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Protein-disulfide Isomerase Displaces the Cholera Toxin A1 Subunit from the Holotoxin without Unfolding the A1 Subunit*§

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Protein-disulfide isomerase (PDI) has been proposed to exhibit an “unfoldase” activity against the catalytic A1 subunit of cholera toxin (CT). Unfolding of the CT A1 subunit is thought to displace it from the CT holotoxin and to prepare it for translocation to the cytosol. To date, the unfoldase activity of PDI has not been demonstrated for any substrate other than CT A1. An alternative explanation for the putative unfoldase activity of PDI has been suggested by recent structural studies demonstrating that CT A1 will unfold spontaneously upon its separation from the holotoxin at physiological temperature. Thus, PDI may simply dislodge CT A1 from the CT holotoxin without unfolding the CT A1 subunit. To evaluate the role of PDI in CT disassembly and CT A1 unfolding, we utilized a real-time assay to monitor the PDI-mediated separation of CT A1 from the CT holotoxin and directly examined the impact of PDI binding on CT A1 structure by isotope-edited Fourier transform infrared spectroscopy. Our collective data demonstrate that PDI is required for disassembly of the CT holotoxin but does not unfold the CT A1 subunit, thus uncovering a new mechanism for CT A1 disassociation from its holotoxin.

Cholera toxin (CT) is an AB5 protein toxin that consists of a catalytic A moiety and a cell-binding B moiety (1, 2). The B subunit is pentameric ring-like structure that adheres to GM1 gangliosides on the plasma membrane of a target cell. The A subunit is initially synthesized as a 26 kDa protein that undergoes proteolytic nicking to generate a disulfide-linked A1/A2 heterodimer. The 21 kDa CT A1 polypeptide is an ADP-ribosyltransferase that modifies and activates Gs in the host cell cytosol. CT A1 can be divided into three subdomains: the A1 subdomain contains the catalytic core of the toxin; the A2 subdomain is a short extended linker that connects the A1 and A3 subdomains; and the A3 subdomain is a globular structure with many hydrophobic residues as well as a cysteine residue involved with the single disulfide bridge between CT A1 and CT A2 (3). The 5 kDa CT A2 polypeptide maintains numerous non-covalent interactions with the central pore of the B pentamer and thereby anchors CT A1 to CT B5. A ribbon diagram of the CT holotoxin which highlights the subdomain structure of CT A1 is provided in supplemental Fig. S1.

To reach its cytosolic Gs target, CT moves from the cell surface to the endoplasmic reticulum (ER) by retrograde vesicular traffic (4). A C-terminal KDEL sequence in the CT A2 subunit is thought to target and/or retain CT in the ER (4, 5). Conditions in the ER lead to reductive cleavage of the CT A1/CT A2 disulfide bond and chaperone-assisted dissociation of CT A1 from CT A2/CT B5 (6–9). Unfolding of the free CT A1 subunit then activates the quality control system of ER-associated degradation (ERAD), thereby promoting CT A1 translocation to the cytosol (10, 11). Most exported ERAD substrates are degraded by the ubiquitin-proteasome system, but it was hypothesized that CT A1 and the A chains of other ER-translocating toxins avoid this fate because they lack substantial numbers of lysine residues for ubiquitin conjugation (12). Subsequent experimental studies verified the paucity of lysine residues protects CT A1 and other toxin A chains from ubiquitin-dependent proteasomal degradation (13–15). The translocated pool of CT A1 instead interacts with ADP-ribosylation factors and possibly other host factors to regain an active, folded conformation in the cytoplasm (16–18).

The reduced form of protein-disulfide isomerase (PDI), an ER-localized oxidoreductase and molecular chaperone (19, 20), was originally proposed to unfold the holotoxin-associated CT A1 subunit and to thereby promote the separation of CT A1 from CT A2/CT B5 (9). It was further posited that unfolding of the CT A1 polypeptide was directly coupled with toxin delivery to the Derlin-1 pore (21, 22). Release of CT A1 from PDI in preparation for translocation through the Derlin-1 pore was thought to involve the oxidation of PDI by Ero1p (23). Thus, PDI is viewed as a redox-dependent chaperone that binds to holotoxin-associated CT A1 in a reduced state, actively unfolds the toxin, and then releases the dissociated CT A1 subunit upon its oxidation by Ero1p.

Many aspects of PDI-CT A1 interactions remain controversial. Lumb and Bulleid demonstrated that PDI does not act in a redox-dependent fashion when assisting the folding of other polypeptide chains (24). Recent experiments using zebrafish and mammalian cells have questioned the role of Derlin-1 in CT A1 translocation (25). Our structural studies have indicated...
that CTA1 will unfold spontaneously after dissociation from the holotoxin at physiological temperature (17, 26, 27). It is therefore possible that, in terms of cholera intoxication, the primary function of PDI is to simply dislodge CTA1 from CTA2/CTB$_5$. The dissociated CTA1 polypeptide would then spontaneously unfold and consequently trigger the ERAD translocation mechanism.

