Properties of *Bacillus cereus* hemolysin II: A heptameric transmembrane pore

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(RECEIVED February 5, 2002; FINAL REVISION April 24, 2002; ACCEPTED April 24, 2002)

Abstract

The gene encoding hemolysin II (HlyII) was amplified from *Bacillus cereus* genomic DNA and a truncated mutant, HlyII(ΔCT), was constructed lacking the 94 amino acid extension at the C terminus. The proteins were produced in an *E. coli* cell-free in vitro transcription and translation system, and were shown to assemble into SDS-stable oligomers on rabbit erythrocyte membranes and liposomes. The hemolytic activity of HlyII was measured with rabbit erythrocytes yielding an HC50 value of 1.64 ng mL⁻¹, which is over 15 times more potent than staphylococcal α-hemolysin. HlyII(ΔCT) was about eight times less potent than HlyII in this assay. Limited proteolysis of the oligomers formed by HlyII and HlyII(ΔCT) on red cell membranes showed that the C-terminal extension is sensitive to digestion, while HlyII(ΔCT) is protease resistant and migrates with an electrophoretic mobility similar to that of digested HlyII. HlyII forms moderately anion selective, rectifying pores (I⁺/I⁻ = 0.57, 1 M KCl, pH 7.4) in planar lipid bilayers of diphytanoylphosphatidylcholine with a unitary conductance of 637 pS (1 M KCl, 5 mM HEPES, pH 7.4) and exhibits no gating over a wide range of applied potentials (−160 to +160 mV). In addition, it was demonstrated that HlyII forms a homoheptameric pore by using gel shift electrophoresis aided by a genetically encoded oligoaspartate tag. Although they share limited primary sequence identity (30%), these data confirm that HlyII is a structural and functional homolog of staphylococcal α-hemolysin.

Keywords: β-Barrel; hemolysin; membrane protein; pore-forming toxin; staphylococcal α-hemolysin; subunit stoichiometry

In this work, we examine the properties of hemolysin II (HlyII), a β-barrel pore forming toxin (β-PFT) from *Bacillus cereus*. The β-PFTs consist of several subfamilies of bacterial exotoxins that are related by sequence (Gouaux et al. 1997; Menestrina et al. 2001) and structure (Song et al. 1996; Gouaux et al. 1997; Olson et al. 1999; Pédelacq et al. 1999). These polypeptides are secreted as water-soluble molecules that bind to the surfaces of susceptible cells and assemble into oligomeric transmembrane pores leading to cell permeation and lysis (Bhakdi et al. 2000; Menestrina et al. 2001). The crystal structure of a heptameric staphylococcal α-hemolysin (αHL) pore has been determined in detergent at 1.9-Å resolution (Song et al. 1996). αHL has also been shown to form heptamers on red cell membranes (Gouaux et al. 1994), in planar lipid bilayers (Krasilnikov et al. 2000), supported bilayers (Fang et al. 1997), and after spontaneous assembly in solution (Cheley et al. 1997). In a...
working scheme for the assembly of αHL, the 293-residue polypeptide first binds to the membrane as a monomer, associates to form a heptameric prepore, and finally inserts into the bilayer to form the transmembrane pore. This scheme is supported by numerous biophysical and biochemical experiments (Cheley et al. 1997), and has been refined to accommodate recent structural data (Olson et al. 1999; Pédelacq et al. 1999).

Bacillus cereus is an opportunistic pathogen. It is associated with a wide range of clinical symptoms, but encountered primarily in cases of severe food poisoning (Drobeniuk 1993; Lund et al. 2000). Over 20 exotoxins are produced and secreted by B. cereus, including several nonhemolytic enterotoxins (Alouf and Freer 1999). Two distinct hemolytic proteins produced by this bacterium, hemolysin II (HlyII) (Baida et al. 1999) and cytotoxin K (CytK) (Lund et al. 2000), have recently been cloned and shown to be homologous with the β-PFTs. HlyII has the longest polypeptide chain in the β-PFT family, with 412 residues, and contains a C-terminal 94 amino acid extension that has no homology with any other known β-PFT. The remainder of HlyII shares 30% sequence identity with αHL, and the level of identity with other members of the β-PFT family does not exceed this value. Although regions of similarity are dispersed throughout the aligned sequences (Fig. 1A), the majority are concentrated in the strands that comprise the “cap” domain (Song et al. 1996), and are probably necessary to preserve the fold of the cap (Fig. 1B).

