Cytolethal Distending Toxin: A Unique Variation on the AB Toxin Paradigm

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Some of the most potent toxins produced by plants and bacteria are members of a large family known as the AB toxins. AB toxins are generally characterized by a heterogenous complex consisting of two protein chains arranged in various monomeric or polymeric configurations. The newest class within this superfamily is the cytolethal distending toxin (Cdt). The Cdt is represented by a subfamily of toxins produced by a group of taxonomically distinct Gram negative bacteria. Members of this subfamily have a related AB-type chain or subunit configuration and properties distinctive to the AB paradigm. In this review, the unique structural and cytotoxic properties of the Cdt subfamily, target cell specificities, intoxication pathway, modes of action, and relationship to the AB toxin superfamily are compared and contrasted.

1. Introduction

Bacteria secrete a myriad of different types of proteins that exhibit a cell damaging or “cytotoxic” activity. Examples include but are not limited to membrane-damaging proteins such as RTX (repeats in toxin) and CDC (cholesterol-dependent cytolysins), cell surface interacting proteins such as ST (heat-stable enterotoxins), and superantigens, as well as other proteins that attack cytosolic activities. One of the recurring organizational themes among secreted bacterial protein toxins that target intracellular processes in mammalian cells is a complex composed of at least two heterogeneous polypeptide chains or subunits. Each chain makes a distinct contribution to cell intoxication or toxic activity. The superfamily of cytotoxins that exhibit this structural arrangement have been generally labeled as the AB toxins [1]. In this arrangement, the A chain or subunit typically functions as an enzyme that disrupts a specific cell process or pathway and the B chain or subunit, in monomeric or polymeric form, and promotes binding of the holotoxin to the target cell surface.

The newest member of the AB toxin superfamily, discovered by Johnson and Lior in 1987 [2], is the cytolethal distending toxin (Cdt). This toxin was named so because Chinese hamster ovary (CHO) cells became stretched or enlarged when exposed to cell-free filtrates of enteropathogenic Escherichia coli (EPEC) isolated from young children diagnosed with gastroenteritis. Nucleotide sequences of the E. coli cdt genes were first reported in 1994 by Pickett et al. [3] and Scott and Kaper [4] and revealed that the toxin was composed of three heterogeneous subunits (CdtA, CdtB, and CdtC). Publication of this sequence led to the rapid discovery of a family that represents a related subgroup of the AB toxins. The various Cdts, like some of the other AB toxins, are produced by bacteria that are associated with specific diseases.

The objective of this treatise is to review current knowledge of the Cdt family and discuss how this subgroup compares and contrasts the AB toxin paradigm. Contributions from my laboratory are derived from studies of the Cdt produced by the human oral pathogen Aggregatibacter actinomycescomitans (formerly Actinobacillus actinomycetemcomitans) [5].

2. AB-Type Toxins

2.1. General Structural Features. Members of the AB toxin superfamily are presented in Table 1. There are currently at least 17 confirmed members, and two additional potential
**Table 1: The superfamily of AB toxins.**

(a)

<table>
<thead>
<tr>
<th>Class/Group</th>
<th>Phylogeny</th>
<th>AB</th>
<th>Gram-negative</th>
<th>Bacteria (prokaryotes)</th>
<th>Gram-positive</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Common name</strong></td>
<td>ricin</td>
<td>abrin</td>
<td>abrin subfamily</td>
<td>viscoxin</td>
<td>diphtheria toxin (Dtx)</td>
</tr>
<tr>
<td><strong>Source</strong></td>
<td>Ricinus communis</td>
<td>Roary pea/Jequirity pea</td>
<td>Modeca digitata</td>
<td>Viscum album L</td>
<td>Corynebacterium diphtheriae</td>
</tr>
<tr>
<td><strong>Toxic chain or subunit</strong></td>
<td>N-glycoside hydrolase [type II ribosome inhibiting protein (RIP)]</td>
<td>NAD⁺-diphthamide ADP-ribosyltransferase</td>
<td>protease</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Cellular target</strong></td>
<td>28S rRNA</td>
<td>elongation factor-2 (EF2)</td>
<td>cleavage of SNARE protein SNAP-25</td>
<td>cleavage of SNARE protein synaptobrevin II</td>
<td></td>
</tr>
<tr>
<td><strong>Toxic activity</strong></td>
<td>inhibits protein synthesis by hydrolysis of the N-glycosidic bond of an amine residue in the sarcin-ricin loop of the 28S rRNA</td>
<td>inhibits protein synthesis by ribosylation of EF2</td>
<td>neurotoxicity due to prevention of release of the neurotransmitters acetylcholine and γ-aminobutyric acid</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Cell receptor</strong></td>
<td>ganglioside GM3/glycoprotein/glycolipid</td>
<td>heparin-binding EGF-like (HB-EGF) receptor</td>
<td>lipoprotein receptor-related proteins LRP 1 and LRP 1B</td>
<td>GT1b</td>
<td></td>
</tr>
</tbody>
</table>
**Class/Group**

<table>
<thead>
<tr>
<th>Phylogeny</th>
<th>A_B_2</th>
<th>A_B_5</th>
<th>A_B_7</th>
<th>A_B_2</th>
<th>A_B_7</th>
<th>A_B_7</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gram-negative</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Phylogeny</strong></td>
<td>Bacteria (prokaryotes)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Gram-positive</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Common name</strong></td>
<td>cytolethal distending toxin (Cdt)</td>
<td>cholera toxin (Ctx)</td>
<td>Shiga toxin (Stx)</td>
<td>subtilase cytotoxin</td>
<td>heat labile enterotoxin (LT-I and LT-II)</td>
<td>enterotoxin</td>
</tr>
<tr>
<td><strong>Source</strong></td>
<td>multiple genera</td>
<td><em>Vibrio cholerae</em></td>
<td><em>Shigella dysenteriae</em></td>
<td><em>Bordetella pertussis</em></td>
<td><em>Escherichia coli</em></td>
<td><em>Campylobacter jejuni</em></td>
</tr>
<tr>
<td><strong>Toxic chain or subunit</strong></td>
<td>mammalian type I</td>
<td>N\textsuperscript{A}pyridine nucleotide-binding protein (PTEN)</td>
<td>N\textsuperscript{A}pyridine nucleotide-binding protein (RIP)</td>
<td>N\textsuperscript{A}pyridine nucleotide-binding protein (G proteins)</td>
<td>N\textsuperscript{A}pyridine nucleotide-binding protein (G proteins)</td>
<td>N\textsuperscript{A}pyridine nucleotide-binding protein (G proteins)</td>
</tr>
<tr>
<td><strong>Toxic activity</strong></td>
<td>cell cycle arrest at G1 or G2/M</td>
<td>alters signal</td>
<td>alters signal</td>
<td>cell cycle arrest at G1 or G2/M</td>
<td>alters signal</td>
<td>altered signal</td>
</tr>
<tr>
<td><strong>Cellular target</strong></td>
<td>nuclear</td>
<td>N\textsuperscript{A}pyridine nucleotide-binding protein (G proteins)</td>
<td>N\textsuperscript{A}pyridine nucleotide-binding protein (G proteins)</td>
<td>N\textsuperscript{A}pyridine nucleotide-binding protein (G proteins)</td>
<td>N\textsuperscript{A}pyridine nucleotide-binding protein (G proteins)</td>
<td>N\textsuperscript{A}pyridine nucleotide-binding protein (G proteins)</td>
</tr>
<tr>
<td><strong>Toxic receptor</strong></td>
<td>β-D-galactopyranoside</td>
<td>ganglioside GM1</td>
<td>N-glycolylneuraminic acid terminal glycan</td>
<td>ANTXR1 (tumor endotethelial marker-8 [TEM8])</td>
<td>ANTXR2 (capillary morphogenesis protein 2 [CMG2])</td>
<td></td>
</tr>
</tbody>
</table>

\textsuperscript{a}See Table 2.

\textsuperscript{b}Toxic activities are dependent on the species-specific Cdt and target cell.

\textsuperscript{c}SNARE, soluble N-ethylmaleimide-sensitive factor attachment.
candidates (*Campylobacter jejuni* enterotoxin and exfoliatin B), which are distinguished by the specific arrangement of the A and B chains or subunits. For example, some AB toxins have a monomeric organization for both the A and B chains (AB class) while others contain one or two A chains and polymeric homo- or heterogeneous B chain complex (AB$_2^-$, AB$_3^-$, AB$_5^-$, and AB$_7^-$). Genes for all members of the AB class identified to date are found in either specific species of plants or Gram-positive or Gram-negative bacteria. Genes for the other classes (AB$_2^-$, AB$_7^-$, and AB$_2^-$B$_7^-$) are not found in either Gram-positive or Gram-negative bacterial species or, in some unusual cases, in the genome of bacteriophages [6] or on conjugative plasmids [7].

The A chains or toxic components of the holotoxin are represented by enzymes including an N-glycosylase, NAD$^+$-diphthamide ADP-ribosyltransferase, deoxyribonuclease/phosphatidylinositol-3,4,5-triphosphate 3-phosphatase (PTEN, cation-dependent metalloenzyme), protein kinase, and Zn$^{2+}$ metalloprotease. The most common terminal mode of action in susceptible cells is the disruption of protein biosynthesis. However, in a few examples, the A chain acts as a potent neurotoxin or genotoxin/cyclomodulin.

