

Sequence Variability of P2-Like Prophage Genomes Carrying the Cytolethal Distending Toxin V Operon in *Escherichia coli* **O157**

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Cytolethal distending toxins (CDT) are potent cytotoxins of several Gram-negative pathogenic bacteria, including *Escherichia coli***, in which five types (CDT-I to CDT-V) have been identified so far. CDT-V is frequently associated with Shiga-toxigenic** *E. coli***(STEC), enterohemorrhagic** *E. coli***(EHEC) O157 strains, and strains not fitting any established pathotypes. In this study, we were the first to sequence and annotate a 31.2-kb-long, noninducible P2-like prophage carrying the** *cdt-V* **operon from an** *stx-* **and** *eae***-negative** *E. coli* **O157:H43 strain of bovine origin. The** *cdt-V***operon is integrated in the place of the** *tin* **and** *old* **phage immunity genes (termed the TO region) of the prophage, and the prophage itself is integrated into the bacterial chromosome between the housekeeping genes***cpxP* **and** *fieF***. The presence of P2-like genes (***n* **20) was investigated in a further five CDT-V-positive bovine** *E. coli* **O157 strains of various serotypes, three EHEC O157:NM strains, four strains expressing other variants of CDT, and eight CDT-negative strains. All but one CDT-V-positive atypical O157 strain uniformly carried all the investigated genomic regions of P2-like phages, while the EHEC O157 strains missed three regions and the CDT-V-negative strains carried only a few P2-like sequences. Our results suggest that P2-like phages play a role in the dissemination of***cdt-V* **between** *E. coli* **O157 strains and that after integration into the bacterial chromosome, they adapted to the respective hosts and became temperate.**

Cytolethal distending toxins (CDT) are considered prototypic inhibitory cyclomodulins [\(1,](#page-5-0) [2\)](#page-5-1). Genes encoding CDTs are widely disseminated among Gram-negative pathogenic bacteria, including *Escherichia coli*, *Campylobacter* spp., *Aggregatibacter actinomycetemcomitans*, *Haemophilus ducreyi*, *Salmonella enterica* serovar Typhimurium, and *Shigella* spp. [\(3\)](#page-5-2).

The holotoxin is a heterotrimer of three protein subunits, CdtA, CdtB, and CdtC. These are encoded by three adjacent, sometimes slightly overlapping genes [\(4\)](#page-5-3). CdtB is the active subunit, possessing DNase activity and sharing homology with the mammalian DNase I [\(5\)](#page-5-4). There is evidence that in case of *Campylobacter* spp., *A. actinomycetemcomitans*, and *H. ducreyi*, the other two subunits play a role in the transport of CdtB into the target cell [\(3\)](#page-5-2). Upon entering the eukaryotic cell, CdtB causes DNA damage, which in turn causes cell cycle arrest between the $G₂$ and M phases [\(6\)](#page-5-5). Double-stranded DNA damage leads to the distension and subsequent death of the target cell [\(7\)](#page-5-6). The characteristic distending transformation has been demonstrated on multiple cell lines by several studies [\(7,](#page-5-6) [8\)](#page-5-7).

CDT was first identified in *E. coli* (Johnson and Lior) [\(9\)](#page-5-8), and its production has been associated with several pathotypes, e.g., enterohemorrhagic (EHEC) and enteropathogenic (EPEC) [\(3\)](#page-5-2). So far, five types have been associated with *E. coli*, termed CDT-I to CDT-V [\(4,](#page-5-3) [6,](#page-5-5) [10,](#page-5-9) [11,](#page-5-10) [12\)](#page-5-11). The genomic localization of *cdt* alleles and their association with mobile genetic elements was reported by several groups. Accordingly, Pérès et al. localized *cdt-III* to a large conjugative virulence plasmid [\(6\)](#page-5-5). *cdt-I* and *cdt*-*IV* are encoded by lambdoid prophages [\(13,](#page-5-12) [14\)](#page-5-13), while the *cdt-V* operon is flanked by P2-like phage sequences [\(11,](#page-5-10) [14,](#page-5-13) [15,](#page-5-14) [16\)](#page-5-15), but the corresponding nucleotide sequences deposited show only small portions of the phage genes *repA* (replication gene A) and *Q* (a capsid gene).

