Bacteriophages Contribute to the Spread of Antibiotic Resistance Genes among Foodborne Pathogens of the Enterobacteriaceae Family – A Review

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Foodborne illnesses continue to have an economic impact on global health care systems. There is a growing concern regarding the increasing frequency of antibiotic resistance in foodborne bacterial pathogens and how such resistance may affect treatment outcomes. In an effort to better understand how to reduce the spread of resistance, many research studies have been conducted regarding the methods by which antibiotic resistance genes are mobilized and spread between bacteria. Transduction by bacteriophages (phages) is one of many horizontal gene transfer mechanisms, and recent findings have shown phage-mediated transduction to be a significant contributor to dissemination of antibiotic resistance genes. Here, we review the viability of transduction as a contributing factor to the dissemination of antibiotic resistance genes in foodborne pathogens of the Enterobacteriaceae family, including non-typhoidal Salmonella and Shiga toxin-producing Escherichia coli, as well as environmental factors that increase transduction of antibiotic resistance genes.

Keywords: bacteriophage, transduction, antibiotic resistance, foodborne pathogens, horizontal gene transfer, Escherichia coli O157:H7, non-typhoidal Salmonella, environment

INTRODUCTION

The World Health Organization (WHO) estimates that in 2010, foodborne illnesses affected 600 million people and caused 420,000 deaths globally (World Health Organization [WHO], 2015). In Canada, 4 million cases of foodborne illnesses, 11,600 hospitalizations and 238 deaths are estimated to occur annually (Thomas et al., 2015). In the United States, 47.8 million cases of foodborne illness are estimated to occur annually with 127,839 hospitalizations and 3,037 deaths (Scallan et al., 2011). The economic impact of foodborne illness in the USA is estimated at approximately $10–83 billion (USD) annually (Mclinden et al., 2014). A major contributor to the costs associated with foodborne disease is healthcare costs. The continued emergence of antibiotic resistance, and especially multidrug resistance, among foodborne pathogens may contribute...
to unsuccessful treatment outcomes, thereby increasing costs associated with foodborne disease. For example, in the United States, antibiotic resistant foodborne infections cause 430,000 illnesses annually (Centers for Disease Control and Prevention [CDC], 2015a). Therefore, there is a growing need to better understand the mechanisms, frequency, reservoirs, and vectors governing the transfer of antibiotic resistance determinants in major foodborne pathogens in hopes of controlling the dissemination of antibiotic resistance.

Non-typhoidal Salmonella is the most prevalent bacterial foodborne pathogen resulting in hospitalization in the western world (Thomas et al., 2015). Shiga toxin-producing Escherichia coli (STEC) are also important causes of foodborne illness that frequently result in hospitalization (Majowicz et al., 2014). Outbreaks linked to these pathogens have been associated with various foods. Salmonella outbreaks have been linked to contaminated poultry, eggs, cheese, ice cream, fresh produce, and chocolate, while STEC outbreaks have been primarily linked with fresh produce and ground beef (Kapperud et al., 1990; Voetsch et al., 2004; Werber et al., 2005; Centers for Disease Control and Prevention [CDC], 2015b; Herman et al., 2015).

In 5% of gastroenteritis cases caused by NTS, the patient will develop bacteremia or other extraintestinal complications such as urinary tract infections, pneumonia, endocarditis, meningitis, and cellulitis (Eng et al., 2015). In the 1980s, NTS was treated with ampicillin, chloramphenicol and TMP-SMZ but by the 1990s widespread resistance emerged (Acheson and Hohmann, 2001). Currently, fluoroquinolones and third-generation cephalosporins are mainly administered but resistance to these antibiotics is also emerging (Eng et al., 2015). Even more concern materialized when multidrug resistant strains of Salmonella Typhimurium phage type DT104 emerged. This strain has been associated with poultry, cattle, and swine, and is defined by ACSSuT (Threlfall, 2000). Data from the National Antimicrobial Resistance Monitoring System (NARMS) indicated that 8.5% of NTS isolated from humans between 1999 and 2004, and 3.8% of retail meat isolates from 2002 to 2004 had the ACSSuT phenotype and Salmonella Typhimurium and Salmonella Newport were the most common serovar with the penta-resistance pattern (Whichard et al., 2010). In fact, Salmonella was the most prevalent foodborne pathogen involved in antibiotic resistant outbreaks between 1973 and 2011, and S. Typhimurium was the most frequent serovar (Dewaal and Grooters, 2013).

In contrast, antibiotic treatment for STEC is controversial since some studies have observed certain classes of antibiotics to be effective while others increase the production of Stx due to the induction of Stx carrying prophages, which could trigger complications such as hemolytic uremic syndrome (HUS) (Kurioka et al., 1999; Bielaszewska et al., 2012). Still, antibiotic resistance among STEC strains has significantly increased since the first identification of STEC as a foodborne pathogen in 1982 (Kim et al., 1994; Meng et al., 1998; Galland et al., 2001; Schroeder et al., 2002; Vidovic and Korber, 2006), likely due to overuse of antibiotics in food producing animals such as ruminants, which are the animal reservoir for STEC (Hunt, 2010). For example, in 1988, one study indicated that 2.9% of E. coli O157:H7 isolates (the predominant STEC serotype, accounting for the majority of STEC cases in North America) from human cases were resistant to antibiotics (Ratnam et al., 1988). As of 2007, 79.8% of E. coli O157:H7 isolates from bovine and human feces, bovine milk products, ground beef, and cider carried one or more antibiotic resistance genes (Srinivasan et al., 2007). Although high percentages of antibiotic resistance have not been historically associated with STEC, additional studies support the increasing presence of antibiotic resistance in E. coli O157:H7 isolates. For example, Ferreira et al. (2015) isolated 90 E. coli O157 isolates from health sheep, and observed that 75 of the isolates (83.3%) were resistant to at least one antibiotic. In another study, antibiotic profiles of 95 fecal isolates of E. coli O157:H7 collected from two commercial dairy farms in South Africa demonstrated that bla\textsubscript{ AmpC } (90%), strA (80%), tetA (70%), bla\textsubscript{ CMY } (70%), and bla\textsubscript{ CTX-M } (65%) were predominant (Iwieriebor et al., 2015). El-Shatory et al. (2015) also demonstrated that of 44 E. coli O157:H7 isolates, 100% were resistant to amoxicillin and 77% were resistant to clarithromycin.

