J Anim Sci Technol 2024;66(2):266-278 https://doi.org/10.5187/jast.2023.e129



Received: Oct 4, 2023 Revised: Nov 6, 2023 Accepted: Nov 14, 2023

[#]These authors contributed equally to this work.

*Corresponding author

Ju-Hoon Lee Department of Agricultural Biotechnology, Seoul National University, Seoul 08826, Korea. Tel: +82-2-880-4854 E-mail: juhlee@snu.ac.kr

Hyeun Bum Kim Department of Animal Biotechnology, Dankook University, Cheonan 31116, Korea. Tel: +82-41-550-3653 E-mail: hbkim@dankook.ac.kr

Copyright © 2024 Korean Society of Animal Sciences and Technology. This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (http:// creativecommons.org/licenses/bync/4.0/) which permits unrestricted non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited.

ORCID

Sriniwas Pandey https://orcid.org/0000-0002-6947-3469 Hyunok Doo https://orcid.org/0000-0003-4329-4128

Antibiotic resistance in livestock, environment and humans: One Health perspective

Journal of Animal Science and Technology

pISSN 2672-0191 eISSN 2055-0391

Sriniwas Pandey^{1#}, Hyunok Doo^{1#}, Gi Beom Keum^{1#}, Eun Sol Kim¹, Jinok Kwak¹, Sumin Ryu¹, Yejin Choi¹, Juyoun Kang¹, Sheena Kim¹, Na Rae Lee¹, Kwang Kyo Oh², Ju-Hoon Lee^{3,4,5}* and Hyeun Bum Kim¹*

¹Department of Animal Biotechnology, Dankook University, Cheonan 31116, Korea ²Microbial Safety Division, National Institute of Agricultural Sciences, Rural Development Administration, wanju 55365, Korea

³Department of Food Animal Biotechnology, Seoul National University, Seoul 08826, Korea ⁴Department of Agricultural Biotechnology, Seoul National University, Seoul 08826, Korea ⁵Center for Food and Bioconvergence, Seoul National University, Seoul 08826, Korea

Abstract

Antibiotic resistance (AR) is a complex, multifaceted global health issue that poses a serious threat to livestock, humans, and the surrounding environment. It entails several elements and numerous potential transmission routes and vehicles that contribute to its development and spread, making it a challenging issue to address. AR is regarded as an One Health issue, as it has been found that livestock, human, and environmental components, all three domains are interconnected, opening up channels for transmission of antibiotic resistant bacteria (ARB). AR has turned out to be a critical problem mainly because of the overuse and misuse of antibiotics, with the anticipation of 10 million annual AR-associated deaths by 2050. The fact that infectious diseases induced by ARB are no longer treatable with antibiotics foreshadows an uncertain future in the context of health care. Hence, the One Health approach should be emphasized to reduce the impact of AR on livestock, humans, and the environment, ensuring the longevity of the efficacy of both current and prospective antibiotics.

Keywords: Antibiotic resistance, Livestock, Human, Environment, One Health

INTRODUCTION

Antibiotic resistance (AR) is a complex, multifaceted issue that poses a serious threat to the health of livestock and humans, as well as the surrounding environments, including soil, air, and water. Its intricacy results from various interrelated elements that contribute to its development and spread, making it a challenging issue to address. AR is on the rise globally, and if current trends continue, there could be up to 10 million annual AR-associated deaths from a variety of untreatable infections by 2050 [1,2]. It is one of the many concerns of the 21st century identified by the Lancet Commission that threatens to hinder future progress in global health [3].

AR is one of the significant global health issues that best exemplifies the One Health concept [4]. This is because bacterial species such as *Escherichia coli* (*E. coli*), *Salmonella* and *Campylobacter* carry

Gi Beom Keum

https://orcid.org/0000-0001-6006-9577 Eun Sol Kim https://orcid.org/0000-0001-8801-421X Jinok Kwak https://orcid.org/0000-0003-1217-3569 Sumin Rvu https://orcid.org/0000-0002-1569-3394 Yejin Choi https://orcid.org/0000-0002-7434-299X Juyoun Kang https://orcid.org/0000-0002-3974-2832 Sheena Kim https://orcid.org/0000-0002-5410-1347

Na Rae Lee https://orcid.org/0009-0003-7230-9891 Kwang Kyo Oh https://orcid.org/0000-0001-5300-2091 Ju-Hoon Lee https://orcid.org/0000-0003-0405-7621 Hveun Bum Kim https://orcid.org/0000-0003-1366-6090

Competing interests

No potential conflict of interest relevant to this article was reported.

Funding sources

This work was carried out with the support of "Cooperative Research Program for Agriculture Science & Technology Development (Project No. PJ0162982023)" Rural Development Administration, Korea.

Acknowledgements

Not applicable.

Availability of data and material

Upon reasonable request, the datasets of this study can be available from the corresponding author.

Authors' contributions

Conceptualization: Pandey S, Doo H, Keum GB, Oh KK, Lee JH, Kim HB.

- Data curation: Doo H.
- Formal analysis: Keum GB.

Validation: Kim ES, Kwak J, Kim S. Investigation: Ryu S, Choi Y, Kang J, Lee

- NR. Writing - original draft: Pandey S, Doo H,
- Keum GB, Kim HB. Writing - review & editing: Pandey S, Doo H, Keum GB, Kim ES, Kwak J, Ryu S, Choi Y, Kang J, Kim S, Lee NR, Oh KK, Lee JH, Kim HB.

Ethics approval and consent to participate

This article does not require IRB/IACUC approval because there are no human and animal participants.

antibiotics resistant genes and can spread between livestock, humans, and the environment [5–7]. The One Health approach recognize the interconnection and mutual influence of these three domains. One Health is defined as "a collaborative effort of multiple-discipline health science professions, together with their allied disciplines and institutions-working locally, nationally, and worldwide to achieve optimal health for domestic animals, humans, wildlife, plants, and the environment as a whole" [7]. AR is closely linked to each of these factors due to the reckless and excessive use of available antibiotics in agriculture, livestock farming and human medicine [8,9]. The interconnections between livestock, humans, and environmental compartments create pathways for the transmission of bacteria, mobile genetic elements (MGEs), and antibiotics themselves [10]. These pathways facilitate the movement of these entities between these compartments, thereby promoting the spread of AR.