The B chain is required for the binding of the holotoxin to susceptible cells which is an essential step for internalization of the A chain. There is significantly greater heterogeneity in the composition and structure of the B chain most likely due to the fact that this component of the holotoxin has evolved to recognize a broad range of target cells. These differences are evident upon a comparison of the crystal structures of several representative examples of the AB superfamily (Figure 1). Ricin, the most well-known representative of the AB class [8], is very similar to the other AB toxins of plant origin. The A and B chains are created by the cleavage of a single polypeptide chain. Following cleavage, the B chain

**Figure 1:** Structure comparisons of representative AB toxins. Structures of the A and B chains are shown as ribbon backbone models generated with UCSF Chimera 1.8.1. The anthrax B chain is depicted as the assembled heptamer. Three to four A1 or A2 chains or combinations of both chains are assembled in the holotoxin. The boxed structure shows the B chain monomer. Protein data bank (PDB) files: 2AAI (ricin and viscumin), 1R4Q (Shigatoxin), 1J7N (LF), 1XFY (EF), 1TZN (PA), 2F2F (AaCdt), and 1SR4 (HdCdt).
becomes bound to the A chain (N-glycosylase) through a disulfide bridge \([9, 10]\) and functions as a lectin that binds to terminal \(\beta\)-D-galactopyranoside residues on the cell receptor. In a slight variation on the ricin structure, the viscinin holotoxin forms a dimer, through a noncovalent linkage, to create an active toxin \([11]\). The B subunit of Shiga toxin is a homopentamer that binds to the membrane glycolipid globotriaosylceramide (Gb3), a gal-\(\alpha\)(1-4)-gal-\(\beta\)(1-4)-glu trisaccharide linked to ceramide \([12–14]\). The B subunit of anthrax toxin is a heptamer of the polypeptide termed protective antigen (PA) which binds to the cell surface anthrax toxin receptors (ATR or ANTXR) \([15, 16]\). Either one or both A chains, lethal factor (LF), and edema factor (ED) become attached as a 3-4 chain unit to the PA complex.

2.2. Modes of Action. Various A chain enzyme activities of AB toxins, other than those of the Cdt, result in the inhibition of protein synthesis, apoptosis, neurotoxicity, and alteration of cell signaling pathways (Figure 2).

2.2.1. Inhibition of Protein Synthesis. The A chain of all of the plant AB toxins and Shiga toxin is an N-glycosylase that removes adenine from the 28S rRNA \([17]\). This depurination reaction completely inactivates the ribosome. Therefore,

**Figure 2:** Comparison of the mode of action of the A chain of selected AB toxins. The A chain of ricin and Shiga toxin is a ribosomal inhibiting protein (N-glycoside hydrolase) that carries out a depurination by removing an adenine from the 28S rRNA. The A chain of anthrax toxin is represented by two polypeptides known as the lethal factor (LF), a \(\text{Zn}^{2+}\)-dependent endoprotease, and edema factor (EF), a polypeptide that forms a \(\text{Ca}^{2+}\)- and calmodulin-dependent adenylate cyclase. The A chain of choler toxin is enzymatically cleaved to create a polypeptide (A1 chain) that ribosylates the guanosine nucleotide-binding protein \(G_\alpha\). The A chain of botulinum and tetanus toxins degrades specific soluble N-ethylmaleimide-sensitive factor attachment proteins (SNARE) which inhibits vesicle fusion and delivery of neurotransmitters such as acetylcholine. Additional details are provided in the text. MAPKK: mitogen-activated protein kinase kinases; CFTR: cystic fibrosis transmembrane conductance regulator.
these toxins are known as an RIP (ribosome inhibiting protein). Diphtheria, cholera and pertussis toxins, exotoxin A, and the heat-labile (LT) enterotoxins also disrupt protein synthesis. The A chain of these toxins is an ADP-ribosyltransferase. In diphtheria toxin and exotoxin A, the enzyme catalyzes the transfer of NAD\(^+\) (nicotinamide adenine dinucleotide) to the elongation factor-2 (EF2). EF2, which is required for translocation of the nascent peptides on the polyribosomes, is inactivated resulting in the inhibition of RNA translation. The ADP-ribosyltransferases of the other toxins catalyze the transfer of different substrates.

2.2.2. Edema and Apoptosis. Anthrax toxin contains two heterogeneous A chains, known as the edema factor (EF) and lethal factor (LF) [16]. The EF forms a Ca\(^{2+}\) - and calmodulin-dependent adenylate cyclase that significantly increases the level of cAMP in the intoxicated cell. Calmodulin is a Ca\(^{2+}\) -binding protein that modifies the interaction of this divalent cation with various other Ca\(^{2+}\) -binding proteins. This action alters the water homeostasis of the cell and destabilizes intracellular signaling pathways which leads to high fluid accumulation in the tissues causing swelling. Macrophages are particularly susceptible to the toxin resulting in impaired immune function. The LF is a Zn\(^{2+}\) -dependent endoprotease that cleaves the N-terminal portion of the mitogen-activated protein kinase kinases (MAPKK). This action alters signaling pathways that require the intact kinases and contributes to apoptosis.

2.2.3. Ribosylation of G Proteins. The A chain of cholera toxin is activated by a cleavage step. The enzymatically active A1 fragment ribosylates the guanosine nucleotide-binding protein G\(\alpha\), This reaction maintains the G protein in a GTP-bound form that activates adenylate cyclase. As a consequence, high levels of cAMP are produced, activating the cystic fibrosis transmembrane conductance regulator (CFTR), resulting in excessive efflux of ions such as Cl\(^-\) and water from the intoxicated cells [18]. The high rate and volume of water released from epithelial cells (enterocytes) cause severe diarrhea.

2.2.4. Neurotoxicity. The A or light chain of the botulinum toxin functions as a protease that degrades the SNARE (soluble N-ethylmaleimide-sensitive factor attachment) protein SNAP-25 (synaptosomal-associated protein 25) [19]. The SNARE complex is a large superfamily of proteins, including syntaxin, synaptobrevin, synaptogamin, and SNAP-25, required for normal vesicle fusion that controls exocytosis of cellular transport vesicles with the cell membrane. Vesicle fusion is the process by which neurotransmitters, such as acetylcholine, are released for the transmission of signals between neurons. Cleavage of the SNARE proteins disrupts vesicle fusion resulting in the inhibition of signal transmission leading to muscle paralysis. The activity of tetanus toxin is similar to that of botulinum toxin [20].

2.3. The Cdt Subfamily of AB Toxins. The Cdt is produced by a handful of facultative or microaerophilic Gram-negative bacteria that are key pathogens in diseases that involve the perturbation of a mucosal (enteritis, gastric ulcers, chancroid) or epithelial (periodontal diseases) layer (Table 2). According to convention, the species-specific Cdts are represented by an abbreviated genus and species prefix such as AaCdt, [21]. To date, eight species-specific Cdt gene loci have been confirmed. The EcCdt is represented by a subfamily in which the various members include the cdt genes located on the chromosome, on plasmids or in bacteriophages. Five types of the EcCdt have been identified based on nucleic acid sequence differences [22, 23]. In comparison to the other classes of AB toxins, the Cdt subgroup exhibits several unique properties:

(i) cdt genes are carried and expressed by multiple genera of Gram-negative bacteria;
(ii) cdt genes are most likely spread across genera and species by horizontal gene transfer;
(iii) the deduced amino acid sequences of all three subunits have phylogenetic relationships to eukaryotic proteins or polypeptides;
(iv) the Cdt complex contains two heterogeneous A subunits—CdtA and CdtC (equivalent to the B chain in other AB toxins);
(v) the CdtB subunit (equivalent to the A chain in other AB toxins) has the potential to exhibit multiple enzymatic activities and therefore affect different cell processes or pathways;
(vi) a major mode of action of the Cdt is that of a genotoxin since the toxin enters the nucleus of cells and damages the host DNA;
(vii) the Cdt indirectly affects regulation of the cell cycle thereby behaving as a cyclomodulin.

3. Properties of the Cytolethal Distending Toxin

3.1. Taxonomic Relationships among the Cdt. The cdt operon, composed of a promoter and the three constitutively expressed structural genes, is usually found on the bacterial chromosome. However, it is likely that the operon was disseminated among the various Cdt-producing species by horizontal gene transfer. Cdt genes have been found in the genome of a bacteriophage from the aphid symbiont Hamiltonella defensa [6]. This gram-negative bacterium is related to the Enterobacteriaceae. Other E. coli bacteriophages also carry the cdt genes [24, 25]. These findings support the possibility of cross-species spread by transduction or transposition. It is interesting that the cdt locus in some E. coli strains is flanked by lambda prophage genes [23] or is adjacent to a putative transposase gene as in Providencia alcalifaciens [26]. Cdt-like genes have also been identified on a virulence plasmid (pVir) isolated from a strain of E. coli obtained from a septicemic calf [7]. Taken together, these observations strongly support an extrachromosomal mechanism of gene dissemination across genus and species lines for the Cdt.
Deduced amino acid sequence comparisons indicated that the \textit{Aa}c\textit{d}t locus may reside in an extant genomic or pathogenicity island (GIY4-1) marked by a bacteriophage attachment (att) sequence and the integrase/resolvase gene (\textit{xerD}). This putative genomic island may have been passed in succession from \textit{Haemophilus somnus} to \textit{H. influenzae} to \textit{H. ducreyi} to \textit{A. actinomycetemcomitans} \cite{27}. More recently, the \textit{Aaad}t operon has been found on a genomic island (\textit{cdt-}\textit{island}) distinct from GIY4-1 \cite{28}. The 2.5 kb \textit{cdt} operon and 1.3 kb region immediately upstream are the only regions of sequence homology between GIY4-1 and \textit{cdt-}island. These findings suggest that the \textit{cdt} genes may have been acquired by \textit{A. actinomycetemcomitans} as a result of at least two unrelated recombinational events. The %G+C content of the \textit{Aaad}t gene sequences indicate that they are foreign to this bacterial species. However, the deduced amino acid sequences \cite{29} are over 90\% similar to those from \textit{H. ducreyi} \cite{30}. This relationship is expected since \textit{A. actinomycetemcomitans} and \textit{Haemophilus} sp. are closely related members of the taxonomic family \textit{Pasteurellaceae} \cite{28}. The distribution of the \textit{cdt} genes among taxonomically unrelated Gram-negative genera such as the \textit{Enterobacteriaceae} and \textit{Pasteurellaceae} fits a model of spread by horizontal gene transfer. It is curious that genes for the other classes of bacterial AB toxins do not appear to have crossed species lines. However, there are similarities among the A chains of toxins produced by several different bacterial genera. For example, an ADP-ribosyltransferase is present in toxins produced by some strains of \textit{E. coli}, \textit{Corynebacterium diphtheriae}, \textit{Pseudomonas aeruginosa}, \textit{Vibrio cholera}, \textit{Bordetella pertussis}, and \textit{Clostridium} sp. (see Table 1). However, the cytotoxic end-points are different.

