Evidence for an Adaptation of a Phage-Derived Holin/Endolysin System to Toxin Transport in Clostridioides difficile

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The pathogenicity locus (PaLoc) of Clostridioides difficile usually comprises five genes (tcdR, tcdB, tcdE, tcdA, tcdC). While the proteins TcdA and TcdB represent the main toxins of this pathogen, TcdR and TcdC are involved in the regulation of their production. TcdE is a holin family protein, members of which are usually involved in the transport of cell wall-degrading enzymes (endolysins) for phage-induced lysis. In the past, TcdE has been shown to contribute to the release of TcdA and TcdB, but it is unclear whether it mediates a specific transport or rather a lysis of cells. TcdE of C. difficile strains analyzed so far can be produced in three isoforms that are initiated from distinct N-terminal ATG codons. When produced in Escherichia coli, we found that the longest TcdE isoform had a moderate effect on cell growth, whereas the shortest isoform strongly induced lysis. The effect of the longest isoform was inhibitory for cell lysis, implying a regulatory function of the N-terminal 24 residues. We analyzed the PaLoc sequence of 44 C. difficile isolates and found that four of these apparently encode only the short TcdE isoforms, and the most closely related holins from C. difficile phages only possess one of these initiation codons, indicating that an N-terminal extension of TcdE evolved in C. difficile.

All PaLoc sequences comprised also a conserved gene encoding a short fragment of an endolysin remnant of a phage holin/endolysin pair. We could produce this peptide, which we named TcdL, and demonstrated by bacterial two-hybrid analysis a self-interaction and an interaction with TcdB that might serve to mediate TcdE-dependent transport.

Keywords: Clostridioides difficile, toxins, holins, endolysins, protein transport

INTRODUCTION

Clostridioides difficile is an opportunistic human pathogen that causes antibiotic-associated diarrhea (Schäffler and Breitrück, 2018). Strains of C. difficile usually produce two large protein toxins, TcdA and TcdB, and hypervirulent strains in addition often produce a third toxin, CDT (Shen, 2012). TcdA and TcdB are encoded in a genomic region termed pathogenicity locus (PaLoc) together with the regulatory proteins TcdR and TcdC, and the holin family protein TcdE.
Holins are known to mediate phage lysis by transporting endolysins across the cytoplasmic membrane (Saier and Reddy, 2015). Already in 2001, it had been postulated that TcdE may function as a lytic protein that mediates toxin transport (Tan et al., 2001). Evidence for a direct involvement of TcdE in toxin transport has been obtained for strains that produce high toxin levels (Govind and Dupuy, 2012; Govind et al., 2015) whereas strains that produce low toxin levels release the toxins by TcdE-unrelated lysis that most likely involves a recently discovered peptidoglycan-degrading transglycosylase (Olling et al., 2012; Govind et al., 2015; Wydau-Dematteis et al., 2018). There is not much known about the holin mechanism. Phage holins have been demonstrated to permit the membrane passage of endolysins and to depolarize the cytoplasmic membrane (Catalão et al., 2013), but it is unknown how a holin can mediate the translocation of very large toxins such as TcdA (308 kDa) or TcdB (270 kDa) without non-specific cell lysis. However, there is clear evidence for a TcdE-mediated transport that does not involve cellular lysis (Mukherjee et al., 2002; Govind et al., 2015). C. difficile TcdE is known to self-interact (Govind and Dupuy, 2012) and has been shown to trigger lysis of C. difficile and Escherichia coli (Olling et al., 2012; Govind et al., 2015). In the most recent studies on TcdE, the presence of differential translational start codons came into the focus (Govind and Dupuy, 2012; Olling et al., 2012; Govind et al., 2015), as previous studies on lambda phage holins showed a regulatory function of the co-existence of holin isoforms with close-by but distinct translational starts (Bläsi et al., 1990; Chang et al., 1995; Barenboim et al., 1999). In this well-studied case, only the short isoform is lytic, whereas the isoform extended by two residues is not lytic and believed to serve as antiholin whose production regulates the timing of lysis (Barenboim et al., 1999). The situation with TcdE is complicated by the existence of a third translational start. The three isoforms, named here by their translational start TcdE-M1, TcdE-M25, and TcdE-M27, have been studied with contradictory results. While one study found that the longest construct induced cellular lysis in the absence of shorter isoforms whose production was abolished by mutation of the alternative start codons (Govind et al., 2015), another study found that the longest isoform is non-lytic and attributed the cellular integrity of the tested C. difficile strains to the production of this non-lytic TcdE-M1 (Olling et al., 2012). Moreover, the latter study showed that the short isoform was lytic, at that time disregarding the translational start at M25 and focusing the analyses on the two starts at M1 and M27. Although TcdE is required for toxin secretion in strains with high toxin production, it is unknown why this holin is conserved also in the other strains (Govind et al., 2015). TcdE could be shown to be able to transport endolysins (Govind and Dupuy, 2012) and nothing is known about recognitions or interactions that could mediate a toxin specificity.