The local and global spread of AR can be attributed to various factors, including inadequate antibiotic use in both livestock and humans, contaminated environments, and ineffective infection control policies [11–13]. Careless human use of antibiotics contributes to the development and spread of antibiotic resistant bacteria (ARB) and antibiotic resistance genes (ARGs). These resistant bacteria can be transmitted to other humans through the direct contact, as well as through food, water, and environmental surfaces [14]. Similarly, the use of antibiotics in livestock can also contribute to the emergence of antibiotic-resistant bacteria. These bacteria can be transmitted to humans through direct contact with livestock, via food products, and can also be present in the environment due to livestock waste [15]. However, it is not yet clear to what extent livestockderived transmission routes contribute to the spread of AR compared to human-derived routes [10]. Environmental contamination is also a major factor in the dissemination of antibioticresistant bacteria and genes. Antibiotic-resistant bacteria can persist in soil and water, potentially spreading to animals and humans when they come into contact with contaminated environmental sources [16]. While selective pressure and horizontal gene transfer are known factors in the spread and dissemination of antibiotic-resistant bacteria and genes, the absence of proper policies and regulations for the use and disposal of antibiotics only exacerbates this issue.

This review aims to enhance understanding of the interdependence between livestock, human, and environmental health in terms of the emergence, spread, and concerns related to AR in each of these domains.

ANTIBIOTIC RESISTANCE IN LIVESTOCK

ARB and ARGs in food animals are regarded as emerging contaminants that pose a serious threat to global public health [17]. Multiple reports have indicated a 35% increase in global antibiotic consumption during the first decade of the 21st century [18]. A study conducted by Muchandani et al. using statistical models estimated antimicrobial usage at 99,502 tons in 2020 and projected an 8% increase to 107,472 tons based on the latest trend [19].

Livestock farms indeed represent a significant source of ARB and ARGs. These ARBs often display resistance to several major classes of antibiotics, such as aminoglycosides, tetracyclines, sulfonamides, chloramphenicols, and β-lactams [20,21]. Livestock farms can serve as reservoirs for pathogenic zoonotic bacteria such as Staphylococcus aureus (S. aureus), Salmonella species, Campylobacter species, E. coli, and Listeria species. These pathogens can be transmitted into environment through air, slurry, and feces from livestock, leading to the potential outbreak of infections and acting as a source of ARGs [22,23].

The causes of AR are indeed very complex, involving various factors such as the overuse and misuse of antibiotics in both human medicine and agriculture, as well as environmental

contamination [24]. The administration of antibiotics in livestock is primarily motivated by three main factors: to treat infections, to prevent animals at risk from getting infection or as feed additives to boost growth [25]. Numerous studies have demonstrated a link between the use of antibiotics in food-producing animals, especially for growth promotion, and AR in people who either live on or near farms [15]. Additionally, drug-resistant strains can emerge if the prescribed antibiotics are not changed in a timely manner [26]. These bacteria generally develop resistance to antibiotics through several different resistance mechanisms, including blocking the antibiotic's target sites, altering the permeability of cell membranes, and producing antibiotic-inactivating enzymes [27]. The resistance observed in food-related bacteria reflects the broader condition of bacterial resistance within the environment from which food for human consumption originates [28]. Despite an increase in the number of relevant studies on livestock and AR, as of 2020, they still accounts for only 17% of all the studies published on AR [29]. Therefore, there is a pressing need to promote more research in this area to better understand and address the complex issues surrounding AR in livestock and its implications for human health.

ANTIBIOTIC RESISTANCE IN HUMANS

The human community is facing an escalating crisis of antibiotic-resistant infections in patients, presenting a persistent and increasing threat to public health [30].

The World Health Organization (WHO) released a list of 12 families of bacteria in 2017 that pose a significant threat to human health [31], categorizing them into three priority groups based on the urgency of developing novel antibiotics to combat them [32]. The most critical group comprises multidrug-resistant bacteria such as *Acinetobacter, Pseudomonas*, and certain *Enterobacteriaceae* like *Klebsiella pneumoniae* (*K. pneumoniae*), *E. coli*, and *Enterobacter* species. These bacteria can cause serious and potentially fatal infections, especially in patients admitted to hospitals and nursing homes or those requiring ventilators and catheters, leading to conditions such as pneumonia and bloodstream infections [33–35]. The high-priority category, which includes bacteria such as *Enterococcus faecium* (*E. faecium*) and *S. aureus*, exhibits resistance to a number of antibiotics such as vancomycin and fluoroquinolones [33]. The medium-priority category includes *Streptococcus pneumoniae* and *Shigella*, which may have some resistance but can still be effectively treated with the antibiotics currently available [36]. Table 1 lists the priority pathogens identified by the WHO for the development of new antibiotics to combat antibiotic-resistant bacteria (Table 1).

AR is indeed a natural process, but its uncontrolled spread has escalated into a public health emergency largely due to the overuse and misuse of antibiotics. This excessive usage accelerates the development and dissemination of antibiotic-resistant bacteria [37]. Among the most influential human behaviors contributing to the development and spread of AR are the failure of patients to adhere to prescribed antibiotic regimens and their tendency to engage in self-medication. These behaviors have a significant impact on the proliferation of antibiotic-resistant strains [38]. The public's perception that antibiotics offer a quick and effective solution to common health problems has led to actions that undermine a doctor's prescribing control [39]. The use of broad-spectrum antibiotics as a substitute for accurate diagnostics or to increase the chances of therapeutic success also contributes to the selection of resistant bacteria [40]. According to a report from the Centers for Disease Control and Prevention (CDC), more than 50% of antibiotic prescriptions are considered unnecessary in the United States [41]. While the increase in the prevalence of AR cannot be solely attributed to antibiotic overuse, other factors known as "socioeconomic determinants" also play a significant role. These determinants include poor community hygiene, unsafe food practices, inadequate infection control measures in healthcare settings, the presence of