In spite of the evidence for genomic plasticity, there is no obvious selective pressure on the \textit{Aa}c\textit{d}t-producing species to prevent shedding and maintain expression of the \textit{cdt} genes. Indeed, fresh clinical isolates of \textit{Aa}c\textit{d}t-producing strains do not appear to readily lose expression of the \textit{cdt} genes after extensive laboratory cultivation. In contrast, the shedding of nonessential genes is readily apparent as in the case of the \textit{tad} (tight adherence) locus of \textit{A. actinomycetemcomitans} \cite{31}. All clinical isolates of this bacterium start out with an adherent “rough” colony morphology or phenotype that is rapidly lost, resulting in a nonadherent “smooth” phenotype, upon repeated subculturing in the laboratory. It has been proposed that propharytic genomes, especially those of pathogens, remain relatively small and generally lack nonfunctional sequences even though they are continuously evolving by acquiring genes due to horizontal gene transfer \cite{32-35}. This contradiction can be explained by a concept known as deletional bias in which genetic change is skewed towards deletions rather than insertions. Interaction of a bacterium with a host leads to low genetic diversity and

### Table 2: The cytotoxic distending toxin subfamily of AB toxins.

<table>
<thead>
<tr>
<th>Source(^a)</th>
<th>Habitat</th>
<th>Associated affliction or disease(^b)</th>
<th>Cell receptor</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{Escherichia coli}(^c)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cdt-I (EPEC)</td>
<td>Lower intestine</td>
<td>Gastroenteritis</td>
<td>Terminal (\beta)-D-galactopyranoside glycoprotein/TMEM181</td>
</tr>
<tr>
<td>Cdt-II (EPEC)</td>
<td>Lower intestine</td>
<td>Gastroenteritis</td>
<td></td>
</tr>
<tr>
<td>Cdt-III (septicemia)</td>
<td>Blood</td>
<td>Septicemia</td>
<td></td>
</tr>
<tr>
<td>Cdt-IV (pVir)</td>
<td>Lower intestine</td>
<td>Gastroenteritis/urinary tract infection</td>
<td></td>
</tr>
<tr>
<td>Cdt-V (STEC)</td>
<td>Lower intestine</td>
<td>Gastroenteritis</td>
<td></td>
</tr>
<tr>
<td>\textit{Campylobacter} sp.</td>
<td>Lower intestine</td>
<td>Gastroenteritis</td>
<td>Unknown</td>
</tr>
<tr>
<td>\textit{Salmonella enterica} subsp. \textit{enterica serovar Typhi}</td>
<td>Lower intestine</td>
<td>Typhoid fever</td>
<td>Unknown</td>
</tr>
<tr>
<td>\textit{Shigella dysenteriae}/\textit{Shigella boydii}</td>
<td>Lower intestine</td>
<td>Dysentery</td>
<td>Unknown</td>
</tr>
<tr>
<td>\textit{Providencia} \textit{alcalifaciens}</td>
<td>Lower intestine</td>
<td>Gastroenteritis</td>
<td>Unknown</td>
</tr>
<tr>
<td>\textit{Helicobacter} sp.</td>
<td>Stomach/liver</td>
<td>Duodenal ulcers/stomach cancer/chronic hepatitis</td>
<td>Unknown</td>
</tr>
<tr>
<td>\textit{Haemophilus ducreyi}</td>
<td>Genitalia</td>
<td>Chancroid</td>
<td>Unknown</td>
</tr>
<tr>
<td>\textit{Aggregatibacter actinomycetemcomitans}</td>
<td>Oral cavity</td>
<td>Localized aggressive and possibly chronic periodontitis/infectious endocarditis</td>
<td>GM3 (also possibly GM1 and GM2)</td>
</tr>
<tr>
<td>bacteriophage</td>
<td>\textit{Hamiltonella defensa}</td>
<td>Aphid symbiont</td>
<td>Not applicable</td>
</tr>
</tbody>
</table>

\(^a\)All facultative or microaerophilic Gram-negative species.
\(^b\)All diseases involving a mucosal or epithelial layer.
\(^c\)EPEC: enteropathogenic \textit{E. coli}; STEC: Shiga toxigenic \textit{E. coli}; pVir: conjugative \textit{E. coli} plasmid.
3.2. Regulation of the cdt Operon. The cdt genes are expressed from a polycistronic operon and are well conserved among the Cdt+ species [38–40]. The operon typically consists of a promoter and the three structural genes CdtA, CdtB, and CdtC. It has been reported that in some strains of A. actinomycetemcomitans two small open reading frames, orf1 (263 bp) and orf2 (258 bp) immediately upstream of cdtA, are cotranscribed in vitro with the three structural genes [41]. It does not appear that expression of orf1 and orf2 is essential for Cdt functions but may possibly have regulatory roles in transcription. In a rare finding, introns or intervening sequences have been located in the AacdC transcript [42]. Introns, when present in prokaryotic cells, are usually present in genes that function in the transfer of conjugative transposons or recombination. Their presence in the cdt transcript may be a phylogenetic holdover from its relationship to the plant (eukaryotic) members of the AB family and may represent a mechanism for transcriptional control. Aggregatibacter actinomycetemcomitans is a facultative bacterium and it has been reported that cdtB gene expression was upregulated 2-fold in stationary phase during anaerobic growth [43]. Bacteria such as A. actinomycetemcomitans associated with periodontal diseases colonize sites of relatively low oxygen tension in the human oral cavity [44, 45].

3.3. Holotoxin Structure. Biologically active Cdt is a trimer having the AB2 configuration (Figure 3). This holotoxin configuration is consistent in all Cdt-producing species examined to date except for that in Salmonella enterica serovar Typhi [21] which lacks the cdtA and cdtC genes [46]. Deduced amino acid sequence comparisons among representative species-specific Cdts reveal as much as 45% sequence identity between the most distantly related CdtB subunits [25, 47, 48]. CdtA sequence identities range between 93 and 26% and CdtC sequence identities range between 95 and 19%. In spite of this diversity, there appears to be a commonality of function among the species-specific Cdt subunits. Similar function in the case of significant sequence diversity is not uncommon based on studies of “highly divergent proteins.” A good example is triosephosphate isomerase (TIM) which functions in glycolysis and is therefore found throughout the universal phylogenetic tree [49]. TIM amino acid sequences are highly divergent. For example, the E. coli and Pseudomonas aeruginosa TIMs differ in 120 of 255 residues. In spite of this high divergence, TIM structures are very well conserved, being nearly identical, as illustrated by superimposing the structures of the E. coli and chicken TIM proteins [50]. It is not surprising that the Cdt proteins are highly divergent since they most likely have been disseminated among species by horizontal gene transfer which, as a consequence, results in a relatively high frequency of genetic change.

3.3.1. B Chain Activity. The “B chain” of the Cdt is a heterodimer composed of the products of the cdtA and cdtC genes. The mature CdtA and CdtC proteins are made up of, on average, 220 and 165 amino acids, respectively. Both CdtA and CdtC contain four cysteines that have the potential to
form two intramolecular disulfide bridges [48, 51]. There is no evidence of disulfide bond formation between the subunits based on the location of the cysteine residues in the CdtA and CdtC structures (see Figure 3).

It was predicted from the crystal structure of the HdCdt that CdtA and CdtC form a ricin-like lectin binding domain composed of a deep groove, between the two subunits, and a region of aromatic and heterocyclic amino acids exposed on the surface of CdtA [52]. The same features are evident in the crystal structure of the AaCdt [53]. However, it is predicted that the AaCdt holotoxin, unlike the HdCdt, forms a dimer due to differences in the amino acid composition of a nonconserved loop in CdtB. There is functional similarity between a sequence of 60–88 amino acids in CdtA and the α nonconserved loop in CdtB. There is functional similarity between a sequence of 60–88 amino acids in CdtA and the B chain of the ricin/abrin subfamily based on amino acid sequence threading analysis [54] and a reverse position specific- (RPS-) BLAST search of the conserved domain (CD) database [39]. The ricin B chain differs from CdtA such that it was formed by a series of gene duplications that created two sugar-binding domains. Each domain is made up of three copies of a 40 residue galactose-binding peptide [10]. An interesting parallel relationship between the binding properties of the ricin B chain and CdtA is evident from experiments performed using the EcCdt. It was observed that the binding of CdtA and CdtC to the surface of HeLa cells could be blocked with glycoproteins that specifically contain N-linked fucose moieties [55]. My laboratory took advantage of this in vitro binding specificity to use a modified enzyme-linked immunosorbent assay (ELISA), with thryoglobulin-coated plastic plates to study the binding kinetics of wild-type and mutated AaCdtA and AaCdtC subunits [56]. We found that CdtA alone bound with saturation kinetics to thyroglobulin. CdtC also bound to thyroglobulin, but at a much lower ratio of ligand to receptor. Using a series of glycan-deficient CHO cell mutants [57], Eshraghi et al. [58] were unable to confirm a role for N-linked glycans and glycolipids in the binding of either EcCdt or AaCdt. However, binding studies employing the thyroglobulin assay to examine the effect of single amino acid substitutions in CdtA empirically confirmed that surface-exposed heterocyclic amino acids in the "aromatic patch" region and on the CdtA face of the heterodimer groove [52] were essential for Cdt binding [59]. Li et al. [60] subsequently found that a heterocyclic amino acid (W115) that resides outside of the aromatic patch is also critical for receptor binding.

We found that CdtC binds poorly to CHO cells and does not enhance the binding of CdtA [56, 59]. These results suggested to us that the CdtA subunit may have a significantly greater role than CdtC in toxin binding in vivo. The results of binding studies based on immunofluorescence [39, 55, 61] and an enzyme-linked immunosorbent assay on live cells (CELISA) [56, 62] reinforced our hypothesis. Other studies have shown that CdtC can bind to HeLa and HEp-2 cells in culture and in a CELISA [55, 62, 63]. The presence of an obvious groove in the CdtA-CdtC heterodimer structure and some amino acid sequence similarity between regions in the two subunits [59, 62] suggests, but does not confirm, that the two subunits may work cooperatively such that CdtC facilitates binding of CdtA.