Members of the β-PFT family other than αHL have not been thoroughly investigated. In particular, limited data exist on the molecular architecture of the pores formed by them. The bicomponent leukocidin pore has a relatively large unitary conductance (2.5 nS in 1 M KCl) compared to other members of the β-PFT family (Miles et al. 2001), and it has been recently demonstrated to form an octameric transmembrane pore (Miles et al. 2002). The clostridial β-toxin has been shown to oligomerize into an SDS-stable multimer of unknown composition on human endothelial cells (Steinthorsdottir et al. 2000), and forms cation-selective channels with two conductance states of 60 and 110 pS (100 mM NaCl, 10 mM HEPES, pH 7.4, +60 mV) (Shatsky et al. 2000). Recently, it was shown that CytK purified

Fig. 1. Comparison of Bacillus hemolysin II and staphylococcal α-hemolysin. (A) Primary sequence alignment of Bacillus cereus hemolysin II (HlyII) and staphylococcal α-hemolysin (αHL). Residues highlighted with a blue background are identical, while similar residues (Blosum62 similarity scoring matrix; Henikoff and Henikoff 1992) are shown as blue characters. The 94 residue C-terminal extension of HlyII is shown in maroon characters; it is the portion that was deleted to form the truncation mutant HlyII(ΔCT). The figure was generated using ClustalW 1.81 (Thompson et al. 1994) and rendered using ESPript 2.0 (Gouet et al. 1999). (B) Structure of one protomer taken from the crystal structure of the αHL heptamer (7aHL.pdb). Areas in dark blue and light blue correspond to identical and similar residues, respectively, as shown in (A). The image was created with SPOCK 6.3 (Christopher 1998) and rendered with Molascript (Kraulis 1991).
from *B. cereus* supernatants forms pores in planar lipid bilayers, and is cytotoxic towards intestinal epithelial cells (Hardy et al. 2001). The subunit stoichiometries of the *Bacillus* toxins, HlyII and CytK, have not been determined.

In this study, we demonstrate that HlyII, produced by in vitro transcription and translation, forms a heptameric transmembrane pore in red cell membranes, which is resistant to SDS. In planar lipid bilayers, the pores are rectifying and lack voltage-induced gating. HlyII with the C-terminal extension removed, HlyII(ΔCT), has similar properties. Knowledge of the subunit stoichiometry and channel properties of this relative of αHL adds to the understanding of the β-PFTs family, and will be helpful in protein engineering aimed at the construction of pore-forming proteins with new properties (Bayley 1999; Bayley and Cremer 2001).

**Results and Discussion**

**HlyII and HlyII(ΔCT) form SDS-resistant oligomers on red cell membranes and liposomes**

HlyII includes a 94-amino acid C-terminal extension, which is not homologous with any known β-PFT. The two most similar candidates from a BLAST search (Altshul et al. 1997) were a 46-amino acid segment of the pX01–124 gene product from *Bacillus anthracis* (39% identity, 67% similarity, one gap) and a 78-amino acid segment of orf16 from *Streptococcus* phage Cp-1 (34% identity, 49% similarity, three gaps). The genes for these proteins are both associated with genetic material linked with virulence, but the roles of the proteins have not been fully defined (Welkos 1991; Martín et al. 1996). We constructed a truncation mutant, HlyII(ΔCT), which lacks the extension (residues 289–382) (Fig. 1A). 35S-labeled HlyII and HlyII(ΔCT) were produced in an *E. coli* S30 transcription and translation system. When HlyII was translated in the presence of rabbit red cell membranes (rRBCM) or incubated with small unilamellar vesicles (SUVs) composed of egg yolk phosphatidylcholine, cholesterol, and phosphatidic acid at a molar ratio of 55:25:20, a single high-molecular mass band appeared upon SDS-polyacrylamide gel electrophoresis of unheated samples (Fig. 2). The extent of oligomerization on rRBCM is comparable to that of αHL, with 74% of the total membrane-bound protein in the oligomeric form (compared to 83% seen with αHL). In