Priority level	Pathogens	Antibiotic Resistance
Critical	Acinetobacter baumannii	Carbapenem resistant
	Pseudomonas aeruginosa	Carbapenem resistant
	Enterobacteriaceae (including K. pneumoniae, E. coli, Enterobacter spp.)	Carbapenem resistant
		3rd generation cephalosporin resistant
High	Enterococcus faecium	Vancomycin resistant
Medium	Staphylococcus aureus	Methicillin resistant
		Vancomycin intermediate and resistant
	Helicobacter pylori	Clarithromycin resistant
	Campylobacter	Fluoroquinolone resistant
	Salmonella spp.	Fluoroquinolone resistant
	Neisseria gonorrhoeae	3rd generation cephalosporin resistant
		Fluoroquinolone resistant
	Streptococcus pneumoniae	Penicillin nonsusceptible
	Haemophilus influenzae	Ampicillin resistant
	Shigella spp.	Fluoroquinolone resistant

Table 1. World Health Organization (WHO) priority pathogens list (2017): key antibiotic-resistant Bacteria [94]

antibiotics in the environment, and their use in animals and agriculture [42]. AR in humans is also influenced by entrenched medical practices, economic pressures, and vaccine hesitancy [43].

ANTIBIOTIC RESISTANCE IN ENVIRONMENT

In recent years, AR in the environment has emerged as a significant public health concern [44]. Several studies have reported the presence of ARB and antibiotic residues in various environmental samples, including soil, water, and wildlife. These findings underscore the need for effective strategies to mitigate the further spread of AR in the environment [44,45]. AR can arise through mutations in a bacterium's pre-existing genome or through the uptake of foreign DNA, both of which contribute to the development of AR. Mutations can easily occur in bacteria in a livestock or patient receiving antibiotic treatment and can become permanently established. However, such intense selection pressure on pathogens is less common in other environmental settings. Therefore, external environmental factors are less likely to influence mutation-based evolution of resistance in most pathogens. In contrast, the uptake of novel resistance factors is facilitated by the diverse ecological niches of water, soil, and other environments, creating a broad gene pool that surpasses the microbiota of livestock and humans [46,47].

The detection of ARGs in diverse environmental settings, including sludge, wastewater, river water and sediment, air, and soil has been well documented [48]. Soil, in particular, serves as a natural habitat for a diverse range of microorganisms and is considered the primary and natural reservoir of ARGs. This presence of ARGs in bacteria without prior exposure to antibiotics is commonly referred to as intrinsic resistance. It highlights the fact that these resistance genes can be naturally occurring and not solely the result of antibiotic use or pollution [49]. However, due to human activities, soil has transformed from being primarily an intrinsic reservoir of ARGs to also serving as a repository and accumulation site for numerous acquired ARGs [41]. A comparative study of soil microbiome in natural and agroecosystems revealed that natural ecosystems had a relatively lower presence of ARB. The study also revealed that *Pantoea agglomerans* and *Bacillus cereus* showed the highest levels of resistance. Approximately 23% of the isolated bacteria exhibited

resistance to antibiotics [50]. Among all classes of ARGs found in wastewater samples worldwide, tet genes exhibited the highest detection frequency, with researchers identifying approximately 20 different types of *tet* genes in wastewater [51]. Untreated sewage has also been found to have a variety of ARGs encoding aminoglycosides, β -lactams, tetracyclines, and vancomycin [52–55]. Water has also been affected by ARG contamination. Several types of ARGs have been identified in microbial populations of natural water sources [56,57]. Das et al. discovered the presence of 139 ARGs belonging to 11 different ARG types in the Yamuna riverine ecosystem [58]. A study conducted by Pruden et al. clearly demonstrated the presence of ARGs in numerous environmental compartments in Northern Colorado, including river sediments, irrigation ditch water, dairy lagoon water and even in drinking water treatment plants [44]. Other studies have also identified many different ARBs in environmental samples, including E. coli, K. pneumoniae, and Pseudomonas aeruginosa (P. aeruginosa) [51,59]. Beyond these commonly found bacteria, a study in China isolated New Delhi metallo-\u03b3-lactamase 1 (NDM-1) producing Acinetobacter baumannii (A. baumannii) in hospital sewage [60]. Even the air has not escaped AR contamination, as a global survey of ARGs conducted by Li et al. revealed that urban air also contains ARGs, raising various health concerns in cities [61].

Antibiotics make their way into the environment through various pathways, including municipal sewage, hospital waste, animal husbandry refuse, antibiotic production facilities, agricultural runoff, and other sources. A significant portion of antibiotics is excreted in urine and feces within 8–24 hours after consumption, with excretion rates ranging from 30% to 90% [62]. Antibiotics have a variable half-life, ranging from hours to hundreds of days, and even after degradation, antibiotic residues persist in the environment, contributing to the selection for AR [63,64]. Antibiotics are excreted from the human body in the form of antibiotic metabolites, glucuronic acid, and sulfuric acid conjugates. Studies have shown that approximately 70%–80% of antibiotics in sewage remain in a consistent form [45,65,66]. In addition to household and hospital sewage, pharmaceutical production facilities also contribute to the antibiotic burden in sewage treatment plants, as the effluent from the pharmaceutical industry is often inadequately treated before discharge [67,68]. Animal husbandry waste and agricultural runoff are additional sources of antibiotics in the environment [69]. The extensive use of antibiotics in agriculture results in their detection in agricultural runoff, which eventually mixes with water bodies, further contributing to environmental contamination.

DISSEMINATION OF ANTIBIOTIC RESISTANCE AT LIVE-STOCK-HUMAN-ENVIRONMENT INTERFACE

AR is a global issue at the livestock-human-environment interface, providing suitable conditions for the rapid spread and evolution of bacteria. The excessive and irresponsible use of antibiotics in medicine, agriculture, livestock farming, and aquaculture has resulted in a rise in resistant bacteria in these areas.