In contrast to the glycoprotein data, the individual AaCdt subunits were tested for binding to gangliosides Galβ(1,3)GalNAcβ(1,4)NeuAca(2,3)-Galβ(1,4)Glc-ceramide (GM1), GalNAcβ(1,4)NeuAca(2,3)-Galβ(1,4)Glc-ceramide (GM2), NeuAca(2,3)Galβ(1,4)Glc-ceramide (GM3), Galα(1-4)Galβ(1-4)Glc-ceramide (Gb3), and GalNAc-β(1-3)Gal-α(1-3)Gal-β(1-4)Glc-ceramide (Gb4) [64]. AaCdtA and AaCdtC bound primarily to GM1 and GM2. However, GM3 exhibited the greatest inhibition of Cdt-induced toxicity in the human monocye cell line U937. Gangliosides such as GM1, GM3, and Gb3 have been implicated as receptors for other AB toxins such as ricin [65], LT-I/LT-II [66], cholera toxin [67], and Shiga toxin [68]. Interestingly, GM3 is a component of membrane lipid rafts [69]. Lipid rafts are membrane domains enriched in glycosphingolipids and cholesterol and have been implicated as the binding site in AaCdt-mediated cell cycle arrest of Jurkat cells, a T-cell leukemic cell line [70, 71]. Those studies implied that CdtC-mediated binding to lipid rafts, via a cholesterol recognition/interaction amino acid consensus (CRAC) sequence (LIDYKGK) within the protein, is important for Cdt uptake.

The G-protein-coupled receptor, TMEM181, has been identified as a third potential Cdt cell receptor [72]. This membrane protein was implicated in the binding of the EcCdt using a novel haploid genetic screening method in a derivative of a chronic myeloid leukemia cell line.

The identity of the Cdt receptor(s) has been a challenging problem. Based on the structural analyses and in vitro binding kinetics, it seems apparent that a CdtA-CdtC heterodimer carries out the B chain function of other classes of AB toxins. However, more information is needed to establish the precise role of each subunit in the attachment of the Cdt to susceptible cells. One implication is that evidence is accumulating to support the idea that the species-specific Cdts, or subgroups such as Aggregatibacter/Haemophilus (Pasteurellaceae subgroup) and Escherichia/Salmonella/Campylobacter/Helicobacter (GI subgroup), may have distinct host cell receptors or that multiple receptors exist in specific cells or cell lines. Based on intoxication experiments using wild-type CHO cells treated with tunicamycin or loaded with cholesterol and CHO cell mutants containing a sialic acid galactose transporter or glycolipid deficiency, the species-specific Cdts separated into three subgroups, AaCdt/HdCdt, EcCdt, and CjCdt, requiring different host cell properties [58]. As noted by this group of investigators, the precise molecular conformation and presentation of the Cdt receptor on the cell surface may be essential to obtain Cdt binding. The implications of these observations mean that binding to a native receptor may only be observable with intact cells since purification of the receptor may significantly alter the conformation required for correct interaction with the toxin.

3.3.2. A Chain Activity. The biologically active or toxic component of the Cdt, comparable to the A chain of the other classes of AB toxins, is also an enzyme. Based on a position-specific iterated (PSI) BLAST analysis [73], EcCdtB and CjCdtB matched to type I deoxyribonucleases
However, a broader deduced amino acid sequence comparison showed that CdtB belongs to a superfamily of enzymes that includes the endonucleases, exonucleases, spheromylinases, and inositol polyphosphate 5-phosphatases [76]. The phylogenetic relationships imply that CdtB appears to be most closely related to prokaryotic and bacteriophage deoxyribonuclease I and sphingomylinase C with broader similarities to PTEN and eukaryotic deoxyribonuclease I. Therefore, CdtB has the potential to exhibit multiple enzymatic activities in vitro and in vivo.

1. Deoxyribonuclease-Like Activity. Recombinant CdtB exhibits nicking or relaxation activities, in vitro, when incubated with supercoiled DNA [74]. DNA damage in the form of double-strand breaks, as assessed by pulsed-field gel electrophoresis (PFGE), has also been observed in sensitive cell types exposed to the heterotrimer [61, 77, 78]. CdtB requires divalent cations for DNA-nicking activity [74, 79] with an optimum concentration of 50 mM MgCl₂, CaCl₂, or MnCl₂ for the AaCdtB [80]. Mutating the conserved deoxyribonuclease active site residues His160 and His274 in the HidCdt [48], or their equivalent His residues in the other species-specific CdtBs, results in the loss of both in vitro nuclease activity and cell cycle arrest [74, 75]. Chimeric proteins composed of the amino terminal half of the AaCdt and the carboxy terminal half of human deoxyribonuclease I (CdtB/DNaseI) exhibited DNA-nicking activity, in vitro, comparable to that of CdtB, and formed an active heterotrimer complex with CdtA and CdtC [80]. The reverse chimeric construct, DNaseI/CdtB, also retained DNA-nicking activity but failed to form an active heterotrimer. Therefore, it appears that a primary activity of CdtB is to act as a neutral, cation-dependent deoxyribonuclease I-like nuclease [61, 74]. Pulsed-field gel electrophoresis of HeLa and immortalized labial epithelial cells (GMSM-K) exposed to the AaCdt [61] and HeLa cells treated with the HidCdt [78] showed that the toxin causes DNA double-strand breaks in vivo. The induction of DNA double-strand breaks occurs by 6 hours after intoxication and leads to the formation of actin stress fibers by way of an ATM-dependent activation of the small GTPase, RhoA [81, 82]. This DNA-damaging activity is unusual for an AB toxin and supports the in vivo activity of the Cdt as a genotoxin [83]. The EcCdt induces DNA double-strand breaks primarily in the S phase of HeLa cells that are exposed to a low dosage of the toxin [84]. Single-strand breaks were converted to double-strand breaks during the S phase and the cells attempted to repair the damage by homologous recombination. These data indicate that the S phase may play a more important role in the toxicity of the Cdt in some cell types and have important implications for cells that are deficient in homologous recombination. The other members of the AB toxin superfamily do not target host cell DNA. One exception, Shiga toxin has been reported to damage single-stranded DNA [85]. However, the damage caused by this toxin is due to depurination rather than cleavage of phosphodiester linkages.

2. Consequences of DNA Damage. DNA damage results in the recruitment of protein complexes for DNA repair and leads to growth arrest at specific “checkpoints” in the cell cycle. When DNA damage is detected by cells, the goal is to prevent DNA replication and mitosis. The cell cycle checkpoints block progression from the G₁ to the S phase or from the G₂ to the M phase (Figure 4(a)). Cell cycle arrest at these checkpoints can be detected and quantified by flow cytometry using fluorescence-activated cell sorting (FACS). A population of cells having a 2n DNA content indicative of cells in the G₁ phase can be separated from that having a 4n DNA content. Thus, quantitative detection of the accumulated cells having an increased DNA content is indicative of G₂ arrest (Figure 4(b)).

During normal growth, the protein kinases Myt1 and Weel phosphorylate the effector kinase Cdc2/Cdk1 at positions Thr14 and Tyr15, respectively (Figure 3(c)). The phosphorylated Cdc2 forms a complex with cyclin B. This complex is required for the cells to enter mitosis and is activated by the removal of the phosphates from Thr14 and Tyr15 by the protein phosphatase Cdc25. Once the cells enter mitosis, the cyclin B is degraded and Cdc2 is recycled by dephosphorylation of Thr161.

The pathways for cell cycle arrest in response to the Cdt appear to be cell-type specific and have been presented in various reviews [86–91]. Cdt-induced DNA damage can trigger pathways leading to the inhibition of cell proliferation, cell death, detrimental changes in cell morphology or alterations in cell signaling. In human cells, when DNA damage occurs, the sensor proteins Mre11, Rad50, Rpa1-3, and Nbs1 (MRN complex) are recruited to the site [92, 93]. Depending on the cell-type various checkpoints, DNA repair and/or signaling pathways are initiated (Figure 4(d)). Arrest at the G₁ phase of the cell cycle has been reported in primary lung fibroblasts after exposure to the HidCdt [94, 95] and in the human fetal lung fibroblast cell line IMR-90 following treatment with the CjCdt [96]. The classical G₁ arrest pathway initiated by DNA damage occurs when a phosphorylated p53 (tumor suppressor protein) complex mediates the upregulation of p21. The p21 inactivates a Cdc2-cyclin E complex that prevents the cells from progressing to the S phase and also binds to PCNA (proliferating cell nuclear antigen) which results in an inhibition of DNA replication [97]. AaCdt-induced expression of p21 may drive G₂ arrest in B cells [98] and p53-dependent and independent pathways may regulate the transition from G₂ to M phase under genotoxic stress [99].

