Resistance Monitoring System (NARMS) is the surveillance system tracking antimicrobial resistance in foodborne pathogens from different sources, including poultry and chicken products. NARMS was established in 1996 after an expert panel convened by the US Food and Drug Administration (FDA) recommended, in 1994, establishing a national surveillance system to monitor resistance among selected enteric bacteria of animals that can cause disease in humans [169–171].

NARMS is a collaborative effort of three federal agencies, CDC, FDA, and the United States Department of Agriculture (USDA), as well as state and local health departments in all 50 states. It was designed to help assess the consequences to human health arising from the use of antimicrobial drugs in food animal production with a view towards mitigation [172]. The main goals of NARMS are to monitor trends in antimicrobial resistance among enteric bacteria from humans, retail meats, and animals, in addition to disseminate timely information on antimicrobial resistance in pathogenic and commensal organisms to stakeholders in the USA and abroad to promote interventions that reduce resistance among foodborne bacteria. Moreover, other goals include encouraging research to better understand the emergence, persistence, and spread of antimicrobial resistance and to provide data that assist FDA in making decisions related to the approval of safe and effective antimicrobial drugs for animals [173].

Since 2016, under the coordination of the Ministry of Health, Brazil has been implementing its National Action Plan (PAN-BR). The Plan is also in line with the Global Action Plan, defined by the tripartite alliance among the World Health Organization (WHO), the OIE, and the Food and Agriculture Organization (FAO), on the “One Health” concept and involves coordination with the Brazilian Ministry of Health, the Health Regulatory Agency (ANVISA), the Ministry of Agriculture, Livestock, and Food Supply (MAPA), the Ministry of Science, Technology, Innovations and Communications (MCTIC), the Ministry of Cities, the Ministry of Education, and the Ministry of the Environment [174]. In May 2018, MAPA published the National Action Plan on Antimicrobial Resistance in Agriculture (PAN-BR Agro) which has been integrated into the PAN-BR.

The PAN-BR Agro presents eight main objectives, 15 strategic interventions, and 21 activities to be implemented up to 2022 [175]. The plan targets health education, epidemiological studies, surveillance and monitoring of antimicrobial use, strengthening infection prevention and implementation of control measures, and promotion of antimicrobial rational use in animals. It can be expected that the data gaps on antimicrobial use will be solved by these activities. Moreover, the regulation of antimicrobial use, such as the need of veterinary prescriptions, the research of alternatives for antimicrobials used as growth promoters, and education programs will contribute to the prudent use of antimicrobials. Some of the activities include several workshops, the implementation of surveillance system to monitor AMR in bacteria recovered in official pathogen monitoring programs, and development of the monitoring program on antimicrobial use in animals, among others [176].

Judicious use of medically important antimicrobials in animal agriculture

The WHO has reiterated guiding principles to manage antimicrobial resistance in food animals through measures such as enhanced surveillance and better animal husbandry. Better animal husbandry practices include phasing out the use of medically important antibiotics for the treatment of humans as performance enhancers in animal production. Following joint workshops among the WHO, OIE, and FAO on non-human antimicrobial usage and antimicrobial resistance, OIE has developed standards and guidelines to address antimicrobial resistance issues including the responsible and prudent use of antimicrobials and the development of surveillance and monitoring programs for antimicrobial resistance. In 2005, the Codex Alimentarius Commission issued a code of practice to countries in order to mitigate antimicrobial resistance. In response to the abuse and misuse of antimicrobials in the USA, the Food and Drug Administration (FDA) developed strategies outlined under the guidance document #209 on “The Judicious Use of Medically Important Antimicrobial Drugs in Food-Producing Animals.” Medically important antimicrobial drugs refer to antimicrobial drugs that are important for human therapeutic use. The two guiding principles are that “the use of medically important antimicrobials in food-producing animals should be limited to those uses that are considered necessary for assuring animal health” and “the use of medically important antimicrobial drugs in food-producing animals should be limited to those uses that include veterinary oversight or consultation.”