AR evolves and spreads in a complex parameter space and therefore there is still a lack of concrete explanation regarding the spread of AR and the directions of its spread [5,70]. However, AR bacterial clones, MGEs, and ARGs have been found to be responsible for the dissemination of AR at the human-animal-environment interface [11]. MGEs, classified into 2 groups: the intercellular MGE (e.g., plasmids) and the intracellular MGE (e.g., transposons, integrons) act as vehicles for dissemination of resistance at the human-animal-environment interface [71]. Together, these components have a significant role in promoting horizontal genetic exchange, which helps disseminate resistance genes [72]. It has been found that MGEs may have even a greater role in

the transmission of AR genes from livestock to humans than bacterial cells [73,74]. *K. pneumonia* is regarded as a host of a variety of mobile ARGs and is crucial in the global spread of various extended-spectrum β -lactamases and carbapenemase [75]. MGE is also responsible for resistance in *E. faecium, S. aureus, A. baumanni, P. aeruginosa,* and *Enterobacter* species [76]. The epidemiology of AR in context of human and animal interactions is also inherently complex, involving numerous potential transmission routes and vehicles. Among these, the foodborne route of transmission is of serious concern, involving pathogens such as *Salmonella enterica, Campylobacter coli/jejuni, Yersinia enterocolitica.* Bacteria released into the environment from the feces of livestock and humans can disrupt the normal environmental flora, potentially acting as a reservoir before reintroduction into the livestock-human cycle. Therefore, all three domains—livestock, humans, and the environment—are deeply interconnected, and changes in one domain can have far-reaching effects on the others [77].

CONCERNS FOR LIVESTOCK, HUMAN AND ENVIRON-MENTAL HEALTH ASSOCIATED WITH ANTIBIOTIC RESIS-TANCE

AR has emerged as one of the most pressing threats to livestock, public health, and the environment, posing serious obstacles to the successful prevention and treatment of infectious diseases. The fact that infectious diseases induced by ARB are no longer treatable with antibiotics foreshadows an uncertain future in the context of health care [78]. The reasons behind AR are highly complex and involve human behavior at multiple levels of society, with outcomes that affect everyone worldwide, including livestock, the environment, and humans themselves (Fig. 1) [8].

AR has become an ominous specter for humanity. A study in 2019 estimated 1.27 million deaths resulted from resistance in 88 pathogen-drug combinations that were evaluated [79]. Methicillin-Resistant *S. aureus* (MRSA), a well-known case of AR, has been associated with a high death rate worldwide every year [80]. Additionally, the treatment of various infections, including urinary tract infections and pneumoniae, caused by multi-drug resistant gram-negative bacteria (MDR-GNB) has become more challenging [81,82]. Furthermore, drug resistance to gonorrhea, tuberculosis, and typhoid fever is also on the rise annually, contributing significantly to high healthcare expenses, particularly in developing countries [83].

The worldwide increase in the use of antibiotics as an indispensable part of poultry and livestock production industries, for treating and preventing infectious diseases, as well as for promoting growth, has led to the emergence of AR in bacteria in recent years [84]. This trend significantly impacts the health of both livestock and humans. The extensive use of antibiotics in livestock farming generates antibiotic-resistant bacteria and genes that can potentially transmit to humans through the food chain, posing a threat to the treatment of human infections, particularly fluoroquinolones, which are of major importance in human medicine [85,86]. However, the use of antibiotics from chemical classes different from those used by humans may still be permissible and not contribute to resistance.

The environment serves as a hidden source of antibiotic-resistant organisms carrying resistance genes. This is due to the deposition of antibiotic residues from the pharmaceutical industry, livestock husbandry, hospitals, fecal waste from humans and animals, agricultural runoff, and more. As a result, the environment is responsible for transmitting multi-drug resistant pathogens and ARGs of public health concern [87–90]. Although bacteria exhibited resistance to many drugs before the advent of antibiotics, human activities have exacerbated its spread, leading to the creation of a 'global resistome' [91]. Given the large-scale production of antibiotics each year, environmental niches play

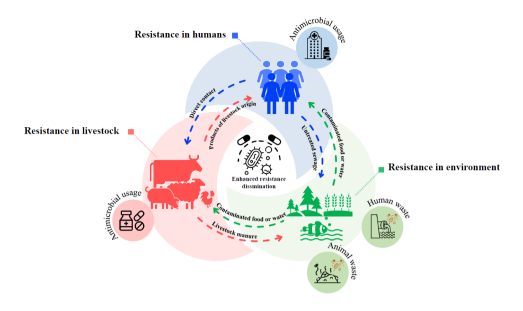


Fig. 1. A One Health perspective on antimicrobial resistance flow: a diagrammatic representation. This figure illustrates the interplay between livestock, humans, and the environment as contributors to antimicrobial resistance. Red arrows represent pathways from livestock to humans and the environment, blue arrows depict pathways from humans to livestock and the environment, and green arrows indicate pathways from the environment to humans and livestock. Each outer diagram highlights factors contributing to antimicrobial resistance.

a significant role in the persistence and spread of AR.

ONE HEALTH STRATEGIES FOR ADDRESSING THE ISSUE OF ANTIBIOTIC RESISTANCE

The One Health paradigm has now become an integral part of global initiatives aimed at confronting AR. Consequently, the major challenges in tackling AR revolve around the differing interests of various social sectors and groups involved in animal, human, and environmental health. These stakeholders have come to a consensus on critical areas of action, the most effective approaches for monitoring AR, containment of infections, and the policies governing the use of antimicrobials [4].

In a collaborative effort to combat AR, the World Organization for Animal Health (OIE), the Food and Agriculture Organization of the United Nations (FAO), and the WHO have introduced the Global Action Plan on AR, facilitating the development of AR-related policies within the One Health framework [92]. These organizations have also established guidelines for AR surveillance to ensure comprehensive monitoring of antimicrobial usage and consumption across animal, human, and environmental settings [93]. Undoubtedly, enhancing global surveillance of drug resistance is essential for effectively combating AR. The medical and scientific community requires a clear understanding of existing and historical AR data to uncover new mechanisms of resistance acquisition, conclusively identify ongoing cases, and anticipate future threats. This task necessitates a deeper understanding of three key aspects: antibiotic consumption in both human and animal contexts, current levels of AR, and the molecular foundations of AR. Additionally, to effectively combat AR through the One Health approach, international collaborations among nations are imperative, as global actions are essential [4].