Here we present data that suggest specific roles for the three translational starts, with TcdE-M27 serving as holin, Tcd-M25 as typical antiholin, and Tcd-M1 as additional non-lytic isoform. The use of M1 provides an N-terminal extension that is inhibitory for lysis even in the presence of abundant short isoforms. This extended N-terminus is not present in the phage holins from which TcdE originated. Genomic analyses indicate that a significant number of C. difficile strains still do not contain this extension, suggesting that it evolved during the radiation of the species, most likely to optimize a toxin-secretion related function. A remnant of an endolysin gene that previously has been believed not to be relevant for the PaLoc system is conserved with a strong ribosomal binding site. First experimental data are presented that support the idea that this fragment might mediate interactions with the endolysin-unrelated large toxin TcdB.

### Materials and Methods

#### Strains and Growth Conditions

Escherichia coli strain ER2566 (NEB, Ipswich, United States) was used for all fractionation studies, and E. coli XL1-Blue Mr’ Kan or Tet (Stratagene, La Jolla, CA, United States), or DH5α were used for cloning. Cells were grown aerobically in LB medium (1% tryptone, 0.5% yeast extract, 0.5% NaCl) at 37°C with the appropriate antibiotics (100 µg/ml ampicillin, 25 µg/ml chloramphenicol, 50 µg/ml kanamycin). 0.5 mM IPTG or 0.1% rhamnose were used to induce P_{lac}Z or P_{rhab} dependent protein production at indicated time points. E. coli strain BTH101 (Euromedex, Souffelweyersheim, France) was used for bacterial-2-hybrid (B2H) studies. For growth curves, 25-ml cultures were inoculated with OD_{600} of 0.1 and grown aerobically. The OD_{600} was determined in 30-min intervals. For induction of gene expression, 0.01% rhamnose was added at indicated time points. 1% glucose was added for repression of rhamnose-dependent gene expression.

#### Genetic Methods and Plasmids

The genes tcdE and tcdL were rhamnose-inducibly expressed from pBW22 based vectors (Wilms et al., 2001) and C-terminally fused to a hexahistidine-tag. The tcdE expression vectors pBW-tcdE-M1-H6, pBW-tcdE(1-142)-H6, pBW-tcdE(1-142)-H6, pBW-tcdE-M25-H6, and pBW-tcdE-M27-H6 were cloned via standard methods (see Table 1 for all primers). A cloned tcdE was used as PCR template, which was amplified from chromosomal DNA using the primers Spel-tcdE-F and BamHI-tcdE-R, and cloned into the corresponding sites of the vector pCR2.1 TOPO (Thermo Fisher Scientific, Waltham, MA, United States). To construct pBW-lepB-H6, the lepB gene of E. coli was amplified with chromosomal DNA as template, using the primers Ndel-lepB-F in combination with BamHI-lepB-R, and the fragment was cloned into the corresponding sites of pBW-tatA-H6 (Berthelmann et al., 2008). For construction of pBW-tcdl-strep, tcdL was amplified by PCR in a first step with overlapping primers tcdl-synth-Ndel-F and tcdl-synth_BamHI-R. The amplified product was used as template for a second step with the primers tcdl_Ndel-F and tcdl_BamHI-F. PCR products were cut with Ndel/BamHI and cloned in the corresponding sites of pBW-tatA-H6 (Berthelmann et al., 2008) or pBW-tatA-strep (Mehner et al., 2012), thereby exchanging the fragments. For all these PCR amplifications, Plu polymerase (Promega, Mannheim, Germany) was used according to the suppliers protocol.
Single amino acid exchanges in TcdE were introduced by QuikChange mutagenesis (Stratagene) of pHIS1522-tcdB in conjunction with reverse primers that cover the identical sequence region. To correct the sequence to TcdE of C. difficile (Wohlan et al., 2014) was used to introduce a silent mutation corresponding sites of pU-strep-

**TABLE 1 | Primers used in this study**.

| Primer used in this study | Sequence |
|---------------------------|----------|
| SpeI-tcdE-F | ACTAGTATGACAGTAGTTCACC |
| BamHI-tcdE-R | GGAATCCTTTTACCTTGGTACATTC |

**Cloning of genomic tcdE**

Cloning of tcdE and its variants

| tcdE-M1-NdeI-F | ATATTGATATCATGAGAAAAACAGTTAGTATTAAATCATATAATTTCAAGAGCAAGAGTGGATCATATTTTGATAGTATTC |
| tcdE-BamHI-R | ATATTGATATCATGAGAAAAACAGTTAGTATTAAATCATATAATTTCAAGAGCAAGAGTGGATCATATTTTGATAGTATTC |
| tcdE-S64-BamHI-R | ATATTGATATCATGAGAAAAACAGTTAGTATTAAATCATATAATTTCAAGAGCAAGAGTGGATCATATTTTGATAGTATTC |
| tcdE-1104-BamHI-R | ATATTGATATCATGAGAAAAACAGTTAGTATTAAATCATATAATTTCAAGAGCAAGAGTGGATCATATTTTGATAGTATTC |
| tcdE-P142-BamHI-R | ATATTGATATCATGAGAAAAACAGTTAGTATTAAATCATATAATTTCAAGAGCAAGAGTGGATCATATTTTGATAGTATTC |
| tcdE-Met25-NdeI-F | ATATTGATATCATGAGAAAAACAGTTAGTATTAAATCATATAATTTCAAGAGCAAGAGTGGATCATATTTTGATAGTATTC |
| tcdE-Met27-NdeI-F | ATATTGATATCATGAGAAAAACAGTTAGTATTAAATCATATAATTTCAAGAGCAAGAGTGGATCATATTTTGATAGTATTC |