Horizontal gene transfer mechanisms responsible for the increased spread of antibiotic resistance to foodborne bacterial pathogens have been well studied. Conjugation, transformation, and transduction are the primary mechanisms by which dissemination of antibiotic resistance genes occurs (von Wintersdorff et al., 2016). The notion that phage mediated transduction is a major driver of horizontal transfer of antibiotic resistance genes between foodborne pathogens, as well as from the environment to animals and humans, is increasingly becoming recognized. Phages are the most abundant organism in the biosphere, and are found in diverse environments including oceans, lakes, soil, urban sewage, potable and well water and plant microbial communities (Clokie et al., 2011). Antibiotic resistance genes are often found on various MGEs, such as plasmids, genomic islands and transposons, and, as such, can be horizontally transferred by phage transduction. This review summarizes the scientific literature in which horizontal transfer of antibiotic resistance genes by phage-mediated transduction has been described, with a specific focus on the role that phages play in horizontal transfer of antibiotic resistance genes among foodborne pathogens, including STEC O157:H7 and Salmonella enterica, as well as between environmental bacteria.

**PHAGE-MEDIATED TRANSDUCTION**

Transduction occurs by means of virulent and temperate phages (Figures 1A–C) (Suzuki and Griffiths, 2000; Feiner et al., 2015). Upon infection, temperate phages integrate their DNA into the host chromosome and the prophage may remain dormant in the host until some stress will induce the excision of the phage from the chromosome leading to subsequent formation of phage particles and lysis of the host cell (Figure 1A). Virulent phages do not integrate their DNA into the host chromosome but induce immediate formation of phage particles and lysis of the host cell (Figure 1B). Under unfavorable growth conditions, some phages can adopt a pseudolysogeny state where their genome does not degrade but instead exists as a plasmid within the cytoplasm and
FIGURE 1 | Phage life cycles and types of phage-mediated transduction. (A) Temperate phage life cycle; (B) Virulent phage life cycle; (C) Pseudolysogeny; (D) Generalized Transduction; (E) Specialized Transduction. Adapted from Feiner et al. (2015).

during bacterial cell division becomes incorporated into only one daughter cell (Figure 1C) (Feiner et al., 2015).

Two types of transduction, generalized and specialized, have been described. Generalized transduction refers to the mis packaging of bacterial DNA into the phage capsid (Figure 1D). The phage may then infect another susceptible host, thereby transferring genetic material to another bacterial cell where it will be integrated by homologous recombination. Specialized transduction is the improper excision of a prophage from the bacterial chromosome, which results in packaging of the bacterial DNA into phages at a higher frequency than generalized transduction (Figure 1E) (Griffiths, 2002). The lateral transfer of genes encoding antibiotic resistance by phage-mediated transduction could be an important contributing factor in the global spread of antibiotic resistance.

SHIGA TOXIN-PRODUCING Escherichia coli

Shiga toxin-producing Escherichia coli serotype O157:H7 was recognized as an important foodborne pathogen following two outbreaks that occurred in 1982 in the United States (Riley et al., 1983). Twenty-six cases were identified in Oregon, and 21 cases were reported in Michigan. The outbreaks were both linked to the consumption of hamburgers from a fast food restaurant chain, and E. coli O157:H7 was isolated from both patient stool samples and hamburger meat from the restaurant (Riley et al., 1983). Following the discovery of E. coli O157:H7, other serotypes of STEC have been identified, with many of these serotypes causing outbreaks and sporadic disease (Johnson et al., 2006). Numerous studies have detected the presence of antibiotic resistance genes in clinical and environmental STEC isolates. For example, resistance to streptomycin, sulfonamide, tetracycline, ampicillin, and cephalothin was found in isolates from human stool samples and in isolates procured from livestock and food (Kim et al., 1994; Schroeder et al., 2002). Multidrug resistant strains have also been observed with resistance to up to 10 different antimicrobials (Walsh et al., 2006).