The presence of CDT-V in Shiga-toxigenic *E. coli* (STEC) strains of various serotypes, both of clinical [\(17,](#page-6-0) [18\)](#page-6-1) and nonclinical [\(19\)](#page-6-2) origin, has been reported. CDT-V has also been associated

with strains of other serotypes and pathotypes associated with human diarrhea [\(16,](#page-5-15) [20\)](#page-6-3), where it is the only known cytotoxin of the respective strains, underlining the importance of CDT-V as a virulence factor. Therefore, it is imperative to obtain more information on the potential mobility of the CDT-V-encoding operon in non-STEC pathotypes and determine whether it is phage associated.

The aim of this study was to characterize the P2-like phage sequence context flanking the *cdt-V*operon in strain T22, an *E. coli* O157:H43 strain of atypical pathotype (*eae* and *stx* negative), and to monitor the presence of characteristic regions in other pathogenic and nonpathogenic CDT-V-positive strains and in additional K-12 *E. coli* strains.

MATERIALS AND METHODS

Bacterial strains. Strains used in this study are listed in [Table 1.](#page-0-0) Strains were grown on lysogeny broth (LB) agar plates or bromothymol blue agar plates.

Cosmid clone library construction and screening. Genomic DNA was isolated from strain T22 of *E. coli* O157:H43 with the Sigma genomic DNA kit (Sigma-Aldrich, St. Louis, MO). The preparation of the cosmid clone library was performed with the pWEB-TNC cosmid cloning kit (Epicentre, Madison, WI) according to the manufacturer's instructions, with the modification that genomic DNA was subjected to a partial digestion with restriction endonuclease MboI (Fermentas, Vilnius, Lithuania). All together, 1,000 transformant colonies were identified (cosmid library) and screened for the presence of *cdt-V* using the primers indicated in [Table 2.](#page-2-0)

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TABLE 1 Bacterial strains used in this study and the presenceof P2-like genes and regions in each of them

 regionand capsid packaging protein gene *Q*, which region spanning replication gene*repB* hypotheticalprotein. , presence; -, absence.

c EPEC, enteropathogenic *Escherichia coli*; EHEC, enterohemorrhagic *Escherichia coli*; ExPEC, extraintestinal pathogenic *Escherichia coli*.

d ND, not determined.

^a The reference sequence for the primers indicated with an asterisk is CP002967 (whole genome of *Escherichia coli* strain W). Positions are given in reference to GenBank accession number KC618326.1, except where indicated otherwise.

Sequencing. Cosmid DNA was isolated by using the alkaline lysis method [\(21\)](#page-6-11), genomic DNA was isolated by using the GenElute bacterial genomic DNA kit (Sigma-Aldrich, St. Louis, MO), and both were sequenced at the Biological Research Center (Szeged, Hungary) by using the Ion Torrent Personal Genome Machine (PGM) next-generation sequencer as well as traditional Sanger-based capillary sequencing. The average coverage for the prophage region was $112\times$. Trimming and assembly were performed manually and by using CLC Genomics Workbench version 6.0.1. Nucleotide sequence analysis and searches for open reading frames (ORFs) and homologous DNA sequences in the EMBL and Gen-Bank database libraries were performed with the tools available from the National Center for Biotechnology Information [\(www.ncbi.nlm.nih.gov\)](http://www.ncbi.nlm.nih.gov) together with Vector NTI and CLC Bio Genomics Workbench softwares.

PCR screening for flanking regions. *E. coli* strains representing different serotypes and pathotypes [\(Table 1\)](#page-0-0) were tested by PCR for the presence of characteristic P2-like phage genes. Primers were designed with the aid of PrimerBLAST, available from the National Center for Biotechnology Information. The general PCR conditions were an initial denaturation of 3 min at 94°C and then 30 cycles of denaturation for 30 s at 94°C, followed by annealing for 30 s at 59 to 60°C (depending on the primer pair) and extension for 60 s at 72°C. The final extension time was 5 min. The primers used in the reactions are listed in [Table 2.](#page-2-0)

Phage induction experiments. Phage induction was carried out for the 8 strains marked in [Table 1,](#page-0-0) using either mitomycin C (0.5 μ g ml⁻¹) or norfloxacin (1.25 μ g ml⁻¹) as inducing agents. Induction was also attempted by UV irradiation as described by Hertman and Luria [\(22\)](#page-6-12), with the modification that doses were 15, 10, and 5 s long with a 30-W lamp. *E. coli* K-12 strains ER2738, C600, and *Shigella sonnei* strain 866 [\(19\)](#page-6-2), kindly provided by Maite Muniesa, were used as indicator strains. As a positive control for phage induction, the *E. coli* O157:H7 strain 34, harboring an inducible Stx2 phage [\(23\)](#page-6-4), was used.