**Cloning of tcdL**

| tcdL-synth-NdeI-F | ATATTGATATCATGAGAAAAACAGTTAGTATTAAATCATATAATTTCAAGAGCAAGAGTGGATCATATTTTGATAGTATTC |
| tcdL-synth-BamHI-R | ATATTGATATCATGAGAAAAACAGTTAGTATTAAATCATATAATTTCAAGAGCAAGAGTGGATCATATTTTGATAGTATTC |
| tcdL-BamHI-F | ATATTGATATCATGAGAAAAACAGTTAGTATTAAATCATATAATTTCAAGAGCAAGAGTGGATCATATTTTGATAGTATTC |

**Cloning of lepB**

| NdeI-lepB-F | TATTACATATGGCGAATATGTTGCTGATCTTGG |
| BamHI-lepB-R | TATTACATATGGCGAATATGTTGCTGATCTTGG |

**Single amino acid exchanges in TcdL**

| tcdL-M25I-L-F | GCTCTGATAAAAAAGATATTGTGAGATTGAGAACGCCTCCTAGGTTTATATAAAAAATATTTCGTACCATTAG |
| tcdL-M25I-L-R | TATGCTCTGATAAAAAAGATATTGTGAGATTGAGAACGCCTCCTAGGTTTATATAAAAAATATTTCGTACCATTAG |
| tcdL-I1151-V-F | GATTAAAGGAAAAATAGAGTTTTACACTAGTACAGATACAG |
| tcdL-I1151-V-R | GATTAAAGGAAAAATAGAGTTTTACACTAGTACAGATACAG |

**Bacterial-2-hybrid screen**

| 2H-tcdE-BamHI-F | TATATGGATCCATGAGAAAAACAGTTAGTATTAAATCATATAATTTCAAGAGCAAGAGTGGATCATATTTTGATAGTATTC |
| 2H-tcdE-BamHI-F | TATATGGATCCATGAGAAAAACAGTTAGTATTAAATCATATAATTTCAAGAGCAAGAGTGGATCATATTTTGATAGTATTC |
| 2H-tcdE-BamHI-F | TATATGGATCCATGAGAAAAACAGTTAGTATTAAATCATATAATTTCAAGAGCAAGAGTGGATCATATTTTGATAGTATTC |

1 Primer sequences are given in 5’-3’ direction. 2 Corresponding reverse primers covered the same sequence.

The results were confirmed by SDS-PAGE with 100 mM Tris pH 9.5, 100 mM NaCl, 50 mM MgCl2, and samples were prepared as described previously (Behrendt and Brüser, 2014). Solubilisation of membrane proteins was achieved by adding 1% (w/v) digitonin. For small proteins, Schägger gels (16% T, 6 M urea) were prepared as described elsewhere (Schägger, 2006). For detection of hexahistidine-tagged proteins, Western blots were developed employing specific mouse monoclonal His-tag antibodies (1:1 mixture of penta-His/tetra-His; Qiagen, Venlo, Netherlands) in combination with the goat polyclonal anti mouse IgG secondary antibodies (Roth, Karlsruhe, Germany) coupled to horseradish peroxidase for enhanced chemoluminescence (ECL). Strep-tagged proteins were detected using Strep-Tactin alkaline phosphatase conjugate (IBA, Göttingen, Germany). For this, Western blots were washed once with 100 mM Tris pH 9.5, 100 mM NaCl, 50 mM MgCl2, and sequencing. The plasmid pLysS was obtained from Promega.

**Biochemical Methods**

SDS–PAGE and subsequent Western blotting were carried out by standard procedures (Laemmli, 1970; Towbin et al., 1979). BN-PAGE was performed with 5 to 13.5% gradient gels, and samples were prepared as described previously (Behrendt and Brüser, 2014). Solubilisation of membrane proteins was achieved by adding 1% (w/v) digitonin. For small proteins, Schägger gels (16% T, 6 M urea) were prepared as described elsewhere (Schägger, 2006). For detection of hexahistidine-tagged proteins, Western blots were developed employing specific mouse monoclonal His-tag antibodies (1:1 mixture of penta-His/tetra-His; Qiagen, Venlo, Netherlands) in combination with the goat polyclonal anti mouse IgG secondary antibodies (Roth, Karlsruhe, Germany) coupled to horseradish peroxidase for enhanced chemoluminescence (ECL). Strep-tagged proteins were detected using Strep-Tactin alkaline phosphatase conjugate (IBA, Göttingen, Germany). For this, Western blots were washed once with 100 mM Tris pH 9.5, 100 mM NaCl, 50 mM MgCl2, and sequencing. The plasmid pLysS was obtained from Promega.