CONCLUSION

AR is a serious global health concern. To reduce its impact on livestock, human, and environmental health and ensure the longevity of the efficacy of both current and prospective antibiotics, the One Health approach should be emphasized. Recognizing the interrelated nature of these three domains, bringing the human, animal, and environmental sectors together is essential for effectively combating AR and securing the health and development gains achieved over the years.

REFERENCES

- Klein EY, Van Boeckel TP, Martinez EM, Pant S, Gandra S, Levin SA, et al. Global increase and geographic convergence in antibiotic consumption between 2000 and 2015. Proc Natl Acad Sci USA. 2018;115:E3463-70. https://doi.org/10.1073/pnas.1717295115
- White A, Hughes JM. Critical importance of a one health approach to antimicrobial resistance. EcoHealth. 2019;16:404-9. https://doi.org/10.1007/s10393-019-01415-5
- Kanem N, Murray CJL, Horton R. The lancet commission on 21st-century global health threats. Lancet. 2023;401:10-1. https://doi.org/10.1016/S0140-6736(22)02576-4
- Velazquez-Meza ME, Galarde-López M, Carrillo-Quiróz B, Alpuche-Aranda CM. Antimicrobial resistance: one Health approach. Vet World. 2022;15:743-9. https://doi. org/10.14202/vetworld.2022.743-749
- Woolhouse M, Ward M, van Bunnik B, Farrar J. Antimicrobial resistance in humans, livestock and the wider environment. Philos Trans R Soc Lond B Biol Sci. 2015;370:20140083. https:// doi.org/10.1098/rstb.2014.0083
- Robinson TP, Bu DP, Carrique-Mas J, Fèvre EM, Gilbert M, Grace D, et al. Antibiotic resistance is the quintessential One Health issue. Trans R Soc Trop Med Hyg. 2016;110:377-80. https://doi.org/10.1093/trstmh/trw048
- Collignon PJ, McEwen SA. One Health—its importance in helping to better control antimicrobial resistance. Trop Med Infect Dis. 2019;4:22. https://doi.org/10.3390/ tropicalmed4010022
- Laxminarayan R, Duse A, Wattal C, Zaidi AKM, Wertheim HFL, Sumpradit N, et al. Antibiotic resistance-the need for global solutions. Lancet Infect Dis. 2013;13:1057-98. https://doi.org/10.1016/S1473-3099(13)70318-9
- Review on Antimicrobial Resistance. Tackling drug-resistant infections globally: final report and recommendations. London: wellcome trust and HM Government; 2016.
- 10. Woolhouse MEJ, Ward MJ. Sources of antimicrobial resistance. Science. 2013;341:1460-1. https://doi.org/10.1126/science.1243444
- Aslam B, Khurshid M, Arshad MI, Muzammil S, Rasool M, Yasmeen N, et al. Antibiotic resistance: One Health One World outlook. Front Cell Infect Microbiol. 2021;11:771510. https://doi.org/10.3389/fcimb.2021.771510
- 12. Marti E, Variatza E, Balcazar JL. The role of aquatic ecosystems as reservoirs of antibiotic resistance. Trends Microbiol. 2014;22:36-41. https://doi.org/10.1016/j.tim.2013.11.001
- Burow E, Käsbohrer A. Risk factors for antimicrobial resistance in Escherichia coli in pigs receiving oral antimicrobial treatment: a systematic review. Microb Drug Resist. 2017;23:194-205. https://doi.org/10.1089/mdr.2015.0318
- Liu YY, Wang Y, Walsh TR, Yi LX, Zhang R, Spencer J, et al. Emergence of plasmidmediated colistin resistance mechanism MCR-1 in animals and human beings in China: a microbiological and molecular biological study. Lancet Infect Dis. 2016;16:161-8. https://doi.

org/10.1016/S1473-3099(15)00424-7

- Marshall BM, Levy SB. Food animals and antimicrobials: impacts on human health. Clin Microbiol Rev. 2011;24:718-33. https://doi.org/10.1128/CMR.00002-11
- Czekalski N, Berthold T, Caucci S, Egli A, Bürgmann H. Increased levels of multiresistant bacteria and resistance genes after wastewater treatment and their dissemination into lake geneva, Switzerland. Front Microbiol. 2012;3:106. https://doi.org/10.3389/fmicb.2012.00106
- Xu C, Kong L, Gao H, Cheng X, Wang X. A review of current bacterial resistance to antibiotics in food animals. Front Microbiol. 2022;13:822689. https://doi.org/10.3389/ fmicb.2022.822689
- Van Boeckel TP, Gandra S, Ashok A, Caudron Q, Grenfell BT, Levin SA, et al. Global antibiotic consumption 2000 to 2010: an analysis of national pharmaceutical sales data. Lancet Infect Dis. 2014;14:742-50. https://doi.org/10.1016/S1473-3099(14)70780-7
- Mulchandani R, Wang Y, Gilbert M, Van Boeckel TP. Global trends in antimicrobial use in food-producing animals: 2020 to 2030. PLOS Glob Public Health. 2023;3:e0001305. https:// doi.org/10.1371/journal.pgph.0001305
- Larsson DGJ, Flach CF. Antibiotic resistance in the environment. Nat Rev Microbiol. 2022;20:257-69. https://doi.org/10.1038/s41579-021-00649-x
- He Y, Yuan Q, Mathieu J, Stadler L, Senehi N, Sun R, et al. Antibiotic resistance genes from livestock waste: occurrence, dissemination, and treatment. NPJ Clean Water. 2020;3:4. https:// doi.org/10.1038/s41545-020-0051-0
- 22. Jeżak K, Kozajda A. Occurrence and spread of antibiotic-resistant bacteria on animal farms and in their vicinity in Poland and Ukraine. Environ Sci Pollut Res. 2022;29:9533-59. https://doi. org/10.1007/s11356-021-17773-z
- Argudín MA, Deplano A, Meghraoui A, Dodémont M, Heinrichs A, Denis O, et al. Bacteria from animals as a pool of antimicrobial resistance genes. Antibiotics. 2017;6:12. https://doi. org/10.3390/antibiotics6020012
- Manyi-Loh C, Mamphweli S, Meyer E, Okoh A. Antibiotic use in agriculture and its consequential resistance in environmental sources: potential public health implications. Molecules. 2018;23:795. https://doi.org/10.3390/molecules23040795
- van den Bogaard AE, Stobberingh EE. Antibiotic usage in animals: impact on bacterial resistance and public health. Drugs. 1999;58:589-607. https://doi.org/10.2165/00003495-199958040-00002
- Guo K, Zhao Y, Cui L, Cao Z, Zhang F, Wang X, et al. The influencing factors of bacterial resistance related to livestock farm: sources and mechanisms. Front Anim Sci. 2021;2:650347. https://doi.org/10.3389/fanim.2021.650347
- Nikaido H. Multidrug resistance in bacteria. Annu Rev Biochem. 2009;78:119-46. https://doi. org/10.1146/annurev.biochem.78.082907.145923
- Kirbis A, Krizman M. Spread of antibiotic resistant bacteria from food of animal origin to humans and vice versa. Procedia Food Sci. 2015;5:148-51. https://doi.org/10.1016/ j.profoo.2015.09.043
- Jian Z, Zeng L, Xu T, Sun S, Yan S, Yang L, et al. Antibiotic resistance genes in bacteria: occurrence, spread, and control. J Basic Microbiol. 2021;61:1049-70. https://doi.org/10.1002/ jobm.202100201
- Spellberg B, Hansen GR, Kar A, Cordova CD, Price LB, Johnson JR. Antibiotic resistance in humans and animals. NAM Perspect. 2016. https://doi.org/10.31478/201606d
- 31. Mulani MS, Kamble EE, Kumkar SN, Tawre MS, Pardesi KR. Emerging strategies to combat ESKAPE pathogens in the era of antimicrobial resistance: a review. Front Microbiol.