In this work we evaluated the role of PDI in disassembly of the CT holotoxin and unfolding of the CTA1 subunit. The prevailing model of PDI-toxin interactions is largely based upon the results of a protease sensitivity assay which demonstrated that CTA1 shifts from a protease-resistant conformation to a protease-sensitive conformation in the presence of reduced PDI (9, 13, 23, 28, 29). As folded proteins are generally more resistant to proteinase than unfolded variants of the same protein, this shift was interpreted to represent the PDI-induced unfolding of CTA1 and the PDI-induced displacement of CTA1 from CTA2/CTB$_5$. Yet this assay, which was typically performed at 30 °C, only provides an indirect measure of protein folding. Here, we developed a surface plasmon resonance (SPR) assay to directly monitor in real time the disassembly of the CT holotoxin. Biophysical methods were used to directly examine the conformation of CTA1 in the presence or absence of PDI. Cell-based assays further probed the role of PDI in CT intoxication and CTA1 translocation. Our collective data indicate that PDI does not unfold the CTA1 polypeptide and that PDI can displace CTA1 from CTA2/CTB$_5$, in a process which does not involve substantial alterations to the structure of CTA1. In further contrast with the current model of PDI-toxin interactions where PDI acts as an unfoldase, we found that the PDI-induced shift of CTA1 to a protease-sensitive conformation does not correlate to the disassembly of the CT holotoxin. Moreover, the release of PDI from CTA1 does not require Ero1p but instead results from the spontaneous unfolding of CTA1 which occurs after its dissociation from the holotoxin. The unstable nature of the isolated CTA1 polypeptide at physiological temperature thus plays a central role in toxin translocation and leads to a new model of CT disassembly in the ER.

**EXPERIMENTAL PROCEDURES**

**Materials**—His-tagged CTA1, CTA1$_{1–168}$ and CTA1$_{1–133}$ constructs were purified as previously described (26). CT and the CTA1/CTA2 heterodimer were purchased from List Biologicals (Campbell, CA), while PDI, chemical and other reagents were purchased from Sigma-Aldrich. Anti-CTA and anti-CTB antibodies were purchased from Sigma-Aldrich; anti-PDI and anti-KDEL antibodies were purchased from Enzo Life Sciences (Plymouth Meeting, PA); and the anti-CTA monoclonal antibody 35C2 (30) was kindly provided by Dr. Randall K. Holmes (University of Colorado Denver School of Medicine).

**SPR**—Experiments were performed with a Reichert (Depew, NY) SR7000 SPR Refractometer as previously described (17). Antibodies were perfused over the plate at the following dilutions in phosphate-buffered saline containing 0.05% Tween 20: anti-PDI antibody, 1:10,000; anti-CTB antibody, 1:15,000; anti-CTA monoclonal antibody, 1:500; anti-CTA polyclonal antibody, 1:5,000; and anti-KDEL antibody, 1:10,000. The flow rate for all steps was 41 μl/min.

**Cytochalasin D (Cyto D) and Calcium and Magnesium Chloride**—Experiments were performed with a Reichert (Depew, NY) SR7000 SPR Refractometer as previously described (17). Antibodies were perfused over the plate at the following dilutions in phosphate-buffered saline containing 0.05% Tween 20: anti-PDI antibody, 1:10,000; anti-CTB antibody, 1:15,000; anti-CTA monoclonal antibody, 1:500; anti-CTA polyclonal antibody, 1:5,000; and anti-KDEL antibody, 1:10,000. The flow rate for all steps was 41 μl/min.

**13C Labeling and Purification of CTA1-His$_{6}$—Escherichia coli strain BL21 pLysS was transformed with pT7CTA1His$_{6}$, an inducible CTA1-His$_{6}$ expression plasmid (31). 13C-labeled toxin was generated by culturing the strain at 37 °C in M9 minimal media supplemented with uniformly 13C-labeled 13C$_{6}$-d-glucose (Cambridge Isotope, Andover, MA) as the sole metabolic carbon source. The culture was shifted to 22 °C when it reached an A$_{600}$ of 0.6, and protein expression was induced overnight by the addition of 1 mM IPTG. The expressed protein was then purified in 8 M urea buffer as described previously (26). The purity of the protein sample was assessed by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) with Coomassie staining. In order to facilitate protein refolding, slide-a-lyzer 3500 MWCO cassettes (Pierce) were used for dialysis as previously described (27). After dialysis, CTA1-His$_{6}$ was lyophilized for FTIR experiments. Control experiments using circular dichroism (CD) confirmed that lyophilization did not alter the structure of CTA1.