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incubated in the same buffer supplied with 0.22 M NaCl in DMF and 0.16 M BCIP in DMF for staining. Images of Western blots were acquired utilizing the Intas Advanced Imager (INTAS, Göttingen, Germany).

For subcellular fractionations, cells were aerobically grown at 37°C in the presence of the appropriate antibiotics. If protein production was rhamnose-dependent, rhamnose was added to a final concentration of 0.1 or 0.01% (v/v), and cultivation proceeded for 1 h. Cells were harvested via centrifugation at 4,500 × g for 10 min at 4°C. Cell densities of corresponding cultures were normalized prior to the centrifugation step. Cell pellets were suspended in 50 mM Tris HCl pH 8.0, 250 mM NaCl, and after adding DNase I and 1 mM PMSF, and cells were homogenized by ultrasonication or French press (two passages, 800 p.s.i.). Cell debris was separated by centrifugation at 14,000 × g for 10 min at 4°C. Fractions of cell debris, supernatant (crude extract) or membranes were analyzed by SDS–PAGE and Western blotting.

In the course of the B2H analyses in E. coli strain BTH101, LacZ activities were determined using the classic activity assay by Miller (1972). In brief, overnight cultures were diluted 1:25 and cells were grown aerobically in the presence 0.5 mM IPTG at 30°C for 4 h. Cells from 0.5 ml of culture were harvested and resuspended in 2 ml Z-buffer (60 mM NaPO₄, 0.4 mM NaH₂PO₄, 10 mM KCl, 1 mM MgSO₄), and the optical density at 600 nm was determined. Cells from 1 ml of this dilution were permeabilized by adding two drops chloroform and one drop 0.1% SDS. After vortexing for 10 s, the reaction was started at 28°C by addition of 0.2 ml ONPG (4 mg/ml). The reaction was terminated after appropriate incubation times by the addition of 0.5 ml of the 1 M Na₂CO₃, the cells were sedimented (13,000 rpm, 5 min, RT), and the absorption at 420 nm was recorded for calculation of the activity as described (Miller, 1972).

**Genomic Sequence Extraction in Between the Genes tcdB and tcdA**
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*Clostridioides difficile* genomes of 38 strains were sequenced using SMRT and Illumina technology, assembled and annotated as described previously (Riedel et al., 2015a,b, 2017; Dannheim et al., 2017). From these genomes and from six additional *C. difficile* reference genomes (M68, CF5, 2007855, B11, R20291, and M120) downloaded from GenBank⁴, the nucleotide sequence regions in between the two genes tcdB and tcdA were extracted (Supplementary Data) and analyzed as described.

**RESULTS**

**Evidence for a Suppression of Translational Initiation at M1**

We analyzed the three TcdE isoforms Tcd-M1, Tcd-M25, and Tcd-M27 in the heterologous *E. coli* system that has been previously demonstrated to be functional with respect to lysis induction (Olling et al., 2012). In addition, we generated TcdE-M1 variants in which the other two translational starts were abolished either by M25L/M27L exchanges (TcdE-M1M25/27L) or by mutation of the second Shine-Dalgarno sequence that serves to initiate translation from M25 and M27 on (TcdE-M1ₐRBS). We also generated a TcdE-M25M27L construct that can only initialize at M25 due to a M27L mutation (Figure 1A). To ensure comparable and regulated expression, we used an identical Shine-Dalgarno sequence and the rhamnose-inducible Prhab Promoter for all constructs. A similar approach was taken in one of the previous studies that used an isopropyl-β-D-thiogalactopyranoside inducible T7 promoter, but in that study the TcdE-M25 construct was not included and the strong T7 promoter might have masked differential effects of the constructs (Olling et al., 2012). We observed primarily a production of the short isoform(s) with the TcdE-M1 construct, and only little full-length TcdE-M1 was detected (Figure 1B). Importantly, the removal of the internal initiation starts at M25/M27 in the construct TcdE-M1M25/27L resulted in a marked decrease of TcdE abundance selectively for the shorter isoforms, indicating that M25 and M27 are predominantly used as translational starts, even in the presence of an M1 start codon in conjunction with a very good Shine-Dalgarno sequence. As the TcdE-M25, TcdE-M25M27L, and TcdE-M27 constructs were highly abundant when produced with the same Shine-Dalgarno sequence (Figure 1B, compare lanes 4,5,6 with lanes 2,3) it is obvious that the translational start at M1 is suppressed in TcdE-M1 as well as in TcdE-M1M25/27L constructs, and the translational start at M25/M27 is therefore not responsible for the suppression of translational start at M1. The lower abundance cannot be explained by degradation, as the amount of full-length TcdE initiated at M1 was only slightly reduced. Nevertheless, the formation of small amounts of TcdE with the size of the small isoforms especially in the TcdE-M1ₐRBS construct indicates some degradation to about the size of the M25/M27 isoforms, which therefore likely represent a protease-resistant core. Taken together, these initial data indicated that the N-terminus 24 residues are only made in small amounts and rather represent some kind of extension of the TcdE core that begins around M25/M27. This suppression of translation at M1 likely results from an mRNA secondary structure around the M1 initiation codon and its Shine-Dalgarno sequence, which is evidenced by the fact that the TcdE-M1ₐRBS construct results in significantly more “full-length” TcdE-M1 than the TcdE-M1M25/27L construct (compare M1 bands in Figure 1B, lanes 2 and 3) and this is supported by the predicted mRNA secondary structures in that region. We used the software RNAstructure (Reuter and Mathews, 2010) to predict the secondary structure formation in a 130 nucleotides region (from −24 to +106) around the first initiation site and found that, in full agreement with the hypothesis, the TcdE-M1 construct formed a quite stable secondary structure including the Shine-Dalgarno sequence, and the stability of secondary structures around that region decreased as a consequence of the exchanges in the TcdE-M1ₐRBS construct due to a competing hairpin (Supplementary Figure S1). This explains the observed increase of translation.