2019;10:539. https://doi.org/10.3389/fmicb.2019.00539

- 32. De Oliveira DM, Forde BM, Kidd TJ, Harris PNA, Schembri MA, Beatson SA, et al. Antimicrobial resistance in ESKAPE pathogens. Clin Microbiol Rev. 2020;33:e00181-19. https://doi.org/10.1128/cmr.00181-19
- Mancuso G, Midiri A, Gerace E, Biondo C. Bacterial antibiotic resistance: the most critical pathogens. Pathogens. 2021;10:1310. https://doi.org/10.3390/pathogens10101310
- Breijyeh Z, Jubeh B, Karaman R. Resistance of Gram-negative bacteria to current antibacterial agents and approaches to resolve it. Molecules. 2020;25:1340. https://doi.org/10.3390/ molecules25061340
- Santajit S, Indrawattana N. Mechanisms of antimicrobial resistance in ESKAPE pathogens. BioMed Res Int. 2016;2016:2475067. https://doi.org/10.1155/2016/2475067
- 36. Abdelaziz SM, Aboshanab KM, Yahia IS, Yassien MA, Hassouna NA. Correlation between the antibiotic resistance genes and susceptibility to antibiotics among the carbapenemresistant Gram-negative pathogens. Antibiotics. 2021;10:255. https://doi.org/10.3390/ antibiotics10030255
- 37. Coculescu BI. Antimicrobial resistance induced by genetic changes. J Med Life. 2009;2:114-23.
- 38. Institute of Medicine, Forum on Emerging Infections, Board on Global Health. The resistance phenomenon in microbes and infectious disease vectors: implications for human health and strategies for containment: workshop summary. Washington, DC: The National Academies Press; 2003.
- Duong DV, Binns CW, Le TV. Availability of antibiotics as over-the-counter drugs in pharmacies: a threat to public health in Vietnam. Trop Med Int Health. 1997;2:1133-9. https:// doi.org/10.1046/j.1365-3156.1997.d01-213.x
- Cars O, Nordberg P. Antibiotic resistance the faceless threat. Int J Risk Saf Med. 2005; 17:103-10.
- Wang F, Fu YH, Sheng HJ, Topp E, Jiang X, Zhu YG, et al. Antibiotic resistance in the soil ecosystem: a One Health perspective. Curr Opin Environ Sci Health. 2021;20:100230. https:// doi.org/10.1016/j.coesh.2021.100230
- Collignon P, Beggs JJ. Socioeconomic enablers for contagion: factors impelling the antimicrobial resistance epidemic. Antibiotics. 2019;8:86. https://doi.org/10.3390/ antibiotics8030086
- Michael CA, Dominey-Howes D, Labbate M. The antimicrobial resistance crisis: causes, consequences, and management. Front Public Health. 2014;2:145. https://doi.org/10.3389/ fpubh.2014.00145
- Pruden A, Pei R, Storteboom H, Carlson KH. Antibiotic resistance genes as emerging contaminants: studies in northern Colorado. Environ Sci Technol. 2006;40:7445-50. https:// doi.org/10.1021/es0604131
- Bombaywala S, Mandpe A, Paliya S, Kumar S. Antibiotic resistance in the environment: a critical insight on its occurrence, fate, and eco-toxicity. Environ Sci Pollut Res. 2021;28:24889-916. https://doi.org/10.1007/s11356-021-13143-x
- Rinke C, Schwientek P, Sczyrba A, Ivanova NN, Anderson IJ, Cheng JF, et al. Insights into the phylogeny and coding potential of microbial dark matter. Nature. 2013;499:431-7. https://doi. org/10.1038/nature12352
- Schulz F, Eloe-Fadrosh EA, Bowers RM, Jarett J, Nielsen T, Ivanova NN, et al. Towards a balanced view of the bacterial tree of life. Microbiome. 2017;5:140. https://doi.org/10.1186/ s40168-017-0360-9
- 48. Tiedje JM, Wang F, Manaia CM, Virta M, Sheng H, Ma L, et al. Antibiotic resistance genes

in the human-impacted environment: a One Health perspective. Pedosphere. 2019;29:273-82. https://doi.org/10.1016/S1002-0160(18)60062-1