![Fig. 2. (a) Autoradiogram of 10% SDS-polyacrylamide gel. Lane 1, HlyII translated in absence of rRBCM; lane 2, HlyII(ΔCT) translated in absence of rRBCM; lane 3, HlyII translated in presence of rRBCM; lane 4, HlyII(ΔCT) translated in presence of rRBCM; lane 5, HlyII incubated in presence of freshly prepared liposomes for 1 hr at room temperature; lane 6, HlyII(ΔCT) incubated in presence of liposomes. In the samples used in lanes 3 and 4, the rRBCM were washed before electrophoresis of the bound protein. In the samples used in lanes 5 and 6, the entire sample was loaded onto the gel. The high molecular weight bands (arrows) represent oligomerized HlyII and HlyII(ΔCT).](image)
the case of HlyII(ΔCT), the extent of oligomerization is reduced to 49%. Oligomers formed by HlyII and HlyII-(ΔCT) are stable in 2.3% SDS (1× Laemmli sample buffer) at room temperature and dissociate at 82°C and 78°C, respectively. αHL dissociates at 70°C in the sample buffer (data not shown).

A deletion variant lacking the extension (and two additional amino acids) was previously shown to retain some hemolytic activity towards human red blood cells (Baida et al. 1999). The removal of the two additional amino acids is expected to lead to a reduction in activity. We found earlier that the removal of three amino acids from the C terminus of αHL causes a dramatic reduction in hemolytic activity that is associated with an almost complete loss of ability to form SDS-stable oligomers (Walker et al. 1992a).

Examination of the HlyII oligomer by limited proteolysis suggests structural similarity to αHL

The susceptibility of HlyII and HlyII(ΔCT) to cleavage by proteinase K was compared with that of αHL (CT) (Fig. 3). The HlyII oligomer shows marked susceptibility to proteinase K, yielding multiple proteolytic fragments at low protease concentrations (5 μg mL⁻¹; Fig. 3, lane 2). In contrast, the oligomeric form of HlyII(ΔCT), like the αHL heptamer, was protease resistant. At the highest concentration of proteinase K (500 μg mL⁻¹), a protease-resistant, SDS-stable form of oligomeric HlyII was formed, which migrated in SDS gels with a similar mobility to oligomers formed from the genetically truncated HlyII(ΔCT) (Fig. 3). These results suggest that the C-terminal extension either has a structure with pronounced sensitivity to proteolytic cleavage, or that the extension is connected to the rest of the protein through a readily accessible linker. The former is favored because the C terminus (TL, see below) expressed as a separate domain is protease sensitive (data not shown). In any case, the stability of the proteolysed pore implies that it is unlikely that the extension contributes significantly to the stability of the cap domain, the formation of which initiates heptamerization at the prepore stage (Cheley et al. 1997). In addition, when the sequences of HlyII from several different B. cereus and B. thuringiensis strains, BGSC 4A1, 4A4, 4A7, 6A3, and 6E2, were compared to the HlyII sequence from Bacillus cereus strain 6A5, the nonredundant sequence differences were concentrated in the C-terminal extension (14 differences in the 94 residues, compared with 9 differences in the 288 residues of the remainder of the polypeptide). This relative lack of conservation again suggests that the extension does not play a major role in oligomerization.

Hemolysin II is a potent hemolytic toxin

The hemolytic activities of HlyII and HlyII(ΔCT), produced by IVTT, were compared with αHL in a quantitative assay. The concentration of HlyII at which 50% of the rabbit erythrocytes were lysed in 1 h at 20°C (HC₅₀) was 1.64 ng mL⁻¹ (Fig. 4A). By contrast, αHL gives an HC₅₀ value of 25 ng mL⁻¹ at 37°C (Walker et al. 1992b) and a similar value at 20°C (unpublished results). The HC₅₀ value for CytK is similar to that of αHL (Lund et al. 2000). Therefore, the specific activity of HlyII is more than 15 times greater than that of αHL or CytK. The HC₅₀ value amounts to about 1000 monomers per cell. HlyII(ΔCT) had a lower specific activity of 4.8 ng mL⁻¹ (Fig. 4B), in keeping with the somewhat less efficient oligomerization seen with this mutant (Fig. 3). The C-terminal extension by itself (TL) showed no hemolytic activity (Fig. 4B), and did not affect the activity of HlyII(ΔCT) or HlyII. We have found that other β-PFTs can accommodate large extensions at their C termini. For example, the mutant αHL-TL was constructed in which the Bacillus tail was spliced genetically onto the C terminus of αHL. The HC₅₀ of αHL-TL was 64 ng mL⁻¹, i.e. αHL-TL was 2.5 times less active than wild type αHL. In the case of the F and S subunits of leukocidin, the hemolytic activities were unchanged when TL was fused to the C termini (Miles et al. 2002). The cytotoxic effects of HlyII against other cell lines has not been examined. It has been shown recently that CytK has potent activity against human intestinal epithelia, and is believed to have caused necrotic enteritis in a food poisoning outbreak resulting in several deaths (Lund et al. 2000; Hardy et al. 2001).