⁴https://www.ncbi.nlm.nih.gov/
initiation at M1 in the TcdE-M1<sub>ΔRBS</sub> relative to TcdE-M1 and TcdE-M1<sub>M25/27L</sub>.

**TcdE-M27 Is the Lytic Isoform, TcdE-M25 Is Hardly Active, and TcdE-M1 Inhibits Lysis**

We then examined the effect of the above described TcdE constructs on growth (Figure 2A). Notably, only TcdE-M25 and TcdE-M27 constructs that allowed a translational initiation at M27 resulted in immediate cell lysis. The construct TcdE-M25/27L showed a very low efficient lysis effect, which may be due to N-terminal degradation to TcdE-M27 over time, indicating that TcdE-M25 is much less lytic than TcdE-M27. Also TcdE-M1, TcdE-M1<sub>M25/27L</sub> and TcdE-M1<sub>ΔRBS</sub> had only weak and retarded effects. There was no lysis when expression from the rhamnose promoter was suppressed by glucose in the medium. Taken together, these data clearly show that initiation at M27 triggers cell lysis in the absence of endolysins, whereas initiation at M1 or M25 does not. Already low amounts of TcdE-M1 inhibit lysis in the presence of large amounts of the small TcdE isoforms (Figure 1), suggesting that the isoform TcdE-M1 is really inhibitory for TcdE-M27 mediated cell lysis. As lysis is not delayed in a strain producing TcdE-M25 together with TcdE-M27 (construct TcdE-M25), TcdE-M25 appears to be not as inhibitory as TcdE-M1 with respect to endolysin-independent TcdE-triggered lysis, but we cannot differentiate the abundances of TcdE-M25 and TcdE-M27 by these Western blots.

**TcdE-Dependent Lysis Can Be Triggered by the Transport of Endolysins and Requires the TcdE C-Terminus**

It has been already demonstrated that TcdE can cause cell lysis in the presence of Lambda endolysin (Govind and Dupuy, 2012; Monot et al., 2015). We thus analyzed the potential effect of an endolysin on TcdE-dependent cell lysis. We constitutively produced T7 lysozyme with pLysS, which is a vector compatible with the rhamnose-induced vector system we used for expression of the tcdE variant genes. Importantly, while TcdE-M25 and TcdE-M27 caused the expected lysis that had already been observed in the absence of endolysins, TcdE-M1 showed a much enhanced cell lysis, indicating that this construct was able to transport T7 lysozyme across the cytoplasmic membrane (Figure 2B). As negative control, we suppressed rhamnose-dependent gene expression by glucose in the medium. As TcdE-dependent transport of lambda endolysin R has been previously demonstrated (Govind and Dupuy, 2012), which is unrelated to T7 lysozyme in sequence, it can be concluded that TcdE is able to mediate the escape of a wide range of endolysins across the cytoplasmic membrane. As expected for a transport-mediating holin, we found by BN-PAGE analysis that TcdE formed large associations of multiple protomers (Figure 2C). Notably, the TcdE-M1 isoform was already able to form large associations, and the smaller isoforms clearly contributed to the formation of associations with masses higher than 500 kDa.

We then examined whether the C-terminal region of TcdE is important for the lytic effect and analyzed TcdE variants truncated behind position P142, I104, or S64 (numbering starting...
production of either TcdE-M1, TcdE-M1\textsubscript{M1ΔRBS}, TcdE-M1\textsubscript{M125/27L}, TcdE-M25, TcdE-M25\textsubscript{M125/27L}, and TcdE-M27. As negative control, glucose instead of rhamnose was added to the strain containing the TcdE-M1 construct. Growth was without (A) or with (B) constitutive production of LysS from pLysS. Induction was with 0.01% rhamnose at the indicated time point. Standard deviations were calculated from three independent cultures. (C) BN-PAGE analysis of TcdE associations formed with the TcdE-M1 construct (1) and with the TcdE-M1\textsubscript{M125/27L} construct (2), showing the presence of large TcdE associations using the E. coli strains ER2566 pBW-tdcE-M1-H6 and pBW-tdcE-M1\textsubscript{M125/27L}-H6, respectively. SDS–PAGE analysis of the TcdE isoforms, showing that the strain with the TcdE-M1 construct contains also highly abundant smaller isoforms, whereas the TcdE-M1\textsubscript{M125/27L} construct contains very low amounts of smaller isoforms. Western blot detection using His-tag antibodies and ECL reaction. Molecular weight markers (in kDa) are indicated on the left.