Human foreskin fibroblasts exposed to the HidCdt arrested at both the G₁ and G₂ phases [94]. Similar results were obtained with HPLF and HGF treated with a Cdt-containing extract from A. actinomycetemcomitans and recombinant HidCdt [100]. Indeed, it appears that the response of most Cdt-sensitive cell types is cell cycle arrest at the G₂/M interphase [7, 75, 91, 94, 95, 100–110]. As in the case of G₁ arrest, Cdt-induced DNA damage triggers a response by the sensor proteins leading to activation of the ATM (ataxia telangiectasia mutated) and ATR (ATM and Rad3-related) checkpoint pathways [111]. In these pathways, the effector kinases Chk1 and Chk2 phosphorylate Cdc25 at Ser216. This inactive form, phospho-Cdc25, fails to dephosphorylate the checkpoint protein kinase Cdc2 (Cdk1) that is complexed
Figure 4: CdtB-induced inhibitory pathways. (a) Schematic representation of stages in the cell cycle showing the locations of the G₁ and G₂ DNA damage checkpoints. (b) Cell cycle arrest at G₂/M can be measured by flow cytometry or cell sorting. Populations that are arrested at G₂/M accumulate cells that have a 4n DNA content. (c) Normal cell cycle pathway leading to mitosis. Protein kinases Myt1 and Wee1 phosphorylate the effector kinase Cdc2/Cdk1 which forms a complex with cyclin B. The complex is activated by the removal of phosphate by the protein phosphatase Cdc25 allowing the cells to enter mitosis. (d) Cdt-induced DNA damage resulting in arrest at either the G₁ or G₂ checkpoint. DNA damage sensor proteins are recruited to the repair site depending on the cell type either the ataxia telangiectasia mutated/ATM and Rad3-related (ATR/ATM) or p53 pathway is initiated. Arrest at the G₁ phase of the cell cycle occurs when the phosphorylated tumor suppressor protein p53 complex mediates the upregulation of p21 which inactivates a Cdc2-cyclin E complex and the proliferating cell nuclear antigen (PCNA). Cdt-induced DNA damage can also trigger a response by the DNA damage sensor proteins.
leading to activation of the ATM and ATR checkpoint pathways. In these pathways the effector kinases Chkl and Chk2 phosphorylate, and therefore inactivate Cdc25. Activation of the ATM pathway can also lead to the initiation of a survival pathway in which the ATM protein kinase dephosphorylates the neuroepithelioma transforming protein, Nett, which activates RhoA which in turn activates the RhoA activated kinases (ROCK). (e) Putative pathway by which CdtB may affect the phosphatidylinositol-3-kinase (PIK) signaling pathway and the production of proinflammatory or anti-inflammatory cytokines. CdtB may function as a phosphatidylinositol-3,4,5-triphosphate 3-phosphatase (PTEN) that removes a phosphate group from phosphatidylinositol-3,4,5-triphosphate (PIP3) present in the plasma membrane. This depletion of PIP3 in lymphocytes inactivates the Akt pathway leading to cell cycle arrest and apoptosis. In macrophages, PIK3 may be activated by toll-like receptors (TLRs) in the cell membrane resulting in the synthesis of PIP3, and the activation of Akt. This activator of anabolic signaling pathways acts on the glycogen synthase kinase-3 (GSK3β). GSK3β and phosphorylated GSK3β direct the pathway toward the synthesis of proinflammatory and anti-inflammatory cytokines, respectively. The mTOR (mammalian target of rapamycin) protein complex is also regulated by the Akt kinase and participates in the control of protein synthesis and cell growth. Additional details are provided in the text. CBP: CREB-binding protein; CREB: cAMP response element binding protein; TLR: toll-like receptor. The pathways were compiled and modified from those described in other studies [89, 90, 95, 119, 177, 212].

with cyclin B. The inhibition of Cdc2 dephosphorylation blocks progression of the cells to mitosis (Figure 4(d)). Toxins that modulate the eukaryotic cell cycle have been termed cyclomodulins [89, 112].

ATM induced Chk2 can also send the cells on an apoptotic pathway due to accumulation of phosphorylated p53 [113]. Guerra et al. [114] found that the transcription factor Myc contributes to the activation of the ATM-dependent induction of apoptosis. The data indicated that Myc stimulates ATM phosphorylation and supports expression of Nbs1 and nuclear translocation. Several groups have published evidence, based on the activation of caspases, mitochondrial damage, decreased cell size, and DNA fragmentation, that the HdcCdt and AucCdt induce apoptosis in lymphocytes in addition to the G2 arrest [107, 115, 116].

Activation of the ATM pathway in response to the HdcCdt in HeLa and HCT116 cells (colorectal carcinoma) can lead to the initiation of a survival pathway in addition to the Chk2-mediated G2/M arrest [81]. In this pathway, the ATM protein kinase dephosphorylates the neuroepithelioma transforming protein, Nett, a RhoA-specific guanine nucleotide exchange factor (GEF). This reaction leads to the activation of the GTPase, RhoA, that is associated with extended cell survival [78]. RhoA activated kinases (ROCK) are required for the induction of actin stress fibers [81, 82, 86]. Activation of a survival pathway by the Cdt implies that some chronic bacterial infections could support the malignant transformation of cells [81].

(3) Phosphatidylinositol-3,4,5-triphosphate 3-Phosphatase Activity. The fact that CdtB is a member of the large metalloenzyme superfamily led to the proposal that the toxin acted upon susceptible cells or cell lines by dephosphorylation of the Weel kinase or Cdc25 phosphatase [117]. These two enzymes are important for regulating the phosphorylation of tyrosine residues in Cdc2 kinase. An increase in Cdc2 phosphorylation leads to cell cycle arrest. This rationale led to the hypothesis that the primary mode of action of CdtB is that of a phosphatase rather than that of a nuclease. Additional support for the phosphatase mode of action came from studies showing that recombinant AucCdtB behaved as a PTEN in lymphoid cell lines [118]. This enzyme is a hydrolase that removes a phosphate group from phosphatidylinositol-3,4,5-triphosphate (PIP3) found in the plasma membrane of cells. PIP3 activates downstream signaling components such as the protein kinase Akt, an activator of anabolic signaling pathways necessary for cell growth. It was proposed that the action of the Cdt in lymphocytes causes a depletion of PIP3 which in turn inactivates the Akt pathway leading to cell cycle arrest and apoptosis. This same group expanded their examination of the PTEN activity of the AucCdt to include human macrophages [119]. Unlike lymphocytes, macrophages did not undergo Cdt-mediated apoptosis. The PIP3 signaling pathway has been implicated in the regulation of the cellular proinflammatory response. PIP3 is made from phosphatidylinositol-4,5-diphosphate (PIP2) via phosphorylation by the lipid kinase phosphatidylinositol-3-kinase (PIK3). In macrophages, PIK3 may be activated by toll-like receptors (TLRs) in the plasma membrane [120]. Inactivation of glycogen synthase kinase-3 (GSK3β) by phosphorylation regulates the transcription factors, CREB (cAMP response element binding protein) and NF-κB, via the transcription activator CREB-binding protein (CBP), thereby altering cytokine synthesis (Figure 4(e)). The mTOR (mammalian target of rapamycin) protein complex also belongs to the PIK3 pathway [121]. This protein complex is involved in the control of protein synthesis and cell growth and is regulated by the Akt kinase [119, 122].

(4) Cell Signaling Activities. It has been reported that the synthesis of interleukin (IL)-1β, IL-6, and IL-8 in human peripheral blood mononuclear cells (PBMC) is stimulated by the AucCdt [38] and IL-8 in human embryonic intestinal epithelial cells (INT407) is induced by the CjCdt [122]. In addition, nitric oxide production was altered in macrophage cultures by the presence of AucCdt and the levels of IL-1β, IL-10, IL-20, and TNF-α increased [123]. The AucCdt may modulate macrophage function in vivo by changing the proinflammatory/anti-inflammatory cytokine balance. However, recombinant AucCdt failed to stimulate IL-1β in human periodontal ligament fibroblasts (HPLF) but upregulated IL-6 and the receptor activator of NF-κB ligand (RANKL) [124, 125]. Another report found that human gingival fibroblasts (HGF) challenged with A. actinomycetemcomitans exhibited increased expression of IL-6 and IL-8 [126]. This increase in expression was not attributed to a specific product, but the Cdt is a reasonable candidate. RANKL was also upregulated in response to the HdcCdt in a T-lymphocyte leukemia cell line (Jurkat) [127]. RANKL is a member of the TNF ligand superfamily and binds to its associated receptor RANK on
osteoclast progenitor cells [128]. This action leads to the differentiation of the progenitor cells into bone-resorbing osteoclasts [129]. Since expression of these molecules is involved in the induction of osteoclast differentiation, they have a central role in the regulation of bone resorption which is a key feature of periodontitis [I30,131]. A recent review by Belibasakis and Bostanci [132] presents a thorough discussion of the role of the Cdt on host inflammatory responses and associated bone remodeling events.

4. AB Toxin Assembly and Cell Intoxication

There are common features of the AB toxins for assembly of the holotoxin complex and intoxication of target cells. Synthesis and assembly of both the A and B chains or subunits are required for these toxins to be effective against susceptible cells. The B chains, in either monomeric or multimeric forms are essential for the delivery of the A chain into the cell. Since these toxins act on internal pathways or processes, unlike pore-forming or membrane-damaging toxins, it is essential that the A chains are transported across the plasma membrane. Endocytosis appears to be a common mechanism for internalization of the A chain. Anthrax toxin can be considered a minor exception to the general mode of AB toxin assembly since active holotoxins are heterogeneous. Holotoxins can consist of the B chain polymer together with various combinations of the A chains (see Figure 1) [15]. The SeCdt represents another deviation from the standard internalization process because S. enterica serovar Typhi carries and expresses only the cdtB gene [46]. It has been hypothesized that delivery of the SeCdt to cells is carried out by internalized bacteria. The bacteria invade cells via a membrane enclosed organelle in macrophages known as a Salmonella-containing vacuole (SCV) [133,134]. The bacteria replicate in this vacuole which may intersect with the Golgi apparatus or endoplasmic reticulum (ER) where the secreted CdtB protein can be released for processing.

Comparison of representative examples of several of the most common classes of AB toxin demonstrates the similarities and differences among the cell intoxication mechanisms (Figure 5). As stated previously, the ricin holotoxin binds to a glycoprotein or glycolipid receptor, containing a terminal β-D-galactopyranoside residue, on the surface of susceptible cells [8]. There are two binding sites in the B chain. The holotoxin enters the cell via endocytic mechanisms enlisting primarily both clathrin-coated pits and clathrin-independent membrane invaginations [135]. Binding of the toxin to the membrane and endocytosis may be concentrated in areas containing lipid rafts since these processes are affected by cholesterol depletion. The results of cholesterol depletion experiments indicated that lipid rafts may also be important for the transfer of ricin from endosomes to the Golgi apparatus. The complex collection of proteins that mediate retrograde transport of proteins from endosomes to the Golgi apparatus has been summarized [136], but their specific roles in ricin transport have not been detailed. Likewise, details of the mechanism of retrograde transport of ricin from the Golgi apparatus to the ER have not been worked out. However, there is some data to suggest that phospholipase A2 contributes to the process [137,138]. Finally, ricin translocation to the cytosol may depend on sequences enriched in hydrophobic amino acids [139].