Despite the non-specific nature of transduction, the phage mediated transfer of antibiotic resistance genes by transduction has been reported to occur in laboratory strains of E. coli (Table 1) (Tanyashin et al., 2003; Kenzaka et al., 2007; Marinus and Poteete, 2014). For instance, induction of Stx-converting generalized transducing phage 933W from E. coli 0157:H7 resulted in efficient transfer of tetracycline resistance genes to the E. coli laboratory strain, K-12 (Marinus and Poteete, 2014). In another study, Serra-Moreno et al. (2006) modified
TABLE 1 | Summary of phage-mediated transduction events demonstrated in E. coli.

| Donor organism | Recipient organism | Phage | Antibiotic resistance gene | Reference |
|----------------|--------------------|-------|----------------------------|-----------|
| E. coli B834 (pBR322) E. coli HMS174 (pBR322) | E. coli B834 | Pseudo-T even phages RB42 RB43 RB49 | Amp and tet determinants of plasmid BR322 | Tanyashin et al., 2003 |
| E. coli O157:H7 EDL933 | E. coli K12 AB1157 | 933W | tet | Marinus and Poteete, 2014 |
| E. coli NBRC 12713 KEN1 | E. coli NBRC 12713 | P1kc T4G7T | bla | Kenzaka et al., 2007 |
| E. coli NBRC 12713 KEN1 | E. coli strains (C600 RK2, HB101, NBRC 12713, and W3110) | EC10 | bla | Kenzaka et al., 2007 |
| N/A | E. coli strains DH5α and C600 | Stx2 producing phages induced from STEC isolated from cattle: 0A9, 0A312, 0A534, 0A549, 0A557, ØVTB55 and 933W | tet and cat | Sierra-Moreno et al., 2006 |
| N/A | E. coli ATCC 13706 | Sixty of 243 coliphages isolated from retail chicken meat could transduce antibiotic resistance genes | 43 phages: φKmr 8 phages: φCmr 4 phages: φTetr 3 phages: φKmrAmpr 2 phages: φKmrCmr | Shousha et al., 2015 |
| N/A | E. coli CSH-2 | P1kc | R factors: N-1: SU, SM, TC N-3: TC, SM, TC N-6: SU, SM, TC N-9: TC N-9: SM R-15: SM, SU S-a: CM, SM, CM S-b: SU, SM, CM, TC S-b: TC K: SU, SM, OM, TC K-R1: SU, SM, OM K-TC: TC | Watanabe et al., 1968 |
| E. coli CFT073 WAM3886 | E. coli CFT073 WAM2909 | Four phages isolated from wastewater effluents: φEB49, φEB47 φEB32, φEB5 | kan | Battaglioli et al., 2011 |

amp: ampicillin; tet, TC: tetracycline; bla: beta-lactamase; kan, φKmr: kanamycin; φCmr, CM: chloramphenicol; φKmr Ampr: kanamycin and ampicillin; φKmr Cmr: kanamycin and chloramphenicol; SU: sulfonamide; SM: streptomycin.

a Stx-converting bacteriophage, to incorporate genes encoding for resistance to tetracycline or chloramphenicol as markers for genetic transfer. Infection of E. coli DH5α and C600 with the recombinant phages resulted in production of transductants resistant to the respective antibiotics. In an attempt to explore the potential of phage mediated transduction as a vehicle of antibiotic resistance from food of animal origin to consumers, Shousha et al. (2015) tested retail chicken meat for the presence of coliphages capable of transducing antibiotic resistance genes. Of 243 coliphages, 24.7% were able to transduce one or more antibiotic genes, encoding for ampicillin, tetracycline, kanamycin and chloramphenicol, to the laboratory strain E. coli ATCC 13706 (Shousha et al., 2015). In another study, four phages isolated from wastewater effluent samples were observed to horizontally transfer a kanamycin resistance gene to strains of uropathogenic E. coli by generalized transduction (Battaglioli et al., 2011). Lastly, Kenzaka et al. (2007) developed a method termed CPRINS-FISH, to investigate the frequency of lateral gene transfer via transduction at the single-cell level, and demonstrated the transfer of the ampicillin resistance gene between E. coli cells at a surprisingly high frequency. Although phage-mediated horizontal transfer of antibiotic resistance genes between strains of STEC has not yet been directly shown, it is clear that Stx-phages can convert antibiotic susceptible E. coli strains to resistant strains in vitro. Therefore, it is reasonable to speculate that the transfer of antibiotic resistance genes via transduction occurs among Stx-producing E. coli serotypes, including O157:H7.

There have also been reports linking antibiotic resistance to specific phage types, suggesting a phage-mediated bias toward antibiotic resistance (Mora et al., 2005; Ziebell et al., 2008; Ziebell et al., 2011). In a Canadian study, Ziebell et al. (2011) characterized 187 isolates of STEC O157:H7 and observed that...
45 of these possessed resistance to streptomycin, sulfonamide, and tetracycline. Of the 45 isolates, 43 belonged to three phage types: PT23, PT45, and PT67. Another study using isolates from Spain also recorded an association between phage type and resistance to the same three antibiotics (Mora et al., 2005). However, phage types found to be linked to antibiotic resistance were different from the Canadian study and included PT21/28, PT23, PT34, and PT2. The proposed link between phage type and antibiotic resistance could suggest a potential role for phages in the dissemination of antibiotic resistance. While Ziebell et al. (2008) believed the resistance genes they studied were stably incorporated into the bacterial genome before the bacterium differentiated into different phage types, it is also possible that certain phage types are prone to acquiring and transferring select resistance genes.