**Determination of Protein Secondary Structure by Fourier Transform Infrared (FTIR) Spectroscopy**—Protein samples for FTIR measurements were prepared in a D$_2$O-based buffer containing 10 mM sodium borate pH 6.6 that corresponds to pH 7.0. The samples contained either unlabeled PDI, uniformly 13C-labeled CTA1, or both PDI and 13C-labeled CTA1 at a 1:1 molar ratio. FTIR spectra were measured on a Jasco 4200 FTIR spectrometer at 0.964 cm$^{-1}$ spectral resolution and a set resolution of 1 cm$^{-1}$. The absorbance spectra were calculated using the respective buffers as reference and corrected by subtraction of H$_2$O vapor contribution, smoothing, and baseline correction in the amide I region. Curve fitting was conducted using the component frequencies from the second derivative spectra, as described in (32, 33) and elaborated upon in the supplemental data. No appreciable signals from the second derivative spectra were detected in the region above 1700 cm$^{-1}$.

**Toxicity Assay**—Cells were seeded into 24-well plates and grown overnight to 80% confluency. The medium was removed and replaced with serum-free medium containing various concentrations of CT. After 2 h, the cells were processed for determination of cAMP levels as described in Ref. 34. The basal levels of cAMP determined from un intoxicated cells were background-subtracted from the values obtained for toxin-treated cells; the maximal response from the parental control cells was arbitrarily set to 100%; and all other results were expressed as ratios of that 100% value. All conditions were performed in triplicate.

**CTA1 Transfection Assays**—Cells seeded to 80% confluency in 6-well plates were transfected with pcDNA3.1/CTA1 (35) using Lipofectamine (Invitrogen) according as per manufacturer’s instructions. For the translocation assay, cells were incubated at 24 h post-transfection in methionine-free medium for 1 h before [35S]methionine was added for another hour. Digitonin was then used to generate separate membrane and cytosolic fractions as previously described (34). Both cellular fractions were immunoprecipitated with an anti-CTA antibody. SDS-PAGE with PhosphorImager analysis was used to visualize and quantify the immunosolated material. The extent of CTA1 translocation for each cell line was calculated with the following equation:

$$\text{Percentage of Translocation} = \frac{\text{Immunosolated Material in CTA1 Transfected Cells}}{\text{Immunosolated Material in Parental Control Cells}} \times 100$$

**RESULTS AND DISCUSSION**

The results presented here suggest that PDI does not unfold the CTA1 polypeptide and that PDI can displace CTA1 from CTA2/CTB$_5$, in a process which does not involve substantial alterations to the structure of CTA1. Furthermore, the release of PDI from CTA1 does not require Ero1p but instead results from the spontaneous unfolding of CTA1 which occurs after its dissociation from the holotoxin. The unstable nature of the isolated CTA1 polypeptide at physiological temperature thus plays a central role in toxin translocation and leads to a new model of CT disassembly in the ER.
equation: \% cytosolic CTA1 = CTA1 supernatant signal/ [CTA1 supernatant signal + CTA1 pellet signal].

For the toxicity assay, cells were exposed to ice-cold acidic ethanol at 3 h post-transfection and were processed for determination of cAMP levels as described in (34). An additional set of transfected cells were chased overnight to determine the relative levels of CTA1 expression via metabolic labeling and immunoprecipitation. Values obtained from the cAMP assay were standardized to CTA1 expression levels before expressing the data as percentages of the signal obtained from the parental control cells.

RESULTS

PDI-mediated Disassembly of the CT Holotoxin—SPR was used to examine the interaction between CT and PDI (Fig. 1). CT was appended to a GM1-coated sensor slide, and anti-CTA or anti-CTB antibodies were perfused over the slide for control measurements (Fig. 1A). Both antibodies generated an increase in the refractive index of the sensor slide, thus indicating direct antibody binding to the CT holotoxin. The anti-CTB antibody produced a stronger signal than the anti-CTA antibody because, given the AB5 structure of the holotoxin, there was 5-fold more CTB than CTA on the sensor slide. Differences in antibody affinity could also account for some of the differences in anti-CTB versus anti-CTA signal strength. The anti-KDEL antibody did not produce a positive signal (Fig. 1A), possibly because the presence of the CTA1 subunit blocked antibody access to the KDEL tag at the C terminus of CTA2 (supplemental Fig. S1). PDI and oxidized PDI did not bind to the CT holotoxin (Fig. 1A), but an interaction between reduced PDI and CT was detected (Fig. 1B). These observations were consistent with a published report on the redox-dependent interplay between PDI and CT (9).