There are many similarities between the mechanism of cell intoxication by ricin and Shiga toxin. This is expected since both contain an A chain comprised of N-glycosidase. The Shiga toxin-receptor Gb3 resides in the outer leaflet of the plasma membrane [13]. Shiga toxin binding to susceptible cells is affected by the concentration of Gb3 in the membrane, the isoforms present, and the types of other glycosphin-golipids adjacent to Gb3 and cholesterol. However, it has not been unequivocally established if lipid rafts are involved [135]. Similar to ricin, multiple endocytic processes, including clathrin-coated pits, bring Shiga toxin into the cell. Unlike ricin, clathrin appears to be required for carrying Shiga toxin from early endosomes to the Golgi apparatus [136]. Two mammalian cell SNARE complexes, containing the proteins GI5, GS28, syntaxin-5, syntaxin-6, and syntaxin-16, YTK6, VTIIa, and VAMP4, play roles in the retrograde transport of the toxin. Also like ricin, Shiga toxin translocates from the ER to the cytosol due to hydrophobic membrane-spanning domains in the protein [140].

Anthrax is an A2,B7 class toxin with each of the LF and EF A chains having distinct modes of action [15]. To initiate the assembly process, a monomer of the PA B chain binds to the ATR (Figure 5). Previously characterized cell surface receptors tumor endothelioma marker-8 (TEM8), capillary morphogenesis protein-2 (CMG2), and integrin β1 have been implicated in PA binding due to the presence of a von Willebrand factor A (vWA) domain in the proteins [141–145]. However, a recent study employing a murine model has shown that TEM8 may be the only receptor candidate to function in vivo [143]. The toxin-receptor complex is associated with lipid rafts and endocytosis is clathrin dependent [146]. Once PA binds to the ATR, a small amino terminal fragment of the former protein is removed by furin-like proteases. This processing triggers PA to spontaneously form a heptamer, or possibly an octamer. The polymerized PA complex then binds 3–4 LF and/or EF chains [147, 148]. The holotoxin enters the cell by endocytosis. The holotoxin forms a membrane channel in the early endosome which allows the translocation of the LF and EF subunits to the cytosol.

Botulinum toxin is made as a single inactive polypeptide of approximately 150 kDa that is internalized by receptor-mediated endocytosis. However, relatively little is known about this process in comparison to the other members of the AB toxin family. The polypeptide appears to bind to a receptor in the early endosome (Figure 5). The toxin recognizes gangliosides and has the greatest affinity for GT1b (Neu5Acα2(2–3)D–Galβ1(3–D–GalNAcβ1(4–)]Neu5Acα2(2–8)Neu5Acα2(2–3)]D–Galβ1(4–D–GlcPβ1(1–1)-ceramide) [149]. The polypeptide is composed of a 100 kDa peptide, representing the B chain (heavy chain) and a 50 kDa A chain (light chain). The heavy and light chains are nicked by a protease but held together by a single disulfide bond. Dissolution of the disulfide bridge releases the light chain into the cytosol while the heavy chain remains in the endosome [150].
Studies of the AaCdt have provided insight into the mechanism by which the Cdt holotoxin is synthesized and assembled in the bacterium. Each of the three subunits are translated with an N-terminal signal or leader sequence that allows the proteins to traverse the cytoplasmic membrane of the Gram-negative bacterium. It has been hypothesized that CdtB and CdtC protein precursors are transported across the membrane via a sec-dependent pathway coordinated with cleavage of the leader sequences by signal peptidase I [151]. This results in accumulation of the mature proteins in the periplasmic space. Precursor CdtA protein is processed by signal peptidase II and modified with a glycerolipid. The modified hydrophobic CdtA is inserted into the periplasmic space side of the outer membrane, via a lipid-binding consensus sequence (lipobox), where it forms a complex with the mature forms of CdtB and CdtC. The precursor holotoxin is then processed by an unidentified protease, possibly signal peptidase II, that removes the N-terminus along with the glycerolipid. This cleavage step allows the now hydrophilic holotoxin to exit the bacterium. Indeed, there is support for a role of a truncated CdtA in the assembly of the mature toxin since both 23- and 17-kDa CdtA peptides were found in H. ducreyi extracts [152]. Similar results were reported for the AaCdt [153].

As discussed in Section 3.3.1, there are three types of molecules, N-linked fucose containing glycoproteins, gangliosides, and transmembrane proteins, that may serve as receptor candidates for the various Cdts and in various types of susceptible cells [55, 64, 72]. The Cdt appears to recognize at least one of these types of receptors and most likely binds to the cell via the CdtA aromatic patch region and the CRAC motif in CdtC [58, 70, 71, 155]. It appears that the cholesterol concentrated in these membrane domains and lipid rafts may also be important for the binding of some Cdts to specific types of cells due to an interaction between the cholesterol concentrated in these membrane domains and the CRAC motif in CdtC [58, 70, 71, 155]. It appears that the AaCdt, as well as several other members of the Cdt subfamily, may require the localization of a specific receptor with lipid rafts to maximize binding to the cell surface.
Experiments performed primarily with the HdCdt suggest that the toxin enters cells by a dynamin-dependent endocytosis that may be clathrin dependent or independent based on the techniques used to assess this property [21, 86]. Dynamin, a GTPase, is believed to be a classical regulator that recruits effectors of endocytosis [156, 157]. Several earlier models of Cdt intoxication predicted that the Cdt holotoxin, rather than a CdtB-CdtC dimer or CdtB alone, is taken up by early endosomes [71, 158, 159]. However, using a precise protein tagging technology in combination with live-cell imaging, we found that CdtA, and a smaller amount of CdtC, from the AaCdt, remains on the cell surface during the intoxication of CHO cells [160]. A significant amount of CdtC was found, in addition to CdtB, inside the cells. These observations supported data from our earlier adhesion studies that indicated that the CdtA subunit may have a more active role than CdtB in the binding of the holotoxin to cells [56, 59] and that CdtC could be detected, by immunofluorescence microscopy, inside cells [39]. Based on data from our experiments and those of Guerra et al. [86, 153, 159], a general model for the Cdt intoxication process is presented (Figure 5). In this model, the Cdt binds to a cell surface receptor via the CdtA and CdtC subunits. This binding may take place in areas of the membrane that contain lipid rafts [70, 71]. However, it should be pointed out that Cdt binding to cells may be either cholesterol-dependent or independent based on the specific cell type or cell line, species-specific origin of the toxin, toxin concentration, and reorganization of the putative receptor for toxin binding within protein components of the cell membrane [58, 161]. After initial binding, CdtA remains on the surface of the cell in association with the specific receptor. A CdtB-CdtC heterodimer undergoes endocytosis due to an association of CdtC with the lipid raft. For this model, we proposed that the endocytosis is clathrin independent because it has been reported that knock-down of the clathrin gene by RNA interference does not reduce Cdt activity in intoxicated cells [86]. However, there is conflicting data suggesting that the endocytic uptake of the HdCdt in HeLa and HEP-2 cells depends on clathrin-coated pits [21]. The early endosome is formed with the help of dynamin II and carries the CdtB-CdtC complex to the Golgi apparatus where it is sulfated. CdtB is then transferred to the ER where it is translocated in a retrograde fashion and glycosylated. CdtB is translocated by a non-ER-associated degradation (ERAD) pathway because the protein does not enter the cytosol but is targeted to the nucleus. We support the possibility that CdtC is also delivered to the Golgi apparatus and ER in complex with CdtB. However, CdtC most likely undergoes degradation by the usual ERAD pathway.

CdtB enters the nucleus of the cell via the ER. In a fluorescent tagging approach combined with the isolation of nuclei from intoxicated CHO cells, we found that the AaCdtB reaches the nucleus within 4.5 hours after intoxication [160]. These results were in good agreement with those of several studies reviewed in Guerra et al. [86]. The CjCdtB was detected in the nucleus of COS-1 and REF52 cells 4 hours after microinjection [75]. CdtB can apparently cross the ER and nuclear membranes due to the presence of a nuclear localization signal (NLS) sequence. Two different NLS sequences have been reported in AaCdtB and EcICdtB. The AaCdtB NLS is comprised of amino acids 48–124 in the amino-terminal part of the protein [83]. In contrast, EcICdtB NLS was located in two regions comprised of amino acids 195–210 and 253–268 [162]. A mutagenesis approach was used to show that both types of NLS sequences contributed to nuclear translocation. Changing tandem arginine residues in the latter putative EcICdtB NLS to threonine and serine had no effect on the in vitro DNA activity of CdtB but reduced CdtB translocation to the nucleus and induction of cell cycle arrest. We obtained similar results when the consecutive arginine residues were mutated in the AaCdtB [160]. When we made amino acid substitutions for the two arginine residues located in the putative NLS site described by Nishikubo et al. [83], there was no effect on the activity of the reconstituted toxin or nuclear localization of CdtB. It is possible that monopartite or bipartite arginine and/or lysine residues necessary for classical NLS function may not be required by this atypical NLS.

5. Cell Tropism of the Cdt

The ability of the Cdts to exhibit a broad cell tropism is one of the intriguing properties of this subclass of Ab toxins. The species-specific Cdts, as a group, usually cause cell distention, cell cycle arrest in either the G1/S or G2/M transition phase, and/or apoptosis in a variety of mammalian cell types [88]. The effects appear to be cell-type specific [89, 90]. In general, rapidly proliferating cells seem to be most sensitive to the Cdt. For example, cells within the deeper layers of epithelium should be good targets for the toxin since they continuously proliferate prior to terminal differentiation as they migrate towards the epithelial surface [163].