Nucleotide sequence accession number. The nucleotide sequence of the P2-like prophage has been deposited into GenBank [\(KC618326.1\)](http://www.ncbi.nlm.nih.gov/nuccore?term=KC618326.1).

RESULTS

Characteristics of the P2-like prophage carrying the *cdt-V* **cluster.**In order to determine the sequence of the *cdt-V* operon and its exact position in the genome of *E. coli* O157:H43 strain T22, a clone of the cosmid library positive for *cdt-V*, as well as the corresponding region of the genomic DNA, was sequenced, annotated, and deposited into GenBank. The length of the prophage containing the *cdt-V* operon is 31.2 kb. The *cdt-V* operon of strain T22 shows the highest nucleic acid similarity to the *cdt* operon of strain AH-10 (GenBank no. AB472839). The sequence of *cdtB*, coding for the active toxin subunit, is highly homologous to the AH-10 specific sequence, with only 4 synonymous single-nucleotide polymorphisms (SNPs). The CDT-V-encoding operon in strain T22 is flanked by P2-like prophage sequences. The majority of P2-like genes carried by T22 have the highest similarity to the bacteriophage L-413C (97% similarity on average at the nucleotide level; GenBank no. AY251033). The GC content of the *cdt-V* operon is 41%, whereas the prophage genes exhibit an average GC content of 53%. The structural organization of the P2-like phage genes is reminiscent of the P2-like prophage in *E. coli* strain W (GenBank no. CP002967). DNA sequence comparison between the P2-like sequences of strains T22 and W revealed that there is a high degree of nucleotide homology (94 to 99% identity) between their genes [\(Table 3\)](#page-4-0). The tail fiber-encoding genes (ORFs 33 to 36) located in the region ranging from bp positions 20720 to 23571 are different from all published P2-like prophages. The respective phage tail fiber genes of the prophage in the genome of *E. coli* strain UMNK88 (accession no. [CP002729;](http://www.ncbi.nlm.nih.gov/nuccore?term=CP002729) with the homologue of ORF 34 being present in two copies) are their closest homologues, with a nucleotide identity of 92 to 99%. The region between bp positions 22000 and 22192 is entirely missing in all other P2-like prophage sequences available in GenBank, whereas it is present in the prophage of UMNK88. The ORFs predicted in the P2-like prophage of *E. coli* O157 strain T22 are listed in [Table 3.](#page-4-0)

PCR scanning of P2-like genes. The dissemination of characteristic *cdt*-V-flanking genes ($n = 20$) in a collection of CDTproducing and nonproducing strains was examined by PCR. At least one gene from each P2-like functional gene cluster was investigated. The list of results, together with the genotype of the strains, is given in [Table 1.](#page-0-0) Fifteen out of 20 primer pairs designed specifically for the flanking regions [\(Table 2\)](#page-2-0) yielded PCR products in all CDT-V-positive strains. The overlapping primer pairs covering the Q capsid gene and the spacer region between the Q gene and *cdtA* yielded only a PCR product in the O157:H43 strains tested, and the tail sheath monomer-encoding FI gene could be amplified only from strain T22 and the O157:NM CDT-V strains. Strains carrying different *cdt* types and the CDT-negative strains carried fewer P2-like genes. Interestingly, the CDT-negative atypical strain B47 and the CDT-IV-positive strain 28C were positive for 15 and 12 of the investigated sequences, respectively. Sequencing of the amplicons produced by the P2_PQ primer pair [\(Table 2\)](#page-2-0) showed that in this 376-bp region of the Q capsid gene, there was a maximum of 14 SNPs, with only 3 of them leading to amino acid changes.

Phage induction experiments from CDT-V-positive strains. In order to induce lysogenic bacteriophages and to isolate corresponding phage particles from the CDT-V-positive strains, phage induction experiments were carried out with 8 CDT-V-positive strains, marked in [Table 1,](#page-0-0) by using UV light as well as mitomycin C and norfloxacin as inducing agents. Among CDT-V-positive strains, only EHEC O157:NM strains 702/88 and 493/89 released phages upon induction with mitomycin C, which were able to infect and lyse *E. coli*strain C600. PCR investigations revealed that none of these phages carried *cdt-V* genes (data not shown). These data indicate that the P2-like prophages harboring the *cdt-V* operon cannot be induced from these *stx*-negative and *cdt-V*-positive *E. coli* O157 strains.