Previous work has shown that reduced PDI can remove CTA1 from the CT holotoxin (9). This event was documented by our SPR system as well (Fig. 1B). After PDI bound to the CT holotoxin, a time-dependent loss of signal was observed. This indicated that PDI was released from the CT sensor slide after its initial binding. If only PDI was lost from the sensor, then the signal strength would revert to the baseline value which was derived from the mass of the bound holotoxin. However, the signal actually dropped to a negative value. This indicated that a portion of the holotoxin had also been released from the sensor slide. To confirm that reduced PDI had displaced CTA1 from the holotoxin, anti-CTA antibodies were perfused over the CT sensor slide. In contrast to our control experiment with the intact CT holotoxin (Fig. 1A), the anti-CTA antibody did not generate an increase in the refractive index of the PDI-treated holotoxin (Fig. 1B). CTA1 was therefore absent from the PDI-treated CT sensor slide. Perfusion of an anti-PDI antibody over the sensor slide also demonstrated that PDI was absent from the plate (Fig. 1B). Previous reports have demonstrated that reduction of the CTA1/CTA2 disulfide bond is not sufficient to remove CTA1 from the holotoxin (9, 36), which was also documented in our SPR system: the CT holotoxin remained intact when exposed to a reducing perfusion buffer containing 1 mM GSH (data not shown). Thus, reduced PDI displaced CTA1 from the CT holotoxin. This effect was documented with a range of PDI concentrations, and the amount of reduced PDI in the perfusion buffer was directly related to the kinetics of holotoxin disassembly (supplemental Fig. S2).

To determine the fate of the CTA2 polypeptide after holotoxin disassembly, we perfused an anti-KDEL antibody over the PDI-treated holotoxin (Fig. 1B). The anti-KDEL antibody did not generate an increase in the refractive index when applied to the intact CT holotoxin (Fig. 1A), but a positive signal was obtained when the antibody was applied to the PDI-treated holotoxin (Fig. 1B). Removal of the CTA1 subunit thus allowed the anti-KDEL antibody to reach the KDEL tag on the CTA2
subunit that remained associated with the sensor-bound CTB pentamer. Reduced PDI thus displaced CTA1 from the CT holotoxin but did not separate CTA2 from CTB$_5$.

Structure-dependent Interactions between PDI and CTA1—Reduced PDI did not bind to CTA2/CTB$_5$ (supplemental Fig. S3), but it did interact with the isolated CTA1 subunit (Fig. 2). As seen with the CT holotoxin, only reduced PDI could interact with CTA1: neither PDI nor oxidized PDI increased the refractive index when perfused over a CTA1 sensor slide (data not shown). Binding of reduced PDI to CTA1 was temperature-dependent. A strong interaction was detected at 10°C, a weaker interaction was detected at 25°C, and little to no interactions were detected at 33°C or the physiological temperature of 37°C (Fig. 2A). The isolated CTA1 subunit is a thermally unstable protein with a partially perturbed secondary structure and a disordered tertiary structure at 37°C (17). It thus appeared that PDI can interact with the folded conformations of CTA1 present at low temperatures but not with the unfolded CTA1 conformations present at physiological or near-physiological temperatures. Thermal instability in the CTA1 polypeptide is only apparent after dissociation from the holotoxin (17, 37, 38), which explains why reduced PDI could interact with CTA1 when it was present in the CT holotoxin at 37°C (Fig. 1B).

The C-terminal A$_1$s subdomain of CTA1 has been proposed to act as the trigger for ERAD-mediated translocation and to serve as the site for PDI binding (12, 39). However, a CTA1 variant lacking most of the A$_1$s subdomain (CTA1$_{1-166}$) can still move from the ER to the cytosol (31). We found that a CTA1$_{1-168}$ variant could also interact with reduced PDI (Fig. 2B). As with the full-length CTA1, PDI binding to CTA1$_{1-168}$ was temperature-dependent: a strong interaction was detected at 10°C, a weaker interaction was detected at 25°C, and no interactions were detected at 33°C or 37°C (Fig. 2B). A similar pattern of temperature-dependent binding was recorded for the interaction between PDI and CTA1$_{1-133}$, a CTA1 variant lacking both the A$_1$s and A$_1$$_s$ subdomains (Fig. 2C). Increasing the temperature from 10 to 37°C results in a progressive loss of structure for CTA1, CTA1$_{1-168}$, and CTA1$_{1-133}$ (27). PDI thus appeared to recognize a folded region in the A$_1$s subdomain of CTA1. The CTA1$_{1-133}$ construct is less stable than either CTA1 or CTA1$_{1-168}$ (27), which explains the weaker binding of PDI to CTA1$_{1-133}$ at 25°C in comparison to CTA1 and CTA1$_{1-168}$.

These experiments highlighted a unique aspect of PDI-CTA1 interactions, as PDI usually binds to unfolded but not folded protein conformations (19, 20).