Fig. 3. Conformational states of HlyII, HlyII(ΔCT) and αHL examined by limited proteolysis. HlyII, HlyII(ΔCT) and αHL were translated in the presence of rRBCM. The membranes were washed, treated with proteinase K, solubilized and subjected to electrophoresis in a 10% SDS-polyacrylamide gel prior to autoradiography. Lanes 1–4, HlyII; lanes 5–8, HlyII(ΔCT); lanes 9–12, αHL. The final proteinase K concentrations were: lanes 1, 5, and 9, 0 μg mL⁻¹; lanes 2, 6, and 10, 5 μg mL⁻¹; lanes 3, 7, and 11, 50 μg mL⁻¹; lanes 4, 8, and 12, 500 μg mL⁻¹.
Properties of Bacillus cereus hemolysin II

Fig. 4. Quantitative hemolysis assays with rabbit erythrocytes. (A) One-hour activity assays on HlyII and αHL synthesized by IVTT. The first well in each panel contained IVTT mix diluted to a final volume of 100 μL in MBSA. The concentration of αHL in an IVTT mix was determined as previously described (Walker et al. 1992b; Miles et al. 2001) and used to calculate the concentration of HlyII produced by IVTT (see Materials and Methods). On this basis, the final concentrations of HlyII or αHL were made equal in the first well (0.21 μg/mL). Twofold serial dilutions from left to right were then made. The final concentration of rRBC in all wells was 0.5%. The arrows indicate ~50% lysis. (B) One-hour activity assays on various constructs synthesized by IVTT. IVTT mixes containing HlyII, HlyII(ΔCT), αHL, and αHL-TL were diluted 10-fold with MBSA before addition (1 μL) to the first well. TL, the C-terminal extension of HlyII was translated as a separate polypeptide, and added where indicated in 32-molar excess as determined by phosphorimager quantitation (5 μL of undiluted IVTT mix in the first well). Twofold serial dilutions were made from left to right.

By comparison with αHL, there is a shorter initial lag period associated with HlyII hemolysis (Fig. 4): HlyII, 5 min to 5% lysis at HC_{50}; αHL, 18 min to 5% lysis at HC_{50}. The short lag period for HlyII suggests that the overall rate of pore formation (membrane binding, oligomerization, and membrane penetration) is greater than that of αHL. By contrast, the rates of hemolysis after the lag period were similar at comparable effective concentrations. Further investigation will be necessary to better define the origin of the difference in lag time.

HlyII forms ionic channels in planar lipid bilayers

Upon insertion into planar lipid bilayers, the gel-purified oligomeric form of HlyII obtained by IVTT (Fig. 2) produced discrete conductance steps of 25.5 pA in 1 M KCl at pH 7.4 at a transmembrane potential of −40 mV (Fig. 5A). If each step is presumed to correspond to a single ionic channel, the unitary conductance of the HlyII pore is 637 ± 51 pS (n = 180), under the prevailing conditions (Fig. 5A). A similar result was obtained with gel-purified HlyII(ΔCT) oligomers, 636 ± 73 pS (Table 1). By comparison, the conductance of the αHL pore was 775 ± 38 pS, under the same conditions (Miles et al. 2001) and that of CytK 627 ± 14 pS (1 M NaCl, 5 mM HEPES, pH 7.2) (Hardy et al. 2001). Therefore, the functional diameters of the HlyII, CytK and αHL pores are similar.