Another instance of phage-mediated transduction of antibiotic resistance in E. coli, stems from the debated use of antimicrobials as growth promoters in food-producing animals (Economou and Gousia, 2015). Carbadox, marketed under the name MECADOX, is an agricultural antimicrobial added to swine feed in the United States to prevent swine dysentery, improve feed efficiency and promote weight gain and growth in swine (Bearson et al., 2014; Food and Drug Administration [FDA], 2015). A laboratory-based study established that Stx-converting prophages were induced by various concentrations of carbadox (0.5–8 ppms) from an E. coli C600 strain containing Stx-phage 933W and three other clinical Shiga toxin-producing E. coli strains (Köhler et al., 2000). The concentrations of carbadox identified to be sufficient for induction were approximately 0.5–100 times lower than that found in the gut content isolated from the intestine of carbadox-fed piglets (Graaf et al., 1988; Köhler et al., 2000). These findings raise concern because, as previously discussed, the Stx-phage 933W has the ability to transfer tetracycline resistant cassettes by generalized transduction, at least in a laboratory setting (Marinus and Poteete, 2014). Moreover, olaquindox, another agricultural antimicrobial permitted in China, was three times more capable of inducing phage 933W than carbadox (Köhler et al., 2000). Although these results were observed in the laboratory, carbadox could potentially induce prophages from E. coli strains in the environment. For example, Allen et al. (2011) demonstrated that carbadox was capable of inducing more prophages in bacterial strains isolated from the gut of carbadox-treated swine compared to that of untreated swine. The implications of phage transfer of antibiotic resistance genes in the food animal environment are clear. For example, once prophages, harbored within E. coli, are released, they could transfer antibiotic resistance genes to foodborne pathogens such as E. coli O157:H7 or Salmonella spp. within the gastrointestinal tract of cattle or swine. These now antibiotic resistant foodborne pathogens would be present in the stool of cattle or swine, potentially leading to environmental contamination of crops, thereby contributing to fresh produce outbreaks (Jung et al., 2014). On the other hand, during slaughter, resistant foodborne pathogens could contaminate meat products destined for human consumption, putting consumers at risk of antibiotic resistant infections.

Conversely, although studies appear to indicate that transduction of antibiotic resistance genes does occur, there is some uncertainty as to how much transduction contributes to overall lateral gene transfer. Using conventional plating methods, the estimates of the frequency of phage-mediated DNA transfer ranged between $10^{-9}$ and $10^{-5}$ per bacteriophage (Weinbauer and Rassoulzadegan, 2004; Kenzaka et al., 2007). However, using the CPRINS-FISH method, Kenzaka et al. (2007) showed that gene transfer by phages in E. coli cells occurs at a much higher frequency (ranging between $10^{-4}$ and $10^{-3}$ for the phages tested) than previously thought. They attributed the previous underestimation to the limitations in traditional culturing and enumeration methods. However, even in the best conditions, modeling of infection dynamics shows that the contribution of generalized transduction to lateral gene spread is small (Volkova et al., 2014). The model developed by Volkova et al. (2014) accounts for many factors that may contribute to transduction such as phage dynamics, lysogens, host range, and bacterial susceptibility to the antibiotic resistance gene of interest. This model was constructed for temperate phages infecting E. coli within the lumen of the bovine large intestine. However, virulent phages may play a role in transduction of antibiotic resistance as well. Furthermore, only phages infecting E. coli specifically were considered when developing the model, and though it may be an indicator, bacteria and phages are highly diverse. Hence, it is difficult to evaluate if predictions derived from the model are applicable across species and types. Finally, the model is restricted to the bovine large intestine and estimates the average E. coli residency time in the intestine to be on the order of 100 h. Though the presence of E. coli makes the intestine an optimal setting for transduction to occur, transduction is not limited to this location, and the possibility of phage mediated spread of antibiotic resistance genes occurring in the environment cannot be discounted. There are also exterior factors, such as the use of antibiotics that may induce excision of prophages. The number of factors that may, or may not, influence transduction and gene transfer is large, and therefore, transduction as a means of lateral gene transfer in E. coli should not be quickly dismissed.

Salmonella enterica

Multidrug resistance amongst Salmonella spp. have globally increased during the past two decades. For instance, in Great Britain, between 1994 and 2010, Salmonella Typhimurium was the most common serovar isolated from swine (Mueller-Doblies et al., 2013). When tested against a panel of antimicrobials, its resistance to six or more antimicrobials increased from 27.2% in 1994 to 58.3% in 2010. In Spain, antimicrobial resistance of different Salmonella serovars isolated from retail poultry increased from resistance to 3.98 antimicrobials in 1993 to 5 antimicrobials in 2006 (Alvarez-Fernandez et al., 2012). In China, 80% of Salmonella isolates from retail meats were resistant to one antimicrobial and 53% were resistant to more than three antimicrobials, with S. Enteritidis being the predominant serovar (Yang et al., 2010).
The prevalence of penta-resistant strains of *S. Typhimurium* phage type DT104 has been increasing exponentially since its emergence in 1989, particularly in Europe and North America (Threlfall, 2000; Helms et al., 2005). In phage type DT104, the penta-resistance genes are clustered on a 43-kb SGI1, which is flanked by two type I integrons (Boyd et al., 2002). Horizontal transfer of the penta-resistance is hypothesized to be facilitated by two P22-like prophages, ST104 or PDT17, harbored within DT104 (Schmieger and Schicklmaier, 1999; Tanaka et al., 2004). P22 is a well-known *Salmonella* phage extensively used in molecular biology for its ability to introduce foreign genes by generalized and specialized transduction (Zinder and Lederberg, 1952; Smith and Levine, 1965; Smith and Stocker, 1966; Smith-Keary, 1966; Wing, 1968). Schmieger and Schicklmaier (1999), determined that when P22-like phages, ES18 and PDT17, were released from DT104, they could transduce antibiotic resistance genes (Table 2). The researchers demonstrated the transduction of *amp* and *amp* by phage PDT17 and *amp*, *cm*, and *tet*, which confer resistance to ampicillin, chloramphenicol, and tetracycline, respectively, by ES18 from a donor DT104 strain into a DT104 recipient strain lacking these resistance genes. Also detected was the co-transduction of selected resistance genes by phage ES18. Most notably, of 145 *cam* transductants, all but one co-transduced *amp* and *tet*, and of 71 *tet* transductants, all but one co-transduced *amp* and *cam*. Furthermore, in 14 of 16 transductants, Schmieger and Schicklmaier (1999) observed that ES18 could co-transduce *sul* and *str*, genes involved in resistance to sulfonamides and streptomycin, respectively, together with *amp*, *cam*, and *tet* to create the ACSSuT resistance phenotype (1). This co-transduction most likely occurs because *amp* and *str* are located on the integrons flanking SGI1, and the phage likely packages the SGI1 and its flanking integrons (Ridley and Threlfall, 1998; Sandvang et al., 1998; Mulvey et al., 2006). In another study, Bearson et al. (2014) investigated whether a mixture of phages induced from *S. Typhimurium* DT104 by carbadox, an agricultural antimicrobial discussed above, could transduce SGI1 to a DT104 strain with sensitivity to ampicillin, chloramphenicol, and tetracycline due to an internal deletion within SGI1. The authors observed the transduction of tetracycline resistance and 100% co-transduction