5.1. Epithelioid Cells. Human epithelial-like cell lines such as HeLa [7, 91, 104, 109, 164] and HEP-2 [38, 165] are arrested at the G2/M phase transition by a variety of the species-specific CdtB. A well-known human colon carcinoma cell line, Caco-2 [91], and two human keratinocyte cell lines, HaCat [108, 166] and Henle-407 (intestinal epithelial cells) [75], are also sensitive to the CjCdt and HdCdt. These epithelioid cell lines typically are arrested at the G2/M interphase of the cell cycle when exposed to the AaCdt. Although Caco-2 cells undergo the characteristic cell cycle arrest at the G2/M transition, it has been reported that no prior cell distention was evident [91]. It is not known why distention fails to occur in some types of cells when exposed to the toxin. Perhaps effects of the Cdt on the expression of scaffolding and structural proteins varies among different cell types.

We have shown that KB cells, thought to have originated from epidermoid carcinomas of the larynx, are sensitive to the AaCdt [165]. The KB cell line has traditionally been used to study A. actinomycetemcomitans invasion [167–169]. We also found that an immortalized human orolabial cell line (GMSM-K), transformed with simian virus (SV40) [170], is particularly sensitive to the toxin [61, 171]. This cell line was employed as an early epithelial cell model for the AaCdt prior
to our ability to culture primary human gingival epithelial cells (HGEC). Interestingly, this cell line was arrested at the S phase of the cell cycle in response to the toxin. Fedor et al. [84] found that a low dose of the EcCdtx progressively induces DNA double-strand breaks in cells in the S phase. Cells isolated from the epithelial layers of human gingival tissue, obtained from crown-lengthening procedures performed on clinically healthy adults, were arrested at the G2/M interphase of the cell cycle when exposed to recombinant AaCdtx [172]. Immortalized human gingival keratinocytes, exposed to a Cdt-containing extract from A. actinomycetemcomitans, also exhibited cell cycle arrest at G2/M [173]. The ability of the AaCdtx to inhibit the proliferation of primary and immortalized HGEC provided supporting evidence that the toxin could have a significant role in damaging the gingival epithelial layer important for protecting the underlying connective tissue and tooth supporting structures from the damaging effects of oral pathogens.

5.2. Mesenchymal Cells. The results of studies examining the ability of the Cdt to inhibit the proliferation of fibroblasts are mixed. In one report, Chinese hamster lung (Don) fibroblasts were reported to be sensitive to the Hdcdt and NIH 3T3 fibroblasts were resistant [166]. Stevens et al. [108] reported only a "modest" inhibitory effect on human foreskin fibroblasts using the Hdcdt. In another study, extracts of A. actinomycetemcomitans containing the Cdt inhibited DNA synthesis in human periodontal ligament and gingival fibroblasts [101]. The cells became distended but remained viable. Aggregatibacter actinomycetemcomitans extracts that did not contain the toxin failed to elicit these effects. In a follow-up study, this same group reported that the periodontal ligament and gingival fibroblasts were arrested at both the G1 and G2/M phases of the cell cycle when treated with the Cdt-containing bacterial extract from A. actinomycetemcomitans or recombinant Hdcdt [100]. We found that primary oral fibroblast-like cells including human periodontal ligament fibroblasts (HPLF) [61], human gingival fibroblasts (HGF) [172] cementoblasts [171], and osteoblasts (unpublished observations) appear to be relatively resistant to the DNA-damaging and cell cycle arresting effects of the Cdt. Cells cultured from the connective tissue layer of human gingival explants and identified as fibroblasts (HGF) failed to enter cell cycle arrest following exposure to recombinant AaCdtx [172]. It is clear that Cdt intoxication of target cells is a very complex set of events that can have different effects on various pathways and may be dependent on the source of toxin, target cell type, and possibly the environment of the cell [90, 100].

5.3. T-Lymphocytes. Human CD4+ and CD8+ T lymphocytes as well as monocytes undergo cell cycle arrest, without cell distention, in the G2 phase when treated with extracts containing the AaCdtx [106, 174]. Lymphocytes treated with recombinant AaCDT underwent apoptosis via activation of the caspase cascade [116]. There was evidence of a role for Bcl-2, an apoptosis regulator, in pathway [175]. Apoptosis induced by the AaCdtx in these cells involved the upregulation of Fas and FasL, downregulation of Bcl-2, and activation of caspase-3 and was dependent on the presence of monocytes [174]. Binding of FasL, the Fas ligand, to the CD95, the Fas receptor, on target cells is one of the pathways that leads to the initiation of apoptosis [176]. Two independent pathways, caspase-dependent (early) apoptotic cell death and caspase-independent (late) cell death, may be involved in Cdt-induced death in some types of T-cells such as the human leukemic cell line MOLT-4 [177, 178].

5.4. Macrophages. Macrophages may be potential in vivo targets of the toxin since it was found that apoptosis in non-proliferating U937 cells required the phosphatidylinositol-3,4,5-triphosphate phosphatase activity of recombinant AaCdtx [179]. However, cell cycle arrest and induction of apoptosis in these cells was dependent on the action of the nucleosome activity of this subunit on proliferating cells. There is additional evidence that the AaCdtx disrupts macrophages by inhibiting phagocytic activity, modulating nitric oxide production, and altering the expression of proinflammatory and anti-inflammatory cytokines [123, 180]. Human macrophages exposed to the AaCdtx did not exhibit Cdt-induced apoptosis but experienced inhibition of the phosphatidylinositol-4,5-bisphosphate 3-kinase signaling pathway and an associated increase in glycogen synthase kinase 3β affecting production and secretion of the cytokines IL-1β, IL-6, and tumor necrosis factor (TNF)-α [119].

6. AB Toxins and Infectious Diseases

The AB toxins can be associated with either noninfectious disease or infectious disease. The noninfectious disease types are the toxins produced by plants. These toxins can promote disease when the toxin itself or cellular material containing the toxin is inhaled or ingested by the host. AB toxins produced by bacteria are of the infectious disease type. Diseases associated with these toxins become established when the host becomes infected with the microorganism or spores (anthrax, botulinum, and tetanus toxins) produced by the microorganism. In most cases, the bacterial AB toxins are produced by one to several species within a single genus of the bacterium and represent the primary "virulence factor" in the disease. For example, the botulinum toxin is produced by Clostridium botulinum, C. butyricum, C. baratti, and C. argentinense and the primary symptoms of botulism are attributed to the activity of the toxin. Cholera toxin is produced by Vibrio cholerae and the primary symptoms of cholera are due to the activity of the toxin. Many of the AB toxins have each been clearly associated with a single specific disease. In contrast to the other AB toxins the Cdt is produced by members of eight different genera of Gram-negative bacteria and can be potentially related to six different diseases (see Table 2). Furthermore, it has been difficult to prove that diseases associated with some Cdt-producing bacteria are due primarily to the activities of the toxin.

6.1. Association of the Cdt with Disease. In site of the genetic diversity of the various Cdts, there is a commonality of
function based on a connection between the Cdt-producing strains of bacteria and diseases that compromise tissue layers comprised of epithelial or epithelial-related cells. Due to the complexity of mucosal related diseases, cause, and effect, relationships between the toxin and clinical symptoms of disease have had to rely on indirect associations. Often the relationship is based solely on the isolation of Cdt-producing strains from a diseased subject as in a case study reporting the isolation of \( E. \) coli 055:K59:H4 from a child having a history of gastroenteritis and encephalopathy [181]. Similar approaches have been used to show that EPEC belonging to serogroup O127, isolated from children with diarrhea, contain \( cdt \) gene sequences detected by PCR [182]. However, not all of the O127 isolates from the diseased subjects produced the Cdt and some Cdt-producing strains were isolated from control subjects. Enterohemorrhagic \( E. \) coli (EHEC) can also be etiological agents for diarrhea and hemolytic uremic syndrome (HUS), a cause of acute renal failure in children. \( Escherichia \) coli 057:H7, a Cdt\(^+\) strain, is a predominant serogroup in HUS and the toxin is capable of contributing to the endothelial or vascular injury associated with the disease [183]. However, disease associations are complicated by the fact that other virulence factors such as endotoxin and Shiga toxin are present in the EHEC.

Attempting to establish a major role for the Cdt in the virulence of \( C. \) jejuni in cases of acute bacterial gastroenteritis has also been challenging. The Cdt appears to be a predominant virulence factor in \( Campylobacter \) sp. [184] and, in addition to endotoxin which is a structural component of all Gram-negative bacteria, it is the only other toxin identified in these species. Cdt\(^−\) mutants of \( C. \) jejuni failed to cause gastroenteritis in a NF-\( \kappa \)-B-deficient mouse model [185]. Other studies have provided evidence that the \( GjCdt \) may support the ability of the bacterium to invade tissues in a severe combined immunodeficient (SCID) mouse model [186]. A study examining the role of \( C. \) jejuni in human gastroenteritis found that disease in patients infected with Cdt\(^−\) strains was clinically indistinguishable from that in patients harboring Cdt\(^+\) strains [187]. It was concluded that Cdt production was not essential for the development of the diarrhea associated with \( Campylobacter \)-related disease.

Other studies have examined the role of the \( HdCdt \) in the formation of ulcers, a characteristic of the sexually transmitted disease (STD), chancroid. Using a rabbit model, it was found that \( cdt \) gene knock-out mutants of \( H. \) ducreyi were just as virulent as the wild-type strain [108, 188]. A combination of a Cdt\(^−\) strain of \( H. \) ducreyi and recombinant \( HdCdt \) produced inflammatory skin lesions that progressed to ulceration in the same rabbit model [189]. Even though a combination of \( Haemophilus influenzae \) and \( HdCdt \) failed to cause ulcers to form, a contributory role for other \( H. \) ducreyi virulence factors could not be ruled out. In a skin pustule formation model employing human volunteers, independent challenges with a \( cdt \) gene knock-out mutant and wild-type strain resulted in the development of papules at most sites of infection [190]. The results of several trials indicated that the \( HdCdt \) was not required for pustule formation in humans.