FIGURE 2 | TcdE-dependent lysis can be triggered by endolysins. Growth curves of E. coli strain ER2566 with recombinant rhamnose-induced production of either TcdE-M1, TcdE-M1\textsubscript{M1ΔRBS}, TcdE-M1\textsubscript{M125/27L}, TcdE-M25, TcdE-M25\textsubscript{M125/27L}, and TcdE-M27. As negative control, glucose instead of rhamnose was added to the strain containing the TcdE-M1 construct. Growth was with or without constitutive production of LysS from pLysS. Induction was with 0.01% rhamnose at the indicated time point. Standard deviations were calculated from three independent cultures. (C) BN-PAGE analysis of TcdE associations formed with the TcdE-M1 construct (1) and with the TcdE-M1\textsubscript{M125/27L} construct (2), showing the presence of large TcdE associations using the E. coli strains ER2566 pBW-tdcE-M1-H6 and pBW-tdcE-M1\textsubscript{M125/27L}-H6, respectively. SDS–PAGE analysis of the TcdE isoforms, showing that the strain with the TcdE-M1 construct contains also highly abundant smaller isoforms, whereas the TcdE-M1\textsubscript{M125/27L} construct contains very low amounts of smaller isoforms. Western blot detection using His-tag antibodies and ECL reaction. Molecular weight markers (in kDa) are indicated on the left.

The N-Terminal Region of TcdE Evolved Within the Clostridia

As initiation at M1 was low efficient, and as the stability of the M25 and M27 isoforms suggested that these shorter isoforms constitute some protease-resistant core, we examined the existence of the differential translational starts in the most closely related phage holins, which are the holins of phiCDHM19, CDMH1, and phiMMP04, all of which are of C. difficile. These phage holins are highly similar among each other (96% identity) and are 79% identical to TcdE-M25. Importantly, all three phage holins clearly lack the N-terminal extension found in TcdE-M1, and all of them are most likely produced from only one translational start, which corresponds to M25 in TcdE, as a Val\textsubscript{GTA} codon is in the phage holins at the position of the M27\textsubscript{ATG} codon (Figures 4A,B). However, we cannot exclude that this GUA codon can be used by C. difficile as rare translational start. All known natural non-AUG start codons differ from AUG only by single bases (Diaz de Arce et al., 2018). For E. coli recI it has been reported that a translational start mutated to GUA could be used with low efficiency, which to our knowledge is the only demonstrated case of a functional translational start at GUA (Haggerty and Lovett, 1997). Having clarified that the translational start at M1 in TcdE is not found in the most closely related phages, we addressed the question whether TcdE proteins lacking this extension exist in C. difficile strains. For that purpose, tcdE genes of 44 distinct C. difficile isolates were analyzed (Figure 4C). Interestingly, in four of these strains, an ATT codon was at the position of the commonly found M1 ATG start codon. ATT is not known as translational start in C. difficile, but in E. coli, ATT has been found as rare non-canonical translational start codon in three out of > 4200 protein-coding genes (ymcF, ynfQ, infC). It thus cannot be excluded that some translation can initiate at this codon also in C. difficile. The tcdE genes of all 44 strains possess the canonical M25 and M27 translational starts (see Supplementary Data). In 11 out of the 44 strains, the Shine-Dalgarno sequence was rather bad due an A/C mismatch, and among these were the four strains that encoded no M1 translational start (Figure 4C). Together, the sequence data indicate that an introduction and optimization of the M1 translational start codon was followed by an improvement of the corresponding Shine-Dalgarno sequence. Most importantly, an
FIGURE 3 | C-terminal truncation abolishes the lytic effect of TcdE. (A) Scheme of the analyzed constructs. The thick regions represent potential trans-membrane domains. (B) Western blot analysis of the constructs produced by the E. coli strains ER2566 pBW-tcdE-M1-H6, pBW-tcdE(1−64)-H6, pBW-tcdE(1−104)-H6, pBW-tcdE(1−142)-H6, and pBW-lepB-H6. Western blot detection using His-tag antibodies and ECL reaction. Molecular weight markers (in kDa) are indicated on the left. The unrelated membrane protein LepB was used as negative control. (C) Effect of the indicated constructs on growth. “Glc” refers to a culture in which rhamnose-dependent expression of the full-length tcdE construct (TcdE-M1) was suppressed by glucose in the medium. (D) The same experiment as in (C) but with strains producing constitutively LysS from pLysS. Note that only the full-length TcdE-M1 construct is capable of mediating lysis, indicating LysS transport into the periplasm.
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![Table](data:image/...)