When exposed to mildly acidic pH, the isolated CTA1 subunit retains a substantial amount of its native structure at physiological temperature (27). Consistent with this observation, we found that an acidic perfusion buffer allowed reduced PDI to effectively bind free CTA1 at 37°C (Fig. 3A). This again suggested that PDI recognizes a structure in the folded conformation of CTA1. It also demonstrated, along with the holotoxin study of Fig. 1B, that reduced PDI was functional in our SPR system at 37°C. Interestingly, PDI was rapidly displaced from CTA1 when the acidic buffer was replaced with neutral buffer (Fig. 3B). The high affinity interaction between CTA1 and PDI that occurred at pH 6.5 was completely lost, as reflected by the return of the signal to baseline value, when the 37°C buffer was switched to neutral pH. The disruption of PDI binding occurred despite the continued presence of PDI in the pH 7.0 perfusion buffer. Furthermore, the loss of PDI interaction was not simply time-dependent: when the perfusion buffer was maintained at pH 6.5, PDI remained stably associated with CTA1 (Fig. 3A). CTA1 would shift from a folded to an unfolded conformation when the buffer was switched from pH 6.5 to pH 7.0. It therefore appeared that the thermal unfolding of CTA1 results in displacement of bound PDI.

To ensure that our pH data resulted from structural alterations to CTA1, we repeated the PDI/CTA1 binding assay with another condition that stabilizes the structure of CTA1: 10% glycerol. Like acidic pH, treatment with 10% glycerol increases the thermal stability of CTA1 and allows the toxin to maintain a substantial amount of its native structure at 37°C (26). Glycerol treatment also allowed reduced PDI to effectively bind CTA1 at 37°C (Fig. 3C). This interaction was lost when glycerol, but not PDI, was removed from the perfusion buffer (Fig. 3D). In contrast, PDI remained stably associated with CTA1 when 10% glycerol was kept in the perfusion buffer for the duration of the experiment (Fig. 3C). CTA1 would shift from a folded to an unfolded conformation when glycerol was removed from the 37°C perfusion buffer. Thus, two independent experimental conditions both indicated that toxin unfolding will displace PDI from its CTA1 binding partner.
**PDI-mediated Disassembly of the CT Holotoxin Does Not Require CTA1 Unfolding**—PDI is thought to remove reduced CTA1 from CTA2/CTB5 by unfolding the holotoxin-associated A1 subunit (9). According to this model, the resulting conformational change in CTA1 would disrupt its non-covalent interactions with CTA2/CTB5 and thereby promote its dissociation from the holotoxin. To determine if PDI could remove CTA1 from CTA2/CTB5 without unfolding CTA1, we added reduced PDI to a CT-coated sensor slide at pH 6.5 (Fig. 4 A). The stable interaction between PDI and CTA1 at 37 °C and pH 6.5 (Fig. 3 A) indicated that PDI does not unfold CTA1 under these conditions—otherwise, as shown in Fig. 4 B, CTA1 unfolding would displace the bound PDI. At both pH 7.0 (Fig. 1 B) and pH 6.5 (Fig. 4 A), reduced PDI could displace CTA1 from the CT holotoxin. Likewise, reduced PDI could displace CTA1 from the CT holotoxin in the presence of 10% glycerol (26). It therefore appeared that PDI can dislodge CTA1 from CTA2/CTB5 without unfolding the A1 subunit. To further examine this possibility, we added PDI to the CT holotoxin at 10 °C or 4 °C. PDI does not induce CTA1 to assume a protease-sensitive conformation at 4 °C, and this has been interpreted to mean that PDI cannot unfold CTA1 at low temperatures (29). However, PDI could still promote the dissociation of CTA1 from CTA2/CTB5 at either 10 °C (Fig. 4 B) or 4 °C (supplemental Fig. S4). This again demonstrated that CTA1 unfolding was not required for its dissociation from the CT holotoxin.

Heating the reduced CT holotoxin to 65 °C, a temperature that results in irreversible denaturation of the holotoxin-associated CTA1 subunit (37), did not result in the displacement of CTA1 from CTA2/CTB5 (supplemental Fig. S5). It therefore appeared that unfolding alone is insufficient to displace CTA1 from CTA2/CTB5. Collectively, these results suggested that PDI removes CTA1 from CTA2/CTB5 by a mechanism that does not involve unfolding of the holotoxin-associated CTA1 subunit.

To determine the fate of PDI after it facilitates the separation of CTA1 from CTA2/CTB5, we repeated the disassembly assay with a holotoxin that was appended to the SPR sensor slide with a rabbit anti-CTA antibody. CTA2, CTB5, and PDI were absent from the sensor slide after PDI-mediated disassembly of the holotoxin at 37 °C (Fig. 5). Use of an anti-CTA monoclonal antibody confirmed that CTA1 was still on the plate. This experiment demonstrated that PDI does not remain associated with CTA1 after holotoxin disassembly.

**PDI Does Not Unfold the CTA1 Polypeptide**—To directly examine the effect of PDI binding on the conformational state...
of CTA1, we performed structural studies using isotope-edited FTIR spectroscopy. This method allows the conformation of a \(^{13}\text{C}\)-labeled protein to be monitored in the presence of a second unlabeled protein. \(^{13}\text{C}\) labeling does not alter the conformation of a protein. However, the heavier nuclear mass of the stable \(^{13}\text{C}\) isotope generates a spectral downshift which allows the FTIR spectrum of a \(^{13}\text{C}\)-labeled protein to be resolved from the spectrum of an unlabeled protein (32, 40). The FTIR spectra of \(^{13}\text{C}\)-labeled CTA1 accordingly exhibited a 50 cm\(^{-1}\) downshift in comparison to the spectra of unlabeled CTA1 (supplemental Fig. S6 A).