DISCUSSION

We sequenced for the first time a whole P2-like prophage from the genome of *E. coli* O157:H43 strain T22 containing a *cdt-V* operon. The *cdt-V* sequence in strain T22 is highly similar to that of the *cdt-V* operon in strain AH-10 [\(16\)](#page-5-15). Our results confirm and expand our knowledge based on previous reports that the *cdt-V* operon is flanked by P2-like phage sequences [\(12,](#page-5-11) [16\)](#page-5-15) and on a more recent finding of an inducible P2-like phage carrying the *cdt-V* operon [\(19\)](#page-6-2). The proximal P2-like regions in *E. coli* T22 are highly similar to the sequences published by the above-mentioned authors; however, in our study, the entire prophage genome was resolved and annotated. Among the lytic P2-like bacteriophages, phage L-413C (induced originally from a *Yersinia pestis*strain [\[24,](#page-6-13) [25\]](#page-6-14)) has the highest homology compared to the prophage sequence of strain T22 (GenBank no. AY251033). Among other available prophage sequences, the P2-like prophage of *E. coli* O157:H7 strain TW14359 (GenBank no. CP001368.1) is the closest homologue. Compared to all the P2-like prophage sequences available in GenBank, the level of homology between the individual structure genes and their closest homologues is between 94 and 100% [\(Table 3\)](#page-4-0). The Z/fun region in the original P2 phage, situated between the tail fiber gene *G* and the tail sheath gene *FI*, is known to host insertions of foreign genes [\(26\)](#page-6-15). In the case of strain T22, the P2-like prophage does not contain a gene homologous to the G gene. Instead, the tail fiber genes between bp positions 20720 and 23571 (ORFs 33 to 36; see [Table 3\)](#page-4-0) seem to have a different origin in this strain, as they show only partial homology to P2-like phages or prophages, their closest homologues being tail fiber genes of a prophage carried by *E. coli* strain UMNK88. Among the flanking ORFs, ORF 32 corresponds to gene *H* (upstream of *G*) and ORF 37 to gene *FI*; therefore, the insertion of ORFs 33 to 36 occurred essentially in the Z/fun region, with the additional deletion of gene *G*. Nilsson and coworkers suggested site-specific recombination as a possible mechanism for the insertion of foreign genes in this region [\(27\)](#page-6-16). The potential recombination of prophages in *E. coli* O157 was already reported [\(28\)](#page-6-17), and it was also suggested that the mosaic structure of prophages is the result of extensive exchange of genetic material among different bacteriophages and also their hosts [\(13\)](#page-5-12), like in the case of lambdoid prophages harboring the *cdt-I* and *cdt-IV* operons [\(13,](#page-5-12) [14\)](#page-5-13). The genome sequencing project of strain T22 [\(29\)](#page-6-18) confirmed that the integration site of the prophage is between the *cpxP* and *fieF* genes. This is the same site where P2-like prophages can be found in a further four *E. coli*strains according to publicly available GenBank entries (accession no. [CP002967.1,](http://www.ncbi.nlm.nih.gov/nuccore?term=CP002967.1) [CP002797.2,](http://www.ncbi.nlm.nih.gov/nuccore?term=CP002797.2) [CP001969.1,](http://www.ncbi.nlm.nih.gov/nuccore?term=CP001969.1) and [CP000970.1\)](http://www.ncbi.nlm.nih.gov/nuccore?term=CP000970.1).

The integration site of the *cdt-V* operon within the prophage, the TO region, is known to be an integration hot spot for foreign genes in P2-like phages [\(26\)](#page-6-15). This region is named after the phage immunity genes *tin* and *old*, encoded by the lytic phages in this region. It has been suggested that this site is a potential carrier of

TABLE 3 Functional annotation of the P2-like prophage in *E. coli* O157:H43 strain T22*^a*

^a Positions refer to GenBank accession number KC618326.1. Homologies of 100% are in bold.

advantageous genes for the host [\(26\)](#page-6-15). The carriage of a potent virulence-associated gene, like *cdt-V*, provides evidence for this notion. It also has to be mentioned that, so far, CDT-V is the only established virulence factor with its genes inserted into the TO region of P2-like prophages. In complete P2-like prophage sequences available in GenBank, the TO region is usually occupied by a gene coding for a hypothetical protein, or in the case of EHEC O157:H7 strain TW14359, the genes of a putative virulence factor, one of which shows partial homology to a eukaryotic serine esterase and a gene of *Bacillus amyloliquefaciens* [\(30\)](#page-6-19). Genes encoding reverse transcriptases have been also reported in the TO regions of P2-like prophages found in the genomes of ECOR strains [\(31\)](#page-6-20). The foreign origin of the *cdt-V* operon is also supported by the considerable difference between the GC content of the operon and the prophage (41% versus 53%).