While legislations and initiatives to limit antimicrobials except for therapy in food-producing animals could be put in place, it seems likely that such changes alone would not solve the problem of antimicrobial resistance. The American Veterinary Medical Association (AVMA) highlighted that the contribution of antimicrobial use in animal production to the global dissemination of antimicrobial resistance is probably limited [177]. Denmark banned the use of several antimicrobial growth promoters in food animals from 1994 to 1999, and a voluntary ban on the use of cephalosporins in pig production was implemented in 2010. According to the Danish Integrated Antimicrobial Resistance Monitoring and Research Programme (DANMAP) report for 2013, the prevalence of antimicrobial-resistant bacteria remained constant or on the rise [178]. The report stated that *S. Typhimurium* isolates among Danish pigs exhibited high levels (range 33–47%) of resistance to ampicillin, streptomycin, sulfonamide, and tetracycline. This co-resistance to four antimicrobial classes refers to the ASSuT profile and could be seen alone or together with resistance to other antimicrobial classes. Resistance to tetracycline increased from 2012 to 2013, while resistances to other antimicrobial agents

were similar to levels reported in 2012 [178, 179]. The level of multidrug-resistance among *E. coli* isolates in Danish broiler chickens and broiler meat has more than doubled within the past 5 years, reaching 16–18% in 2013 [178]. It was presumed to be driven by an increased therapeutic use of tetracycline in poultry production as subtherapeutic use was banned. There were no significant changes in levels of resistance from *Salmonella* infections in humans between 2009 and 2013. In the 10-year period from 2004 to 2013, resistance in *E. coli* human blood isolates to 2nd generation cephalosporins, ciprofloxacin, and aminoglycoside increased steadily [178]. While it is crucial to ban important antimicrobials in animal production, the Danish experience showed that further approaches are needed not only to reduce food-borne pathogens in animal-derived foods but also the transmission of antimicrobial resistance.

There will be or already are stricter rules or bans on antibiotic growth promoters globally, although it is recognized that there cannot be a total ban on antibiotic use in food animals, at the expense of dramatic adverse health consequences on animals and limitation of tools at the disposal of veterinarians [180].

A solution suggested in stewardship bootcamps for and against the use of antimicrobials in food animals [181] is the use of alternative feed additives that could reduce food-borne pathogen carriage and shedding, promote growth, and reduce the incidence of antimicrobial-resistant bacteria in the gut of food animals [182].

The One Health approach to mitigate antimicrobial resistance in the globalized world

Although there is a consensus that actions must be taken to reduce the emergence and spread of antimicrobial resistance in livestock, those actions must consider key premises:

- 1) Antimicrobial resistance is an inevitable biological phenomenon in bacteria that is accelerated by antimicrobial use selective pressure. The actions should address the critical aspects of antimicrobial resistance to human, animals, and the associated ecosystem. The alternatives to control antimicrobial resistance need to be supported by scientific-based knowledge.
- 2) The animal industry plays a key role in the global food supply. Fighting antimicrobial resistance need to consider economically feasible alternatives that warrant the success of the industries and the sustainable development of the regions where they are located.
- 3) All sectors of the society, including the decision-makers in the companies, must understand that antimicrobial resistance is a critical topic for the present and future

of the human civilization. If this is somehow difficult to be grabbed, the companies' lack of commitment on the reduction of antimicrobial-resistant could jeopardize the success of their business.

- 4) Governments must understand that fighting antimicrobial resistance requires coordinated efforts with the private sector to promote disruptive shifts. That requires an adequate environment for research, development actions, and cost compensation.
- 5) There is no villain nor magic bullet for the control of antimicrobial resistance. Its mitigation depends on the coordinate multidisciplinary efforts and strengthens capacity building and education. The One Health approach must be embraced as a disruptive platform to tackle antimicrobial resistance and other complex problems.

As explained in the previous sections of this manuscript, antimicrobial resistance is very complex multi-branched problem that require actions at the human, animal, and environmental levels.

Awareness (or lack thereof) is growing on the overlap of individual, regional, and global public health, as well as the health of humans, animals, and the environment [183]. The "One Health" concept involves an interdisciplinary, cooperative approach and is captured in this statement: "collaborative efforts of multiple disciplines, working locally, nationally and globally, to reach optimal health for people, animals and the environment." [184] The concept of "One Health" or "One Medicine" is not new. It was first mentioned by Dr. Calvin Schwabe who recognized that collaboration between human and veterinary professionals is needed to control zoonotic diseases [185].