### 6.2. Contribution of the AaCdt to Oral Infectious Disease.

Clearly, the variety of cell types and infectious diseases associated with the Cdt subfamily of AB toxins has added layers of complexity in attempts to unequivocally demonstrate a cause and effect relationship between the toxin and disease at the clinical level. Further complications can be attributed to the presence of multiple virulence factors, including endotoxin, in Cdt\(^+\) strains. Application of standard epidemiological, serological, and genetic approaches including: (i) identification of Cdt\(^+\) strains or evidence of \( cdt \) gene sequences in diseased but not in control subjects, (ii) presence of Cdt or Cdt-neutralizing antibody titers in diseased and healthy subjects, respectively, and (iii) use of \( cdt \) gene knock-out mutants in animal and human disease-related models have for the most part failed to provide strong support for an active \textit{in vivo} role of the various enteric- and STD-associated Cdts. These same problems have plagued investigations of the \( AaCdt \). Data from some studies have indicated a correlation between the presence of Cdt\(^+\) \( A. \) \textit{actinomycetemcomitans} strains and periodontal disease while other studies have failed to find a difference between colonization of subjects with disease by Cdt\(^−\) and Cdt\(^+\) strains. A representative collection of studies in which Cdt\(^+\) and Cdt\(^−\) strains of \( A. \) \textit{actinomycetemcomitans} were quantified by either culturing or PCR in localized aggressive periodontitis (LAP) subjects and clinically healthy controls yielded mixed results in relation to a positive correlation between the toxin and disease [19, 191–197]. In one of these studies, a random cohort of 500 Ghanaian adolescents (mean age 13 years) was examined for the presence of \( A. \) \textit{actinomycetemcomitans} and expression of the \( AaCdt \) at baseline and in 397 of the subjects two years later [191]. No statistically significant correlation between Cdt\(^+\) and Cdt\(^−\) strains and attachment loss indicative of progression to LAP was observed in the follow-up examination.

Similar conflicting results were obtained when systemic and neutralizing \( AaCdt \) antibody titers were measured in diseased and healthy patients [198–200]. Based on these studies, a cause and effect relationship has not been confirmed between the \( AaCdt \) and the development of some forms of periodontal disease. To the best of my knowledge, there have not been any reports of studies designed to assess the activities of the Cdt in cases of \( A. \) \textit{actinomycetemcomitans}-involved infectious endocarditis. \textit{Aggregatibacter} \textit{actinomycetemcomitans} is a member of the HACEK group of bacteria that cause infective endocarditis in children [201].

In spite of the conflicting data from epidemiological studies, complexities of polymicrobial diseases like LAP and chronic periodontitis, and the limitations of animal models to adequately represent human periodontal disease, significant progress has been made towards identifying an \textit{in vivo} role for the \( AaCdt \). A significant amount of information has been obtained from studies assessing the effects of the various species-specific CdtS on cells in culture an additional set of new questions can be approached using tissue. The effects of the Cdt on sensitive cells \textit{in situ} are expected to be different, due to the spatial distribution and unique interactions in tissue, than on the same cells grown \textit{in vitro} in tissue culture.
In an elegant study, Ohara and coworkers [202] exposed areas of the gingival sulcus in live rats to recombinant wild-type and mutated AaCdt. The infected tissue was observed in vivo and harvested 1–3 days following exposure to the toxin. Tissue samples were decalcified, embedded in paraffin and sectioned parallel to the long axis of the tooth. Sections were stained with hematoxylin and eosin (H&E) and proliferating cell nuclear antigen (PCNA) antibody. Cells in the junctional epithelium became desquamated and detached by day three of exposure to the wild-type toxin. The biologically inactive mutated AaCdt had no effect on the tissue. The amount of PCNA staining was significantly reduced in the junctional and gingival epithelia. This observation was an indication that proliferation of the epithelial cells in the tissue was inhibited by the Cdt.

My laboratory examined the effect of recombinant AaCdt on human gingival tissue obtained from healthy adults having no clinical signs of periodontal disease [172]. As in the case of rat gingiva, we observed extensive desquamation and detachment of the keratinized surface layer of the human tissue and obvious disruption of the underlying spinous layer and rete pegs in H&E stained paraffin sections by 18 hours of exposure to the toxin. On the gross morphological level, the tissue appeared swollen and the spinous layer became substantially thickened. The epithelial cells were enlarged or distended with an apparent loss of cell junctions as indicated by the increased separation of cells in the tissue. The observed changes in the toxin-treated tissue were dose and time dependent. Tissue exposed to an inactive AaCdt mutated in the CdtB subunit did not exhibit these major morphological changes. Stained sections of untreated gingival tissue from adult subjects with clinical signs of oral inflammation exhibited the same histological appearance as the AaCdt-treated tissue.

We extended the analysis of AaCdt-treated human gingival tissue by assessing effects on cell junctions in situ [203]. Claudin 1, a primary component of tight junction filaments [204], was poorly detected by immunofluorescence microscopy, in untreated tissue sections. E-cadherin, the major membrane protein in adherens junction complexes [205, 206] was readily detected as brightly fluorescent areas surrounding the periphery of the epithelial cells in the tissue. Healthy human gingivae have a relatively high level of E-cadherin in the basal cell layer [207]. Exposure of the tissue to the AaCdt resulted in an observable increase in the relative fluorescence of E-cadherin and redistribution in the cytosol. These changes correlated with an increase in E-cadherin mRNA expression in 50% of the tissue samples tested. There were also indications that the toxin caused detachment of the basal cells from the basement membrane. Gingival tissue exposed to the mutated AaCdt exhibited minimal changes in E-cadherin distribution. Wild-type toxin-treated tissue also showed increases in β-catenin and F-actin staining and in mRNA expression in 25% and 63%, respectively, of the samples examined. Both β-catenin and actin are cytosolic components of the adherens junction apparatus [206]. F-actin assemblies having a morphology similar to that of actin stress fibers accumulated in CHO cells treated with recombinant EcCdt [208].

The morphological changes observed in AaCdt-treated gingival tissue and the cell tropism exhibited by the toxin (see Section 5) support a possible multipronged attack for the activities of the toxin in oral infectious disease. A model has been proposed outlining the putative contributions of the AaCdt to the cascade of events relevant to the development of periodontal disease [209]. In this model, the AaCdt interacts with three different types of cells, gingival epithelial cells, gingival fibroblasts, and infiltrating inflammatory cells (T-lymphocytes and macrophages) in three successive stages during the development and progression of the disease. These interactions result in inhibition of cell proliferation and alteration of cell signaling pathways. The coordinated effects of these Cdt-cell interactions could lead to the direct and indirect destruction of tissues characteristic of the periodontal disease.

7. Conclusions

The Cdt is an intriguing and novel member of the superfamily of AB toxins and continues to pose significant challenges in studies of the biology of cytoxins and their role in disease almost 30 years after the discovery of the first member of this subgroup. One Cdt enigma is that, unlike the other AB toxins, it seems to fail to live up to the tremendous potential it has to be a prominent virulence component in bacterial species associated with major diseases. For example, Campylobacter spp. such as C. jejuni cause campylobacteriosis which is one of the most common gastroenteric infections in developed and developing nations [210]. The bacterium is responsible for 400–500 million cases of diarrhea per year. It is estimated that 2 million cases of campylobacteriosis occur in the United States each year making it the most common bacterial cause of foodborne illness [211]. Yet, it has been difficult to show that the CjCdt, the only toxin other than endotoxin produced by Campylobacter sp., is involved in the disease. Associations between the Hdcdt and chancroid and the AaCdt and some forms of periodontal disease offer very similar scenarios. As discussed in Section 3.1, Cdt-producing organisms do not appear to easily shed their cdt genes even though they have a promiscuous evolutionary history and no obvious selective pressure to enforce maintenance by the producing bacterium. So, a very interesting question that remains unanswered is how can the various species-specific Cdts, which are clearly AB toxins based on their structural and biological characteristics, not play important roles in infectious diseases specific to the Cdt-producing bacterium?

Another Cdt puzzle is the molecular basis for the extensive heterogeneity. The different Cdts exhibit significant deduced amino acid sequence diversity, a broad host cell range exemplified by heterogeneity in cell attachment mechanisms and variability in the end-point of intoxication. Some Cdts appear to bind specifically structured glycan residues on glycoproteins or glycolipids, certain gangliosides, and/or some types of transmembrane G-protein-coupled receptor proteins. Also, intoxication by some Cdts leads to cell cycle arrest at either the G1 or G2/M checkpoint or cell death via an apoptotic pathway. Exhaustive phylogenetic analyses and comparisons within each of the three subunits across
the species-specific CdtS have been performed to attempt to understand this heterogeneity [48, 158]. Unfortunately, this approach has yielded few insights. For example, in spite of the marked sequence diversity among the various species-specific CdtB subunits, the largest genetic distance between clades is 0.8 [158]. The genetic distance is even smaller if the bacteriophage and putative Yersinia CdtBs are removed from the analysis. Furthermore, the amino acid residues important for the DNase-like activity (catalytic site, DNA-binding, and cation binding) are highly conserved. Yet, DNA damage caused by the different CdtBs results in G1, G2/M arrest or apoptosis. Similar to CdtB, the largest genetic distance between clades of CdtA and CdtC is 0.9 and 0.6, respectively. Therefore, CdtA and CdtC are not more diverse than CdtB which is the most highly conserved of the three subunits. All of the CdtAs have a conserved aromatic patch motif and all CdtAs and CdtCs combine to form a groove in the structure predicted to be an important receptor binding site. However, the various holotoxins appear to recognize different receptors on different types of cells. Therefore, the extensive body of work that exists for the Cdt has to be extended in order to solve the mysteries of what is arguably the most fascinating AB toxin. As pointed out by a number of other groups working in this field, the obvious current challenges are to decipher the details of the Cdt intoxication process, determine whether there are truly multiple specific receptors or a defined interplay between components in highly specialized regions of the cell membrane, and identify cause and effect relationships to confirm Cdt-mediated pathogenicity. Although the results of current studies are leading to a consensus of opinion that environmental factors dictate different outcomes for Cdt behavior, there may turn out to be more commonality among the species-specific Cdts than expected.

Conflict of Interests

The author declares that there is no potential conflict of interests with respect to the authorship and/or publication of this paper.

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