| Donor organism | Recipient organism | Phage | Antibiotic resistance gene | Reference |
|----------------|--------------------|-------|---------------------------|-----------|
| *S. Typhimurium* DT104 strain DT17 | *S. Typhimurium* DT104 strain DT16 | ES18 | *amp*, *cam* | Schmieger and Schicklmaier, 1999 |
| Resistance profile | Resistance profile | PDT17 | *amp*, *cam*, *tet* | |
| ACS(sp)SuT | SSu | ES18 co-transduction: *Amp cam* | |
| *S. Typhimurium* DT104 NCTC13348 | *S. Typhimurium* DT104 strain BBS 1012 | Phages induced by carbadox from donor organism | Transduction: *tetG* | Bearson et al., 2014 |
| Internal deletion in SI-1 for ampicillin, chloramphenicol, and tetracycline | | | Co-transduction: *floR bB<sub>osp</sub>-1* | |
| N/A | *S. Typhimurium* LT2 | P22 | R factors: N-1: TC N-3: TC, SU, SM N-3: SU, SM, TC N-3: SU, SM N-6: SU, SM, TC N-9: TC N-9: TC, SU, SM S-a: CM, SU, SM K: TC K-TC: TC | Watanabe et al., 1968 |
| | *S. Heidelberg* S25 | *S. Typhimurium* MZ1262 | P24 | *bB<sub>osp</sub>-2* | Zhang and Lejeune, 2008 |
| | | | | *tet(A)* | |
| | | | | *tet(B)* | |
| | *S. Typhimurium* DT120 (BBS 1162) | *S. Typhimurium* BBS 243 | Fluoroquinolone induced phages from donor organisms | Plasmid conferring kanamycin resistance native to donor organisms | Bearson and Brunelle, 2015 |
| | *S. Typhimurium* DT104 (BBS 1170) | | | | |
| N/A | Soil coliforms | Isolated and enriched biosolid phages | Cefoxitin Sulfamethazine | Ross and Topp, 2015 |

*amp*, *bla<sub>osp</sub>-1*: ampicillin; *tet*, *tet* TC: tetracycline; *cam*, *floR*, *floR* CM: chloramphenicol; *kan*, *Kmr*: kanamycin; *sul*, *SU*: sulfonamide; *str*, *SM*: streptomycin; *bla<sub>CMY-2</sub>*: beta-lactamase; *Kmr Amp*: kanamycin and ampicillin; *Kmr Cmr*: kanamycin and chloramphenicol.
of floR encoding for chloramphenicol and \( \text{bla}_{\text{pse}-1} \) encoding for ampicillin, as seen by Schmieg and Schickmaier (1999). These authors suggested that the carbadox induced phage transduction in \( S. \) Typhimurium DT104 and another ACSSuT resistant strain, DT120, is common after observing frequent transduction events of the \( \text{his} \) operon. Phage P22 was detected within several DT104 and DT120 isolates shown to be capable of generalized transduction. The absence of transduction following carbadox induction of a DT104 strain with a deletion of a P22-like prophage suggests that P22-like prophages are responsible for generalized transduction in DT104. This was further supported by the incapability of transduction in four \( S. \) Typhimurium strains that do not contain P22-like phages. These studies provide evidence that transduction and co-transduction by P22-like prophages of antibiotic resistance genes co-located within SGII in multidrug-resistant \( S. \) Typhimurium strains, is a common phenomenon. Moreover, genome scanning demonstrated that P22-like prophages are common in 18 \( S. \) enterica serovars suggesting that generalized transduction may be underestimated (Bearson et al., 2014).