Following the 2015 World Health Assembly, a global action plan on AMR was adopted [186]. The plan underscores the strong collaboration between the WHO, FAO, and OIE to work with United Nations to combat AMR at the political level. The objective was to have multi-sectorial national action plans by 2017. AMR is also only one of three priority areas that the tripartite organizations have agreed to work on, in addition to rabies and avian influenza. The WHO recommended an overall reduction in the use of medically important antimicrobials in food-producing animals, as well as a complete restriction in the use of such antimicrobials for growth promotion and disease prevention [187]. The FAO action plan of 2016–2020 focuses on improving awareness, developing surveillance and monitoring capacity, strengthening governance, and promotion of good practices [188]. The plan has since been updated to 2021–2025 to cover research and allocation of resources sustainably. The OIE also announced its strategy on AMR in 2016, covering similar areas, with an addition of encouraging the implementation of international standards. The One Health

Global Leaders Group on antimicrobial resistance (Global Leaders Group) has recently been established, which was a recommendation by the Interagency Coordination Group (AMR) on Antimicrobial Resistance. The United Nations Environment Programme (UNEP) is also slated to join the tripartite to combat AMR. This is a positive move to ensure that the One Health approach and political commitment are maintained.

Interestingly, the "One Health concept" has begun to be incorporated in the animal production systems. For instance, a certification program for animal-derived foodstuff produced according to the One Health principles (One Health Certification) has been recently launched. Additionally, other initiatives such as the One Health Poultry Hub, led by the U.K. Royal Veterinary College (RVC), will concentrate efforts in Bangladesh, Sri Lanka, and Vietnam and will look at why intensification of poultry production increases the risk of infectious disease and identify high-risk behaviors, processes, and environments. It will also test and evaluate novel interventions for disease control. With a GBP19.1 million (US\$2.5 million) funding, the major goals of this hub are to assist in the development of policies for the environmental, economic, and social sustainability of poultry production systems and reduce threats to human and animal health and welfare. Bacterial food poisoning, avian influenza, and antimicrobial resistance are among the key areas for collaborators in the hub.

While such initiatives are welcome, poultry companies need to not only apply the principles of One Health, but they need to communicate this compliance with consumers, in order to build consumer trust. Given the inter-sectorial nature of AMR, the poultry industry also needs to join forces with representatives of other sectors, including wildlife and environmental agencies, as these serve as sentinel for AMR [189].

The antimicrobial resistance is probably the clearest example of an increasing multifaceted problem requiring solutions in One Health. The vision that embraced the antimicrobial resistance with a more heuristic vision, including actions at the different levels such as humans, animals, and the environment, is important, but will not solve or enable the civilization to efficiently tackle the problem. One Health initiatives on antimicrobial resistance must include actions addressing the key factors related to the emergence of AMR: political willingness and education on how we interact with the surrounding environment. While we recognize that much must be done in order to better understand the complexity of AMR, recent epidemiological studies agree that the solution is aligned with the principles of global health promotion, reduction of inequity, and investments on research and development to improve our capacity to understand the problems and propose effective control measures. Importantly, investments on research and development must

require interdisciplinary approaches involving social scientists, economists, and natural science components. For example, many aspects of the misuse of antimicrobials in human and veterinary medicine, considered key drivers for the emergence of antimicrobial resistance, are related to social behavior.

Lastly, political willingness is of crucial importance, as only coordinate actions among the many organizational levels could resonate.

Final considerations

In poultry farming, a wide range of bacterial species can infect broiler and layer chickens. As in other sectors, the emergence and dissemination of antimicrobial resistance among those bacteria have been posing a serious threat to the whole society, including the industry itself, because:

- 1) Avian diseases caused by drug-resistant bacteria are more difficult to be treated, leading to aggravated economic losses. Furthermore, animal-derived products have the potential to carry commensal bacteria harboring important antimicrobial resistance genes. The levels of resistance genes or resistant commensal bacteria in foodstuff can be used as non-tariff barriers for exporting countries.
- 2) Some of these bacteria are zoonotic pathogens, increasing the risk for consumers or workers to also acquire hard-to-treat infections, representing a burden to public health.

Antimicrobial resistance represents a long-term and very complex biological problem; therefore, coordinate and accurate actions are necessary in order to reduce its impact on the present and the future of the human civilization. Such measures need to be performed upon science-based knowledge in order to preserve the economic efficiency of the industry along with the promotion of public health. In this sense, Environmental, Social, and Governance (ESG) integrated initiatives in the poultry industry are encouraged to embrace One Health principals.

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Declarations

Conflict of interest The authors declare no competing interests.

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