As expected, no structural differences between labeled and unlabeled CTA1 were detected by CD over a range of temperatures (supplemental Fig. S6, B and C). The down-shifted FTIR spectra of \(^{13}\text{C}\)-labeled CTA1 could be readily distinguished from the spectra of PDI (Fig. 6), allowing us to directly examine the impact of PDI on CTA1 structure.

Analysis of the 10 °C FTIR spectrum from CTA1 alone indicated that the folded toxin contains 37% \(\alpha\)-helical and 45% \(\beta\)-sheet content (Fig. 7, A and B, Table 1). This result was consistent with the 36% \(\alpha\)-helix/51% \(\beta\)-sheet content predicted from the crystal structure of the folded, holotoxin-associated CTA1 subunit (41). The \(\alpha\)-helical, \(\beta\)-sheet, and irregular (often referred to as “random”) structures of CTA1 were unaltered in the presence of PDI (Fig. 7, C and D, Table 1). Thus, the interaction between PDI and CTA1 at 10 °C which displaces CTA1 from the CT holotoxin (Fig. 4 B) did not result in any significant disordering of the CTA1 polypeptide. In contrast, because of its intrinsic thermal instability, heating CTA1 alone at neutral pH (supplemental Fig. S7) shifted the percentage of its irregular structure from 8% at 10 °C to 53% at 37 °C (Table 1). The gain in irregular structure was accompanied by a loss of both \(\alpha\)-helical and \(\beta\)-sheet structure. These observations were consistent with previous CD measurements that documented the partially dis-
ordered conformation of CTA1 at 37 °C (17, 26, 27). Our FTIR data thus demonstrated that physiological temperature alone has a dramatic disordering effect on CTA1 structure, while the interaction with PDI exerts little effect on the conformation of CTA1.

To determine if the putative unfoldase activity of PDI could be detected at temperatures greater than 10 °C, we performed additional experiments in which reduced PDI was added to CTA1 at 30 °C (supplemental Fig. S8) or to an acid-stabilized CTA1 subunit at 37 °C (supplemental Fig. S9). Although a temperature of 30 °C does not promote an optimal interaction between PDI and CTA1 (Fig. 2), we chose this temperature because most of the CTA1 protease sensitivity assays have been performed at 30 °C (9, 13, 23, 29). PDI and CTA1 do not nor-

### TABLE 1

PDI does not induce unfolding of the CTA1 polypeptide

Deconvolution of the conformation-sensitive amide I bands from FTIR data presented in Figs. 7, S7, S8, and S9 was used to calculate, as detailed in supplemental data, the percentages of CTA1 structure under various conditions. The means ± standard deviations from four separate curve-fitting iterations are shown.

| Condition | % of CTA1 structure |
|-----------|---------------------|
|           | α-Helix | β-Sheet | Irregular | Other |
| 10 °C     | 37 ± 1  | 45 ± 3  | 8 ± 2     | 9 ± 2  |
| 10 °C + PDI | 36 ± 1  | 44 ± 2  | 8 ± 1     | 10 ± 2 |
| 30 °C     | 34 ± 4  | 35 ± 2  | 29 ± 4    | 12 ± 2 |
| 30 °C + PDI | 27 ± 2  | 38 ± 3  | 25 ± 2    | 10 ± 1 |
| 37 °C pH 6.5 | 29 ± 2  | 37 ± 4  | 25 ± 3    | 9 ± 2  |
| 37 °C pH 6.5 + PDI | 24 ± 2  | 35 ± 3  | 30 ± 1    | 11 ± 4 |
| 37 °C     | 21 ± 2  | 15 ± 4  | 53 ± 2    | 11 ± 4 |

FIGURE 7. Structure of CTA1 in the absence or presence of PDI as evaluated by FTIR at 10 °C. Curve-fitting (A and C) and second derivatives (B and D) for the FTIR spectrum of 13C-labeled CTA1 recorded in the absence (A and B) or presence (C and D) of PDI are shown. In panels A and C, the dotted line represents the sum of all deconvoluted components (solid lines) from the measured spectrum (dashed line). Note that different wavenumber scales are used in panels A and B than in panels C and D.
Role of PDI in Cholera Intoxication