**FIGURE 4** The N-terminal extension of TcdE evolved within clostridia. (A) Comparison of the N-terminal sequence of TcdE with that of the three most closely related TcdE homologs from *C. difficile* phages. Note that the M1 and M27 isoforms are lacking in the phages, and only M25 (red) is conserved in all cases. “*” indicates identical positions, “:” positions with highly similar exchanges, “.” positions with weakly similar exchanges; (B) DNA sequence of the translational start region in all three phage tcdE homologs. Note that the Val codon that corresponds to the M27 in TcdE is not the rare initiation codon GTG, but rather the valine codon GTA. (C) DNA sequence comparison that shows the lack of the standard M1 translational start for TcdE in four of the analyzed *C. difficile* strains, and a less efficient ribosomal binding site in these four cases and in further seven strains, with DSM 27640 as example. The corresponding sequence of strain 630 is given as example for the 33 analyzed strains that possess a canonical M1 translational start with a good Shine-Dalgarno sequence. The improved Shine-Dalgarno sequence is underlined. Bold letters indicate the variable position in the Shine-Dalgarno sequence and the position of the M1 start codon.

initiation at M1 evolved in *C. difficile*, and not in phages, and therefore the N-terminal 24 residues extension likely relates to the function that TcdE has in *C. difficile*.

**A Remnant of a Phage Endolysin, TcdL, May Mediate Toxin Interactions With the System**

TcdE is known to be a holin that originated from lytic phage holin/endolysin systems (Monot et al., 2015), and our above analyses have indicated the closest phage homologs of TcdE. We wondered, whether a remnant of the endolysin could play a role. Such a remnant has been identified (gene ID CD630_06620) and regarded as pseudogene (Monot et al., 2015). However, a fragment of the endolysin could in principle be produced that does not need to have hydrolytic activity if it serves to mediate toxin transport functions rather than cell lysis in *C. difficile*. The endolysin fragment is unrelated to the endolysins from the three phages that contain holins closely related to TcdE and corresponds to a fragment of an endolysin found in the *C. difficile* phage phiCD481-1. The original endolysin gene is heavily mutated in the second half, whereas there are still very high
FIGURE 5 | Analyses of TcdL. (A) SDS-PAGE/Western blot detection of recombinantly produced Strep-tagged TcdL in cell debris and crude extract from the E. coli strain ER2566 pBW-tcdL-strep using Strep-Tactin alkaline phosphatase conjugate and NBT/BCIP. (B) Bacterial-2-hybrid analysis of TcdL, TcdB, and TcdE interactions, using LacZ activity assays. Data were normalized to the negative control values. Adenylate cyclase domains T18 or T25 were fused N-terminally or C-terminally to the proteins as indicated below the diagram. Note that the N-terminal TcdL-fusion to T18 (= TcdL-18) interacted with the C-terminal TcdL-fusion to T25 (= 25-TcdL), and the N-terminal TcdB-fusion to T18 (= TcdB-18) interacted with the N-terminal TcdL-fusion to T25 (= TcdL-25). (C) Homology-modeled TcdL structure and the corresponding region in the trimeric prophage endolysin (PDB 1YB0). Homology modeling was done using Swiss-Model (Waterhouse et al., 2018), and the images were created using CHIMERA (Petterson et al., 2004).
sequence identities on DNA level in the first half (Supplementary Figure S2). A one basepair deletion has generated an early stop codon, and the CD630_06620 open reading frame starts with an ATG codon that is not in frame of the orginal endolysin. The sequence of TcdL is brought into the correct frame by a frameshift after five codons, which already argues for some potential functionality of this gene. TcdL is a peptide of 53 amino acids (6.3 kDa). A Shine-Dalgarno sequence is conserved in all 44 isolate sequences that we have analyzed. In fact, there was only one A>G point mutation in 13 out of 44 tcdl sequences, and this point mutation included the four strains that also had not the M1 translational start in TcdE. The Shine-Dalgarno sequence differs from the phage endolysin-encoding sequence at one position, which potentially improves ribosome binding (AAGG>GAGG, see Supplementary Figure S2).

To analyze whether the peptide can be stably produced, we cloned it into the rhamnose-inducible expression system and detected it by Western blot in subcellular fractions. The peptide was detectable but largely formed inclusion bodies (Figure 5A). We term this gene product TcdL, which relates to its lysis protein origin. We then examined potential interactions of TcdL with TcdL, TcdE, and TcdB by use of a bacterial two hybrid system that is based on the reconstitution of adenylate cyclase activity from enzyme fragments that are brought in close proximity by interactions of fused proteins (Karimova et al., 1998). Notably, TcdL interacted with itself and with TcdB (Figure 5B). We did not detect any TcdE interaction. The TcdB interaction for the first time related TcdL directly to the toxins and possibly toxin transport.