The fact that no phages were detected from any of the induced

cdt-V-positive strains, neither by induction with UV light, mitomycin C, or norfloxacin, is in harmony with current knowledge that P2-like phages are classified as noninducible in *E. coli* [\(26\)](#page-6-15). This finding also suggests that the inducible prophages found in STEC strains [\(19\)](#page-6-2) may represent an earlier evolutionary stage of the *cdt-V-*carrying P2-like prophages in *E. coli* and that the phages, which we were unable to induce, could have become temperated by adapting to the host organism.

The P2-like prophage sequences seem to be characteristic for the CDT-V-positive strains, with few differences that can be attributed to the adaptation processes in the various hosts. The presence of the L-413C-like C gene in the *stx-* and *eae*-negative O157 CDT-V strains, relative to EHEC CDT-V strains, can be explained with the high variability of this gene, which therefore can be used as a marker to identify and distinguish different P2-like phage variants [\(26\)](#page-6-15). Our results suggest that while the *stx*-negative *cdt-V* O157:H43 strains uniformly contain a P2-like variant with an L-413C-like C gene, the P2-like prophages in the other strains may carry different variants of this gene, as they represent other evolutionary lineages of P2-like phages. It will be an important future task to evaluate whether different P2-like bacteriophage variants may be associated with individual *E. coli* clonal groups and/or with different virulence factors in their TO region. An interesting finding was the carriage of P2-like genes by strain 28C, an extraintestinal pathogenic *E. coli* strain, which harbors the *cdt-IV* operon flanked by lambda phage-like genes [\(14\)](#page-5-13). This observation may indicate that the P2-like genes could be located elsewhere in the *E. coli* 28C genome. Alternatively, recombination between a P2-like and a lambdoid phage cannot be excluded. Another example of P2-like phages serving as vehicles for foreign DNA can apparently be found in the CDT-negative atypical O157:NM strain B47, which also carries most of the P2-like prophage genes investigated and is where the integration site between *cpxP* and *fieF* also seems to be occupied. Strain B47 probably also carries a nearly complete P2-like prophage similar to other investigated O157 CDT-V strains, with as-yet-unknown genes inserted into its integration hot spots. On the other hand, the PCRs specific for the integration site of the P2-like prophage were negative in the case of the O157:NM EHEC strains, suggesting a different integration site for the prophage in these isolates. The atypical O157 strains harboring P2-like genes are phylogenetically distinct from those that do not carry them. While the CDT-negative O157 strains (T4, B20, and B54; all representing the O157:H12 serotype) belong to phylogenetic group A, the CDT-positive O157 strains (T22, T16, T34, T49, T50), as well as strain B47, belong to phylogenetic group B1 [\(Table 1\)](#page-0-0) [\(32\)](#page-6-21).

It has been proposed that the presence of P2-like phage sequences suggests the common acquisition of the *cdt-V* operon in O157:NM EHEC strains after the lineage has diverged from the O157:H7 strains [\(12\)](#page-5-11). While there is indeed a strong association between O157:NM strains and the presence of CDT-V [\(17,](#page-6-0) [19,](#page-6-2) [33\)](#page-6-22), several strains from other serotypes, both from healthy cattle [\(23,](#page-6-4) [33\)](#page-6-22) and from cases of human diarrhea [\(15,](#page-5-14) [16,](#page-5-15) [17,](#page-6-0) [34\)](#page-6-23), express CDT-V. These findings, together with the results of our study, indicate that while the *cdt-V*genes are rather conserved, the carrier P2-like phages became diverse during the evolution of their hosts, and this event in most cases may have resulted in loss of their mobility. These facts suggest a more diverse evolutionary and/or transductional history of the *cdt-V* operon and its carrying of P2 like prophages. The highly conserved *cdt-V* operon within more

variable and potentially inactivated bacteriophage genomes may result from selective pressure to maintain a functional *cdt* gene cluster and to stabilize this cargo determinant by inactivation of this bacteriophage genome. Further investigations of flanking regions and P2-like prophage sequences in CDT-V-positive strains are expected to help in clarifying the evolutionary background of the distribution of these variants.

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