Our collective data suggested that PDI is only needed to displace CTA1 from CTA2/CTB subunit; the dissociated CTA1 subunit can then unfold spontaneously to trigger its ERAD-mediated dislocation. To test this model, we performed a transfection-based CTA1 toxicity assay in the TZM, TZM 1–2, and TZM 5–1 cell lines (Fig. 8B). A plasmid-based system was used to express CTA1 directly in the ER of transfected cells (35). cAMP levels were then quantified at 3 h post-transfection. To control for possible differences in transfection efficiencies between the cell lines, CTA1 was radiolabeled and immunoprecipitated from parallel sets of transfected cells. Results from the cAMP assay were subsequently standardized to CTA1 expression levels. Although TZM 1–2 cells were completely resistant to exogenously applied CT holotoxin, they exhibited no resistance to the ER-localized CTA1 construct. In fact, the cAMP levels in TZM 1–2 cells expressing ER-localized CTA1 were slightly elevated in comparison to the cAMP levels generated from CTA1 expression in either the parental TZM cells or TZM 5–1 control cells. Collectively, this data indicated that PDI function is not required for ER-localized CTA1 to enter the cytosol.

Previous work with metabolically labeled cells has demonstrated that the entire detectable pool of plasmid-expressed CTA1 is inserted into the ER (44). Yet, if the co-translational targeting of our ER-localized CTA1 construct was not completely efficient, then the CTA1 toxicity assay would be compromised by the small fraction of CTA1 that was synthesized in the cytosol rather than in the ER. To control for this possibility, we directly monitored the ER-to-cytosol translocation of transfected CTA1 in the parental TZM, TZM 1–2, and TZM 5–1 cell lines. Transfected, radiolabeled cells were partitioned into separate membrane and cytosolic fractions according to established protocols (26, 29, 34, 45). Material immunoprecipitated from each fraction with an anti-CTA antibody was then resolved by SDS-PAGE and quantified using PhosphorImager analysis. As shown in Fig. 8C, approximately equivalent amounts of CTA1 were recovered from the cytosolic fractions of all three cell lines. Thus, the lack of PDI in TZM 1–2 cells did not inhibit the ER-to-cytosol export of transfected CTA1. The results of Fig. 8 demonstrated that PDI is required for productive intoxication with exogenously applied CT but is not needed.
Role of PDI in Cholera Intoxication

for translocation of the isolated CTA1 subunit. As such, it appears that the only role of PDI in cholera intoxication is to displace CTA1 from the CT holotoxin.

DISCUSSION

For CTA1 translocation, PDI has been proposed to act as a redox-dependent chaperone that actively unfolds the holotoxin-associated CTA1 subunit to separate CTA1 from the rest of the toxin (9). The unfoldase activity of PDI in this process is, to the best of our knowledge, a unique property that has not been reported for PDI interactions with any other substrate. Furthermore, the proposed unfoldase activity of PDI is based upon a protease sensitivity assay that only serves as an indirect measure of protein folding (9, 13, 28, 29). In this work, we demonstrated that PDI is required for disassembly of the CT holotoxin but does not unfold the CTA1 subunit. CTA1 unfolds spontaneously after its dissociation from CTA2/CTB, at physiological temperature (17, 26). PDI-mediated disassembly of the CT holotoxin thus leads to the spontaneous unfolding of dissociated CTA1, but PDI itself does not function as an unfoldase to actively unfold CTA1.

Using SPR, we were able to monitor the PDI-mediated disassembly of the CT holotoxin in real time. The kinetics of holotoxin disassembly were linked to the available concentration of PDI (supplemental Fig. S2). Holotoxin disassembly required the reduced form of PDI and did not occur in the presence of reducing agent alone (Fig. 1). These results were consistent with previous reports (9, 36), as was the observed retention of CTA2 with CTB, after release of CTA1 from the holotoxin (Fig. 1B) (46). We could thus faithfully reconstitute the process of CT disassembly on an SPR sensor slide.

Conditions that block CTA1 unfolding in vitro also block the ER-to-cytosol export of CTA1 in vivo (26, 27). However, conditions that block CTA1 unfolding do not block CTA1 dissociation from the CT holotoxin: reduced PDI could separate CTA1 from CTA2/CTB, in the presence of acidic pH (Fig. 4A) or 10% glycerol (26). Secretion of the free A1 subunit from glycerol-treated cells (26) further suggested that in vivo unfolding of CTA1 is unnecessary for toxin disassembly: only folded proteins exit the ER for secretory transport, so the secreted pool of CTA1 must have been in a folded conformation. CT disassembly could also occur at 4 °C (supplemental Fig. S4), a temperature that maintains CTA1 in a folded conformation and prevents the PDI-induced shift of CTA1 to a protease-resistant conformation (29). Collectively, these results strongly suggested that CTA1 unfolding is not required for CTA1 dissociation from the rest of the toxin. In further support of this interpretation, we demonstrated that the thermal denaturation of reduced but holotoxin-associated CTA1 did not displace CTA1 from its non-covalent assembly in the CT holotoxin (supplemental Fig. S5). A similar observation was previously made by Goins and Friere with the technique of differential scanning calorimetry (37). Thus, PDI apparently removes CTA1 from CTA2/CTB, by a mechanism that does not involve unfolding of the holotoxin-associated CTA1 subunit.