\( R \) factors may represent another form of MGE that can be horizontally transferred via phage-mediated transduction. An \( R \) factor is an episome that harbors one or more antibiotic resistance determinants. \( R \) factors were first discovered in Japan in the 1940s when resistance emerged to sulfonamides, which were being used to treat dysentery caused by \( S. \) shigella spp. (Mitsushashi, 1993). Shortly after, streptomycin, chloramphenicol, and tetracycline became the treatments of choice but resistance to these antibiotics soon emerged as a result of an \( R \) factor harbored within \( S. \) shigella strains (Ochiai et al., 1959; Akiba et al., 1960; Nakaya et al., 1960). There have been numerous studies demonstrating that conjugative \( R \) factors spread antibiotic resistance genes between \( S. \) enterica serovars, as well as \( E. \) coli strains (Ishiguro et al., 1980; Balis et al., 1996; Lázaro et al., 2004), and one study reported that the transfer occurred by transduction in \( S. \) enterica as well as \( E. \) coli (Watanabe et al., 1968). In this study, Watanabe et al. (1968) isolated various \( R \) factors from \( S. \) shigella strains with different MICs of resistance to sulfanilamide, streptomycin, chloramphenicol, and tetracycline, and demonstrated successful transduction of the \( R \) factors by phage P22 to \( S. \) Typhimurium LT-2, as well as to \( E. \) coli CSH-2 by phage P1. Following transduction of the \( R \) factors, the group attempted to transfer the \( R \) factor by conjugation. Many of the transductants were unable to transmit their drug resistance genes suggesting that the \( R \) factor had integrated into the host's chromosome (Watanabe et al., 1968). The P22-like phages epsilon34, ES18, P22, ST104, and ST64T are all inducible and frequently lysogenize \( S. \) enterica (Kropinski et al., 2007). The integration of \( R \) factors into or in close proximity to P22-like prophages would facilitate the packaging of the \( R \) factor into the head of the assembling phage during induction from its host, thus contributing to the spread of antibiotic resistance within bacteria capable of causing foodborne illnesses, in the intestinal flora of livestock and in the environment.

Another aspect that is of concern is the worldwide increase in resistance to third-generation cephalosporins, which are used to treat invasive \( S. \) enterica infection (Eller et al., 2013; Seiffert et al., 2013; Noda et al., 2015; Liakopoulos et al., 2016). Resistance to cephalosporins is mainly due to ESBLs, such as TEM-, SHV-, and CTX-M, or plasmid mediated AmpC \( \beta \)-lactamases (pAmpCs), such as CMY, encoded on transferrable conjugative plasmids (Carattoli et al., 2002; Guerra et al., 2002; Chen et al., 2004). Still, Zhang and Lejeune (2008) demonstrated that they could also be transferred by generalized transduction. In this work, the authors observed that phage P24, induced from an isolate of \( S. \) Typhimurium, was capable of propagating on a multidrug resistant strain, \( S. \) Heidelberg (S25). Therefore, S25 harboring phage P24 was used as transduction donor to transfer ESBL and tetracycline resistance genes to a recipient \( S. \) Typhimurium isolate. PCR confirmed the presence of \( \text{bla}_{\text{CMY}-2} \), \( \text{tet}(A) \), and \( \text{tet}(B) \) in various \( S. \) Typhimurium transductants. Tetracycline genes were not co-transduced with \( \text{bla}_{\text{CMY}-2} \), however, their transduction frequency was equivalent, indicating generalized transduction. This study is the first report of antibiotic resistance genes transferred by phage-mediated transduction between different \( S. \) enterica serovars. Although this may be the first documented report, it is likely that cross-serovar transduction occurs frequently because phages can bind to various surface protein receptors on different species and serovars. The LPS, FlIC, OmpC, OmpF, OmpA, are examples of phage receptors present in STEC and \( S. \) enterica (Rakhuba et al., 2010). As further evidence, Zhang and Lejeune (2008) observed that 13 inducible phages recovered from 31 \( S. \) enterica serovars were capable of propagating on two or more \( S. \) enterica serovars including those often responsible for outbreaks such as \( S. \) Heidelberg, \( S. \) Enteritidis, and \( S. \) Typhimurium as well as \( S. \) Kentucky, which is one of the most common serovars detected in chickens and ground chicken meat in Europe. This study demonstrates that phage-mediated transduction can contribute to the spread of antibiotic resistance in various \( S. \) enterica serovars associated with foodborne outbreaks.

The other antibiotic of choice for treatment of invasive \( S. \) enterica infections is ciprofloxacin, a fluoroquinolone (Eng et al., 2015). Ciprofloxacin resistance has dramatically increased in food and clinical isolates (Lin et al., 2015). Its resistance in \( S. \) enterica is mainly attributed to double mutations in \( gyrA \) and a single mutation in \( parC \). In addition, it’s suggested that \( oqxAB \) operon is responsible for an increase in resistant clinical \( S. \) enterica strains (Wong et al., 2014). Bearson and Brunelle (2015) sought to investigate whether exposure to ciprofloxacin, as well as fluoroquinolones enrofloxacin and danofloxacin, which are used in veterinary medicine, would induce phage-mediated transfer of a native plasmid encoding kanamycin resistance by generalized transduction from \( S. \) enterica DT104 and DT102. The authors observed that all three fluoroquinolones stimulated generalized transduction of the kanamycin resistance plasmid to a strain of \( S. \) Typhimurium. Since \( S. \) Typhimurium frequently colonize cattle, swine and poultry without causing clinical disease, the authors expressed concern that there is potential for a scenario in which fluoroquinolones are administered to animals that are asymptotically colonized. This would induce phage-mediated generalized transduction and increase the horizontal transfer of antibiotic resistance genes. Moreover, the administered fluoroquinolones would be excreted in the
feces and urine of these food producing animals and stimulate horizontal gene transfer in the environment (Bearson and Brunelle, 2015).

As seen by the frequency of multi-drug resistance patterns, the transfer of resistance genes occurs extensively in strains of foodborne Salmonella, and transduction appears to play a prominent role in the successful dissemination of antibiotic resistance among the Salmonellae.