- Davies J, Davies D. Origins and evolution of antibiotic resistance. Microbiol Mol Biol Rev. 2010;74:417-33. https://doi.org/10.1128/mmbr.00016-10
- Symochko L, Demyanyuk O, Symochko V, Grulova D, Fejer J, Mariychuk R. The spreading of antibiotic-resistant bacteria in terrestrial ecosystems and the formation of soil resistome. Land. 2023;12:769. https://doi.org/10.3390/land12040769
- Zhang XX, Zhang T, Fang HHP. Antibiotic resistance genes in water environment. Appl Microbiol Biotechnol. 2009;82:397-414. https://doi.org/10.1007/s00253-008-1829-z
- 52. Taviani E, Ceccarelli D, Lazaro N, Bani S, Cappuccinelli P, Colwell RR, et al. Environmental Vibrio spp., isolated in Mozambique, contain a polymorphic group of integrative conjugative elements and class 1 integrons. FEMS Microbiol Ecol. 2008;64:45-54. https://doi.org/10. 1111/j.1574-6941.2008.00455.x
- Volkmann H, Schwartz T, Bischoff P, Kirchen S, Obst U. Detection of clinically relevant antibiotic-resistance genes in municipal wastewater using real-time PCR (TaqMan). J Microbiol Methods. 2004;56:277-86. https://doi.org/10.1016/j.mimet.2003.10.014
- Auerbach EA, Seyfried EE, McMahon KD. Tetracycline resistance genes in activated sludge wastewater treatment plants. Water Res. 2007;41:1143-51. https://doi.org/10.1016/ j.watres.2006.11.045
- 55. Caplin JL, Hanlon GW, Taylor HD. Presence of vancomycin and ampicillin-resistant Enterococcus faecium of epidemic clonal complex-17 in wastewaters from the south coast of England. Environ Microbiol. 2008;10:885-92. https://doi.org/10.1111/j.1462-2920.2007.01507.x
- Habibur Rahman M, Nonaka L, Tago R, Suzuki S. Occurrence of two genotypes of tetracycline (TC) resistance gene tet(M) in the TC-resistant bacteria in marine sediments of Japan. Environ Sci Technol. 2008;42:5055-61. https://doi.org/10.1021/es702986y
- Mohapatra H, Mohapatra SS, Mantri CK, Colwell RR, Singh DV. Vibrio cholerae non-O1, non-O139 strains isolated before 1992 from Varanasi, India are multiple drug resistant, contain intSXT, dfr18 and aadA5 genes. Environ Microbiol. 2008;10:866-73. https://doi.org/10.1111/ j.1462-2920.2007.01502.x
- Das BK, Behera BK, Chakraborty HJ, Paria P, Gangopadhyay A, Rout AK, et al. Metagenomic study focusing on antibiotic resistance genes from the sediments of River Yamuna. Gene. 2020;758:144951. https://doi.org/10.1016/j.gene.2020.144951
- Martinez JL. Antibiotics and antibiotic resistance genes in natural environments. Science. 2008;321:365-7. https://doi.org/10.1126/science.1159483
- Zhang C, Qiu S, Wang Y, Qi L, Hao R, Liu X, et al. Higher isolation of NDM-1 producing Acinetobacter baumannii from the sewage of the hospitals in Beijing. PLOS ONE. 2013; 8:e64857. https://doi.org/10.1371/journal.pone.0064857
- 61. Li J, Cao J, Zhu YG, Chen QL, Shen F, Wu Y, et al. Global survey of antibiotic resistance genes in air. Environ Sci Technol. 2018;52:10975-84. https://doi.org/10.1021/acs.est.8b02204
- Frade VMF, Dias M, Teixeira ACSC, Palma MSA. Environmental contamination by fluoroquinolones. Braz J Pharm Sci. 2014;50:41-54. https://doi.org/10.1590/S1984-82502011000100004
- 63. Sanderson H, Brown RS, Hania P, McAllister TA, Majury A, Liss SN. Antimicrobial resistant genes and organisms as environmental contaminants of emerging concern: addressing global public health risks. In: Roig B, Weiss K, Thireau V, editors. Management of emerging public health issues and risks: multidisciplinary approaches to the changing environment. San diego,

CA: Academic Press; 2019. p. 147-87.

- 64. Palmer AC, Angelino E, Kishony R. Chemical decay of an antibiotic inverts selection for resistance. Nat Chem Biol. 2010;6:105-7. https://doi.org/10.1038/nchembio.289
- Tran NH, Reinhard M, Gin KYH. Occurrence and fate of emerging contaminants in municipal wastewater treatment plants from different geographical regions-a review. Water Res. 2018;133:182-207. https://doi.org/10.1016/j.watres.2017.12.029
- 66. Milić N, Milanović M, Letić NG, Sekulić MT, Radonić J, Mihajlović I, et al. Occurrence of antibiotics as emerging contaminant substances in aquatic environment. Int J Environ Health Res. 2013;23:296-310. https://doi.org/10.1080/09603123.2012.733934
- Harrabi M, Varela Della Giustina S, Aloulou F, Rodriguez-Mozaz S, Barceló D, Elleuch B. Analysis of multiclass antibiotic residues in urban wastewater in Tunisia. Environ Nanotechnol Monit Manag. 2018;10:163-70. https://doi.org/10.1016/j.enmm.2018.05.006
- Kookana RS, Williams M, Boxall ABA, Larsson DGJ, Gaw S, Choi K, et al. Potential ecological footprints of active pharmaceutical ingredients: an examination of risk factors in low-, middle- and high-income countries. Philos Trans R Soc B Biol Sci. 2014;369:20130586. https://doi.org/10.1098/rstb.2013.0586
- Piña B, Bayona JM, Christou A, Fatta-Kassinos D, Guillon E, Lambropoulou D, et al. On the contribution of reclaimed wastewater irrigation to the potential exposure of humans to antibiotics, antibiotic resistant bacteria and antibiotic resistance genes – NEREUS COST Action ES1403 position paper. J Environ Chem Eng. 2020;8:102131. https://doi.org/10.1016/ j.jece.2018.01.011
- Samples ME, Byom MJ, Daida JM. Parameter sweeps for exploring parameter spaces of genetic and evolutionary algorithms. In: Lobo FG, Lima CF, Michalewicz Z, editors. Parameter setting in evolutionary algorithms. Berlin: Springer; 2007. p. 161-84.
- Jiang Y, Wang Y, Hua X, Qu Y, Peleg AY, Yu Y. Pooled plasmid sequencindg reveals the relationship between mobile genetic elements and antimicrobial resistance genes in clinically isolated Klebsiella pneumoniae. Genomics Proteomics Bioinformatics. 2020;18:539-48. https:// doi.org/10.1016/j.gpb.2020.12.002
- Gu Y, Shen S, Han B, Tian X, Yang F, Zhang K. Family livestock waste: an ignored pollutant resource of antibiotic resistance genes. Ecotoxicol Environ Saf. 2020;197:110567. https://doi. org/10.1016/j.ecoenv.2020.110567
- 73. de Been M, Lanza VF, de Toro M, Scharringa J, Dohmen W, Du Y, et al. Dissemination of cephalosporin resistance genes between Escherichia coli strains from farm animals and humans by specific plasmid lineages. PLOS Genet. 2014;10:e1004776. https://doi.org/10.1371/journal. pgen.1004776
- 74. Hu Y, Yang X, Li J, Lv N, Liu F, Wu J, et al. The bacterial mobile resistome transfer network connecting the animal and human microbiomes. Appl Environ Microbiol. 2016;82:6672-81. https://doi.org/10.1128/AEM.01802-16
- 75. Tzouvelekis LS, Markogiannakis A, Psichogiou M, Tassios PT, Daikos GL. Carbapenemases in Klebsiella pneumoniae and other Enterobacteriaceae: an evolving crisis of global dimensions. Clin Microbiol Rev. 2012;25:682-707. https://doi.org/10.1128/cmr.05035-11
- Partridge SR, Kwong SM, Firth N, Jensen SO. Mobile genetic elements associated with antimicrobial resistance. Clin Microbiol Rev. 2018;31:e00088-17. https://doi.org/10.1128/ CMR.00088-17
- 77. Wegener HC. Antibiotic resistance—linking human and animal health. In: Institute of Medicine, editor. Improving food safety through a One Health approach: workshop summary. Washington, DC: The National Academies Press; 2012. p. 331-49.