To directly examine the putative unfoldase activity of PDI, we employed the structural technique of isotope-edited FTIR spectroscopy. Experiments were performed under three conditions that facilitate PDI-CTA1 interactions and support PDI-mediated disassembly of the CT holotoxin: 10 °C, 30 °C, and 37 °C at pH 6.5. For each condition, deconvolution of the conformation-sensitive amide I bands demonstrated that PDI did not substantially alter the percentage of irregular structure in CTA1 and, thus, did not unfold the CTA1 polypeptide (Table 1). In contrast, heating CTA1 alone at neutral pH shifted the percentage of its irregular structure from 8% at 10 °C to 53% at 37 °C at the expense of both α-helix and β-sheet structures (Table 1). CT disassembly therefore occurs under conditions that do not unfold substantial PDI-induced disordering of the CTA1 polypeptide. These results, which represent the first direct examination of the impact of PDI binding on CTA1 structure, demonstrate that CTA1 is unfolded by physiological temperature (due to its intrinsic thermal instability) but not by PDI.

The unfoldase activity of PDI has only been described with a biochemical protease sensitivity assay. Using this method, the folding state of a protein is inferred from its susceptibility to proteolysis: a folded protein is generally more resistant to proteolysis than an unfolded variant of the same protein. The resistance of CTA1 to trypsin-mediated proteolysis, combined with the sensitivity of PDI-treated CTA1 to proteolysis, thus yielded the conclusion that PDI actively unfolds the CTA1 subunit. Unfortunately, interpretation of these experiments is complicated by the conformational shift in CTA1 that occurs upon its separation from the holotoxin and by the unstable, heat-labile nature of the free A1 subunit (17). Protease concentration can also affect the outcome of the experiment, as can the temperature of the experiment (17, 26, 28). It should be noted that most experiments to monitor the "unfoldase" activity of PDI have been performed at 30 °C (9, 13, 23, 29), but incubation of CTA1 alone at 37 °C will induce the toxin to assume a protease-sensitive conformation (17, 26) (supplemental Fig. S10). Furthermore, the sensitivity of PDI-treated CTA1 to proteolysis depends upon the protease used: the toxin is more sensitive to trypsin (9) than to the thermolysin protease which cleaves bulky and aromatic residues (supplemental Fig. S10). These issues suggest that protease sensitivity is a poor, indirect measure of the purported conformational change required for CTA1 dissociation from the CT holotoxin. More important, however, is the disconnect between the PDI-induced shift of CTA1 to a trypsin-sensitive conformation and the PDI-mediated disassembly of the CT holotoxin. PDI does not induce CTA1 to assume a protease-sensitive conformation at 4 °C (29), yet PDI can still displace CTA1 from CTA2/CTB, at 4 °C (supplemental Fig. S4). These results demonstrate that the PDI-induced shift of CTA1 to a protease-sensitive conformation, which has been interpreted as an unfolding event (9, 13, 23, 28, 29), does not correlate to the PDI-induced separation of CTA1 from CTA2/CTB,.

PDI binds to folded but not unfolded conformations of CTA1 (Figs. 1–3). The spontaneous unfolding of CTA1 that occurs upon holotoxin disassembly at 37 °C thus displaces the toxin-bound PDI (Figs. 3 and 5). This observation is consistent with a previous report that concluded PDI-substrate interactions can be disrupted by changes to the structure of the substrate (24). However, our results conflict with the model of CTA1-PDI
interactions that proposes PDI oxidation by Ero1p is responsible for displacing toxin-bound PDI (23). That model is based upon studies performed at temperatures below 37 °C. While Ero1p clearly regulates the redox status of PDI (22, 23, 47, 48), our results strongly suggest that Ero1p function is not required to release PDI from CTA1 at physiological temperature.

As demonstrated with cell-based assays, CTA1 translocation does not require PDI function after disassembly of the holotoxin. PDI-deficient cell lines were completely resistant to exogenously applied CT, as expected from its essential role in separating CTA1 from CTA2/CTB5. However, the same PDI-deficient cells displayed wild-type sensitivity to a CTA1 construct that was directly expressed in the ER of transfected cells. This experimental condition, which mimicked the status of ER-deficient cells, displayed wild-type sensitivity to a CTA1 construct that was directly expressed in the ER of transfected cells. Our results strongly suggest that Ero1p function is not required for displacing toxin-bound PDI (23). That model is based upon studies performed at temperatures below 37 °C. While Ero1p clearly regulates the redox status of PDI (22, 23, 47, 48), our results strongly suggest that Ero1p function is not required to release PDI from CTA1 at physiological temperature.

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Role of PDI in Cholera Intoxication

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Protein-disulfide Isomerase Displaces the Cholera Toxin A1 Subunit from the Holotoxin without Unfolding the A1 Subunit
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