ENVIRONMENTAL FACTORS ASSOCIATED WITH BACTERIOPHAGE TRANSDUCTION OF ANTIBIOTIC RESISTANCE GENES

A number of factors contribute to the dissemination of antibiotic resistance such as the use and misuse of antibiotics in medicine, agriculture, and aquaculture. The emergence and persistence of antibiotic resistant bacteria impacts the environment through wastewater treatment plants, farm and slaughterhouse runoffs, hospital effluents, manure applications, and aquaculture. Muniesa et al. (2013) proposed a model to explain the mobilization of antibiotic resistance genes by phages in the environment. These researchers proposed that phages harboring antibiotic resistance genes present in different environmental biomes, are mobilized to commensal bacteria of animal and human biomes. From commensal bacteria, antibiotic resistance genes are transferred to pathogens of the Enterobacteriaceae family, such as Salmonella and E. coli O157:H7. Under selective pressure exerted by antibiotics, antibiotic resistance genes are incorporated into MGEs and will continue their mobilization through the animal biomes and human biomes by horizontal gene transfer. Variations of antibiotic resistance genes incorporated on MGEs will also develop through point mutations (Muniesa et al., 2013). In addition to this model, antibiotic resistant foodborne pathogens can be transferred from livestock to humans through improperly cooked meat and cross-contamination (Centers for Disease Control and Prevention [CDC], 2002; Schneider et al., 2011). Irrigation water and manure application contaminate crops, which also leads to antibiotic resistant foodborne illnesses. Finally, antibiotic resistance genes return to the environment through urban sewage and wastewater effluents and the cycle begins again (Figure 2). If left unchecked, this cycle has the potential to greatly compromise the effectiveness of antibiotic therapeutics.

Evidence of phage-mediated transduction events has been investigated in different environmental niches harboring phages

![Figure 2](image-url)
and in different antibiotic resistant bacteria. For example, Colomer-Lluch et al. (2011) investigated whether two prominent β-lactamase resistance genes, blaTEM and blaCTX_M, were present in phages collected from urban sewage or in the Llobregat river, near Barcelona, Spain. The sewage plant in Barcelona services approximately 500,000 inhabitants, and the river contains human and animal runoffs. The authors detected $10^2$–$10^4$ gene copies of blaTEM in DNA of phages isolated from urban sewage. Phages from the river samples contained, on average, 10 less gene copies of blaTEM than phages isolated from sewage. Presence of blaCTX_M was similar to that of blaTEM; 1.5 to 3 log_{10} units were detected in phages isolated from urban sewage samples and less than 1 log_{10} unit was detected in phages from the river samples. The authors then evaluated the viability of these resistance genes by examining whether they could confer resistance to the bacteria when transferred into two E. coli recipients. They detected blaTEM in a greater percentage than blaCTX_M in both strains. The authors suggested that in the presence of susceptible recipient strains, these environmental conditions would facilitate the phage-mediated transduction of ESBLs in the environment. As further evidence, Martí et al. (2014) demonstrated that phages carrying antibiotic resistance genes persist longer in the environment than their bacterial hosts. These authors investigated the presence of ESBLs (blaTEM, blaCTX_M, and blalSHV) and fluoroquinolone resistance genes (qnrA, qnrB, and qnrS) from two wastewater treatment plants and two hospitals located in Catalonia, Spain. A decrease in antibiotic resistance genes was observed in bacterial DNA from wastewater treatment plant samples compared to hospitals samples, which is unsurprising since wastewater treatment decreases the microbial load. However, antibiotic resistance genes, qnrS and blalSHV were detected in the phage DNA of all samples providing evidence of the persistence of phages and their potential to act as vectors for horizontal transfer of antibiotic resistance genes in the environment.

A study in Spain quantified quinolone resistance genes, qnrA and qnrS, in DNA of phage particles isolated from raw urban wastewater, cattle and poultry slaughterhouse wastewater and from the Llobregat river near Barcelona, Spain (Colomer-Lluch et al., 2014). QnrA was most prevalent because it was identified in the phage DNA of 100% of urban waste and river samples and 71.4% of animal wastewater samples. To evaluate the influence of phage inducing factors on the dissemination of antibiotic resistance genes (i.e., qnrA, qnrS, blaTEM, and blaCTX) in the environment, mitomycin C, ciprofloxacin, EDTA, and sodium citrate were used to induce phages from wastewater samples. Surprisingly, these authors observed that mitomycin C and ciprofloxacin, two known inducers of temperate phages, had no effect on the number of phages carrying qnr. However, EDTA and sodium citrate demonstrated a significant increase in phages carrying qnrA, qnrS, blaTEM, and blaCTX (Colomer-Lluch et al., 2014). This is probably attributed to the chelating properties of EDTA and sodium citrate, which disrupt the outer bacterial membrane by chelating the magnesium present within the LPS and cause induction of the phages (Imanovic and Muniesa, 2012). Based on these results, Colomer-Lluch et al. (2014) hypothesized that these phage particles carrying antibiotic resistance genes are generalized transducing phages since they contain fragments of bacterial DNA and no phage DNA. Thus, the phage particles lack the necessary genes for phage growth and lysis making them unable to produce plaques. Furthermore, to further support their hypothesis, they argued that no increase in phages carrying qnr genes was observed when induced with mitomycin C and ciprofloxacin, which are generally effective against specialized transducing phages. Additionally, the authors argued that since they did not observe an increase in virulent phages during induction, this further supports the hypothesis that the mobilization of qnrA, qnrS, blaTEM, and blaCTX was due to generalized transducing phages. This study provides evidence that the presence of phage-encoded antibiotic resistance genes in slaughterhouse run offs and river samples can be hotspots for generalized transduction contributing to the emergence of new resistance strains.