The HlyII pores show rectification when examined at applied potentials from −160 to +160 mV (I_{+80}/I_{-80} = 0.566) (Miles et al. 2001) (Fig. 5B, Table 1), which is the opposite of that observed with the clostridial β-toxin (Shatursky et al. 2000) (note that these authors use the opposite polarity sign convention in their bilayer recordings). The charge selectivity of the HlyII pore was determined by measuring the reversal potential of single-channel currents in asymmetrical KCl solutions. HlyII forms anion selective pores, with a permeability ratio (P_{Cl−}/P_{K+}) of 0.29 ± 0.03 (1000 mM KCl (cis), 200 mM KCl (trans), pH 7.5) (Fig. 5C). The HlyII pore is significantly more anion selective than the αHL pore (P_{Cl−}/P_{K+} = 0.79, under the same conditions) (Gu et al. 2000), whereas the leukocidin pore is cation selective (P_{K+}/P_{Cl−} = 1.64) (Miles et al. 2001). Of the known β-PFTs, HlyII and CytK share the greatest sequence identity within the putative transmembrane domain (48%). In HlyII, each subunit contributes seven charged residues that are predicted on the basis of their hydrophilicity and a sequence alignment with αHL to face the lumen of the transmembrane barrel (K106, E110, K120, D125, K126, D131, K141). However, only two charged side chains (K128, E139) are predicted to project into the lumen of the CytK pore, which is almost nonselective (Hardy et al. 2001). The overall positive charge in the HlyII pore is likely to contribute to the observed anion selectivity.

In addition, the HlyII pore remains predominantly open at the applied potentials tested (−160 to +160 mV) (Fig. 6). This is similar to what is observed with αHL (Korchev et al. 1995), but in contrast with the properties of most porins and many other β-PFTs, which exhibit voltage-induced gating. For example, the leukocidin pore gates at positive potentials above +60 mV, but remains fully open at negative potentials up to −160 mV (Miles et al. 2001). Two rings of charged residues facing the lumen of the HlyII pore form the boundaries of the putative transmembrane region, E110/K141 and K120/D131, based on the alignment of residues in the stem domain (unpublished results) with those in the αHL β-barrel (Song et al. 1996). We speculate that these side chains form salt bridges that strengthen the β-barrel, holding it in a rigid conformation even under high applied potentials. This argument is reinforced by the ruggedness of the HlyII
oligomer, which dissociates in SDS at 82°C, as it is known that it is barrel formation that stabilizes the H9251 HL heptamer to heat denaturation in SDS (Valeva et al. 1996; Cheley et al. 1999).

Finally, we determined the subunit composition of the HlyII pore by gel shift electrophoresis. Whereas the original gel-assembly of the HlyII pore is a heptamer, the results from gel shift electrophoresis suggest that the subunit composition is not a simple heptamer but rather a complex of heptamers and octamers. The subunit composition of the HlyII pore was determined by gel shift electrophoresis and is shown in Table 1.

**Hemolysin II assembles as a heptamer on membranes**

Table 1. Single-channel properties of wild-type and engineered HlyII pores

<table>
<thead>
<tr>
<th>Protein</th>
<th>g (pS)</th>
<th>n</th>
<th>I_{a0}/I_{b0}</th>
<th>V_r (mV)</th>
<th>P_{K+}/P_{Cl−}</th>
</tr>
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<tbody>
<tr>
<td>HlyII(WT)</td>
<td>637 ± 51</td>
<td>180</td>
<td>0.566 ± 0.004</td>
<td>−18.0 ± 1.3; 0.29 (n = 4)</td>
<td></td>
</tr>
<tr>
<td>HlyII(ΔCT)</td>
<td>636 ± 73</td>
<td>142</td>
<td>0.583 ± 0.001</td>
<td>n.d.</td>
<td></td>
</tr>
<tr>
<td>HlyII(ΔCT)-D8</td>
<td>610 ± 54</td>
<td>273</td>
<td>0.591 ± 0.032</td>
<td>−19.3 ± 1.2; 0.26 (n = 3)</td>
<td></td>
</tr>
<tr>
<td>[HlyII(ΔCT)−D8]_1</td>
<td>642 ± 62</td>
<td>144</td>
<td>0.691 ± 0.077</td>
<td>n.d.</td>
<td></td>
</tr>
</tbody>
</table>