DISCUSSION

This study reveals two main aspects with relevance for the field: (1) TcdE-M25 and TcdE-M27 behave similar to the well-studied Lambda S holin that has two translational starts, the first resulting in an antiholin and the second resulting in the lytically active holin (Chang et al., 1995). In contrast to phage holins, an additional isoform with an N-terminal extension evolved in C. difficile which inhibits lysis in the absence of transported substrates, i.e., in the absence of an endolysin (Figure 4). Interestingly, the extended isoform did either not inhibit transport of LysS, or the presence of LysS abrogated the inhibition of the lytic effect of TcdE (Figure 2). The finding that the N-terminal extension evolved most likely in PaLoc-encoded TcdE suggests that it is somehow related to toxins and their secretion. (2) There exists a remnant of a phage endolysin, TcdL, that might play a role in toxin transport as it is definitively inactive in its hydrolytic function but still may mediate interactions. The small protein per se is not soluble, which is expected for a fragment of a globular protein. However, interaction studies support an interaction of TcdL with the toxin TcdB (Figure 5B). It therefore is possible that the conserved fragment mediates transport processes that holins usually carry out with endolysins. Based on the known structure of the closely related Bacillus anthracis prophage endolysin PlyL (PDB 1YB0), we modeled the TcdL structure and analyzed the position of the corresponding region in the trimeric endolysin (Figure 5C).

Interestingly, TcdL is exposed to one face of the trimer and likely mediates the interaction of the endolysin with the associated phage holin. As the catalytic endolysin domain is absent in TcdL, it can be speculated that the TcdL-interacting toxin substitutes this domain, which could be the basis for a holin interaction that results in transport. However, such aspects are difficult to address and need thorough further analyses, including cross-linking approaches and biochemical transport analyses that so far could not be established for any holin. Attempts to transport TcdB into the periplasm of E. coli failed so far but the direct detection of selective transport would be a key to clarify these points.

Interesting is the aspect of the differential initiation at M1, M25, or M27 in C. difficile in comparison to E. coli as heterologous host. Some strains are reported to produce exclusively TcdE-M1, although they have unaltered initiation sites at M25 and M27. This is physiologically relevant, as TcdE-M1 has been shown not to cause cell lysis in several tested strains (Olling et al., 2012), and our data confirm this. However, we also observed that TcdE-M1 can in principle mediate cell lysis in strains that produce endolysins, which can explain results obtained by another study (Govind et al., 2015). The fact that, in E. coli, translation initiation at M1 was strongly inhibited, and the observation that point mutations within the second Shine-Dalgarno sequence could partially release this inhibition (Figure 1B) argues for the involvement of RNA secondary structures. Such structures have been shown to be important for the differential initiation at the two translational starts of the lambda holin S (Chang et al., 1995), and similarly have been suggested to play a role for the initiation sites at M25 and M27 of TcdE (Govind et al., 2015). However, secondary structures have so far not been considered for the initiation of TcdE at M1. Importantly, in the natural system, we found that there exists a RBS-protecting hairpin at M1 that is predicted to be very stable, implying that without additional regulatory components there would not be significant initiation at this translational start site (Supplementary Figure S1). The fact that it has been observed that C. difficile initiates in several tested strains only at this M1 site suggests that most likely a sRNA is involved that promotes initiation at M1, thereby suppressing toxin-substrate-independent lysis. This fits to the finding that TcdE is not responsible for a lytic release of the toxins in strains that release only low amounts of toxins (Govind et al., 2015). In agreement with these and our data, it has been recently demonstrated that a TcdE-independent lytic toxin release exists as an alternative pathway in parallel to the TcdE-dependent non-lytic pathway (Wydau-Dematteis et al., 2018). The lytic pathway depends on a novel transglycosylase that is active on specific media, especially in the stationary growth phase (Wydau-Dematteis et al., 2018). In principle, TcdE could also contribute to lysis when initiation at M1 would be suppressed by the secondary structure, which would induce a switch to lysis. However, initiation at M1 appears to be the rule, and this could be achieved by a regulatory sRNA. A single potential regulatory sRNA has been identified in the PaLoc, which is encoded at position 785940–786181 of the C. difficile 630 reference genome.
(Soutourina et al., 2013). We tested potential influences of this sRNA on secondary structures around M1 using the bimolecular interaction analysis tool of RNAstructure (Reuter and Mathews, 2010) and found no evidence for an influence of this sRNA on the M1 hairpin, which could base-pair only at distant positions. It thus remains unknown why initiation at M1 is so efficient in several C. difficile strains (Olling et al., 2012), albeit a very stable hairpin can be formed that is expected to efficiently suppress initiation. We propose that additional factors must be involved in the regulated use of this translational start.

AUTHOR CONTRIBUTIONS

DM-B and CR carried out all experimental work except genomic sequencing. TR and JO carried out genomic analyses. IJ and RG provided non-toxic tcdB variants and tcdE. TB and DM-B designed the experiments. TB supervised the study and wrote the initial manuscript. All authors contributed to the final manuscript.

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