Agricultural soils have also been investigated as a reservoir for phages, which carry antibiotic resistance genes, as well as the potential of such phages for horizontal gene transfer in the agricultural soil microbiome. Ross and Topp (2015), assessed the abundance of antibiotic resistance genes within the bacterial fraction and phage fraction of soil mixed with raw, digested, dewatered, and composted manure prior to application, in contrast to soil without the application of manure. Antibiotic resistance genes were more abundant in bacterial fractions of soil applied with raw, digested and dewatered manure compared to composted manure, which indicated the efficiency of this practice in reducing the presence of antibiotic resistance gene carrying bacteria. However, the abundance of antibiotic resistance genes remained steady in the phage fraction of all manure applications. In addition, the reservoir of antibiotic resistance genes in the phage fraction remained constant with the application of treated biosolids, defined as nutrient rich organic materials produced from treated sewage sludge and used as a fertilizer in agriculture. Taken together, these results suggested that soilborne phages are a reservoir of antibiotic resistance genes. Moreover, the authors investigated the potential for transduction of antibiotic resistance genes by phages isolated from biosolids to soil coliforms, when selective pressure (1/10 and 1/100 of the breakpoint concentration of cefoxitin and sulfamethazine antibiotics, respectively), was applied. A 3.7- and 7.1-fold increase of cefoxitin-resistant coliforms and sulfamethazine-resistant coliforms, respectively, was observed (Table 2). This study provides evidence that regardless of the type of manure treatment, the soil is a reservoir of antibiotic resistance genes due to the presence of soilborne phages and antibiotics represent a form of selective pressure that promotes the horizontal transfer of antibiotic resistance genes by transduction (Ross and Topp, 2015).

**PHAGES MAY NOT PLAY A LARGE ROLE IN THE SPREAD OF ANTIMICROBIAL RESISTANCE GENES AMONG BACTERIA**

Reasonable evidence exists supporting the concept that phage-mediated transduction is a contributing factor to the
dissemination of antibiotic resistance in foodborne pathogens *E. coli* and *Salmonella* spp. Still, the role that phages play in transduction of antibiotic resistance genes remains controversial, as evidenced by two studies highlighted here. In the first study, Enault et al. (2017) employed several bioinformatic approaches to identify and assess the presence of antibiotic resistance genes within phage genomes, and found that antibiotic resistance gene prevalence in 1,181 phage genomes were vastly over-estimated due to low homologies with antibiotic resistance genes, as well as matches to proteins unrelated to antibiotic resistance. Furthermore, re-analysis of human- or mouse-associated viromes for antibiotic resistance genes suggested that the presence of such genes as attributable to phages in viromes were previously over-estimated. The authors concluded that their findings show that antibiotic resistance genes are rarely encoded in phages.

In the second study, Allen et al. (2011) directly studied prophage induction and horizontal gene transfer in animals. In this work, the authors used metagenomics to evaluate the effect of two antibiotics in feed (carbadox and ASP250 [chlorotetracycline, sulfamethazine, and penicillin]) on swine intestinal phage metagenomes (viromes). They also monitored the bacterial communities using 16S rRNA gene sequencing. It was observed that ASP250, but not carbadox, caused significant population shifts in both the phage and bacterial communities. Of importance to the topic of phage transduction, the authors observed that antibiotic resistance genes, such as multidrug resistance efflux pumps, were identified in the viromes, but in-feed antibiotics caused no significant changes in their abundance. The abundance of phage integrase-encoding genes was significantly increased in the viromes of medicated swine over that in the viromes of non-medicated swine, demonstrating the induction of prophages with antibiotic treatment. This suggests that while prophages were induced in the swine gut, this did not result in horizontal gene transfer of ARGs from the prophages to bacteria in the swine gut.

**CONCLUSION**

Notwithstanding these studies, and given the ubiquity, abundance and resilience of phages and the role of the environment as a reservoir, the contribution of transduction to the spread of antibiotic resistance must be considered and should be further explored in future studies. Furthermore, reasonable evidence supports the concept that phage-mediated transduction is a contributing factor to the dissemination of antibiotic resistance in *E. coli* O157:H7 and NTS. Controlling the spread of antibiotic resistance is an ongoing concern that must be addressed in order to ensure that a primary defense against infectious disease remains effective. Future studies should also address methods by which phage-based spread of antibiotic resistance genes would be mitigated. On this note, several studies have shown that phage-transferable CRISPR-Cas systems are capable of killing pathogens and resensitizing them to antibiotics (Bikard et al., 2014; Citorik et al., 2014). A recent study by Yosef et al. (2015) is noteworthy. In this work, the authors used phages to deliver a CRISPR-associated (Cas) to reverse antibiotic resistance and eliminate the transfer of resistance between strains. Instead of directly killing bacterial pathogens, as in traditional phage therapy, the bacteria were instead resensitized to the antibiotics that they are resistant to. While this work represented a proof of concept, as only bacterial isolates susceptible to the genetically modified phage would be resensitized, this study nevertheless demonstrates the potential for antibiotic resistance to be reversed through the use of novel approaches.

**AUTHOR CONTRIBUTIONS**

AC, BC, AL, and LG drafted and edited the manuscript. All authors revised the manuscript.

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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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