- Chokshi A, Sifri Z, Cennimo D, Horng H. Global contributors to antibiotic resistance. J Glob Infect Dis. 2019;11:36-42. https://doi.org/10.4103/jgid_jgid_110_18
- Antimicrobial Resistance Collaborators. Global burden of bacterial antimicrobial resistance in 2019: a systematic analysis. Lancet. 2022;399:629-55. https://doi.org/10.1016/S0140-6736(21)02724-0
- Founou RC, Founou LL, Essack SY. Clinical and economic impact of antibiotic resistance in developing countries: a systematic review and meta-analysis. PLOS ONE. 2017;12:e0189621. https://doi.org/10.1371/journal.pone.0189621
- Bassetti M, Peghin M, Vena A, Giacobbe DR. Treatment of infections due to MDR Gramnegative bacteria. Front Med. 2019;6:74. https://doi.org/10.3389/fmed.2019.00074
- Annavajhala MK, Gomez-Simmonds A, Uhlemann AC. Multidrug-resistant Enterobacter cloacae complex emerging as a global, diversifying threat. Front Microbiol. 2019;10:44. https:// doi.org/10.3389/fmicb.2019.00044
- Llor C, Bjerrum L. Antimicrobial resistance: risk associated with antibiotic overuse and initiatives to reduce the problem. Ther Adv Drug Saf. 2014;5:229-41. https://doi. org/10.1177/2042098614554919
- Apata DF. Antibiotic resistance in poultry. Int J Poult Sci. 2009;8:404-8. https://doi. org/10.3923/ijps.2009.404.408
- Barton MD. Impact of antibiotic use in the swine industry. Curr Opin Microbiol. 2014;19:9-15. https://doi.org/10.1016/j.mib.2014.05.017
- Pfaller MA. Flavophospholipol use in animals: positive implications for antimicrobial resistance based on its microbiologic properties. Diagn Microbiol Infect Dis. 2006;56:115-21. https://doi. org/10.1016/j.diagmicrobio.2006.03.014
- Queenan K, Häsler B, Rushton J. A One Health approach to antimicrobial resistance surveillance: is there a business case for it? Int J Antimicrob Agents. 2016;48:422-7. https://doi. org/10.1016/j.ijantimicag.2016.06.014
- Ashbolt NJ, Amézquita A, Backhaus T, Borriello P, Brandt KK, Collignon P, et al. Human Health Risk Assessment (HHRA) for environmental development and transfer of antibiotic resistance. Environ Health Perspect. 2013;121:993-1001. https://doi.org/10.1289/ehp.1206316
- Berendonk TU, Manaia CM, Merlin C, Fatta-Kassinos D, Cytryn E, Walsh F, et al. Tackling antibiotic resistance: the environmental framework. Nat Rev Microbiol. 2015;13:310-7. https:// doi.org/10.1038/nrmicro3439
- Banerji A, Jahne M, Herrmann M, Brinkman N, Keely S. Bringing community ecology to bear on the issue of antimicrobial resistance. Front Microbiol. 2019;10:2626. https://doi. org/10.3389/fmicb.2019.02626
- Gaze WH, Krone SM, Joakim Larsson DG, Li XZ, Robinson JA, Simonet P, et al. Influence of humans on evolution and mobilization of environmental antibiotic resistome. Emerg Infect Dis. 2013;19:e120871. https://doi.org/10.3201/eid1907.120871
- 92. World Health Organization. Global action plan on antimicrobial resistance. Geneva: WHO Document Production Services; 2015.
- 93. World Health Organization. Integrated surveillance of antimicrobial resistance in foodborne bacteria: application of a One Health approach: guidance from the WHO Advisory Group on Integrated Surveillance of Antimicrobial Resistance (AGISAR). Geneva: WHO; 2017. p. 88.
- Tacconelli E, Carrara E, Savoldi A, Harbarth S, Mendelson M, Monnet DL, et al. Discovery, research, and development of new antibiotics: the WHO priority list of antibiotic-resistant bacteria and tuberculosis. Lancet Infect Dis. 2018;18:318-27. https://doi.org/10.1016/S1473-3099(17)30753-3