* Recordings were made at −40 mV in 1 M KCl, 5 mM HEPES, pH 7.4. Single-channel conductances were determined by fitting the peaks in amplitude histograms to Gaussian functions (e.g., Figs. 5, 8).
* Rectification ratios were determined from three or more experiments.
* The reversal potentials (V_r) and permeability ratios (P_{K+}/P_{Cl−}) are mean values ± SD determined with the following buffers: cis, 1 M KCl, 5 mM HEPES, pH 7.4; trans, 0.2 M KCl, 5 mM HEPES, pH 7.4. n.d., not determined.
shift method used to evaluate the stoichiometry of the αHL pore was based on a mobility change brought about by site-specific chemical modification (Gouaux et al. 1994; Braha et al. 1997), gel-shift experiments based on genetically engineered truncations or extensions have proved successful in other cases (Heginbotham et al. 1997; Zitzer et al. 1999; Miyata et al. 2001; Miles et al. 2002).

In a similar approach to the one we used here, transthyretin heterotetramers were separated by using an N-terminal charged extension (Flag tag) (Schneider et al. 2001). Similarly, hybrid trimers of the catalytic subunits of aspartate transcarbamoylase were resolved with a six-residue aspartate tail (Sakash and Kantrowitz 2000). The mutant HlyII-D8 was constructed in which an extension encoding eight aspartate residues was incorporated at the C terminus of HlyII (Fig. 7A). Oligomers containing various ratios of HlyII to HlyII-D8 were produced by cotranslation in the presence of rRBCM. Analysis of the oligomers by SDS-polyacrylamide gel electrophoresis and autoradiography lacked sufficient resolution to enable a count of individual bands (Fig. 7B). It is unclear why the D8 tail on full-length HlyII fails to yield sharp bands; it is possible that the various permutations of each combination of subunits (Braha et al. 1997) are spread out on the gel. By contrast, eight distinct bands were obtained by using various mixtures of αHL and αHL-D8 (Fig. 7C). Each downward shift in electrophoretic mobility corresponds to the incorporation of one αHL-D8 subunit into the αHL heptamer. Therefore, we engineered the D8 tail onto HlyII(ΔCT). When assembled with HlyII(ΔCT), eight species were resolved by gel electrophoresis (Fig. 7D). This result indicates that the HlyII(ΔCT) oligomer contains seven subunits.

Because the properties of the pores formed by HlyII and HlyII(ΔCT) are similar, HlyII is also likely to form a heptamer. HlyII and HlyII(ΔCT) produced pores with similar conductance values (Fig. 8A,B; Table 1). Together with the proteolysis experiments and measurements of hemolytic activity, these results support the idea that the 94-amino acid C-terminal extension is neither essential for pore formation nor for determining functional properties. Oligomers composed entirely of HlyII(ΔCT)-D8 subunits and a heteromer containing one D8 subunit [HlyII(ΔCT)]6/[HlyII(ΔCT)-D8] also exhibited similar conductance values, rectification properties and charge selectivities when compared with the pores formed by HlyII and HlyII(ΔCT) (Fig. 8A,B; Table 1). Therefore, despite the additional 56 negative charges in [HlyII(ΔCT)-D8]7, the conductive pathway remains unaffected, in keeping with the location of the C terminus distant from the channel entrance.

Several different N- and C-terminal oligoaspartate extensions have been evaluated in our laboratory for their ability to produce electrophoretic shifts in β-PFTs. For instance, a D8 tail provides about 1.5 times the separation between oligomeric species in SDS gels compared with a D4 extension (G. Miles and S. Cheley, unpubl.). Besides its utility in evaluating the subunit composition of proteins, this approach provides a convenient way to separate functional heteromers (Sakash and Kantrowitz 2000; Howorka et al. 2001a, 2001b).

**General implications and future prospects**

Here, we have shown that the properties of the pore-forming protein HlyII from *Bacillus cereus* conforms, in general, with those of other members of the class of β-PFTs. HlyII has the most potent hemolytic activity yet found in the β-PFTs; 50% lysis of rabbit red cells occurs at a ratio of ~1000 HlyII monomers per cell. HlyII possesses additional properties that might be improved for applications in biotechnology (Eroglu et al. 2000; Bayley and Cremer 2001). For example, the heptameric pore is more stable than that formed by αHL, dissociating in SDS at 82°C. Remarkably, the pore formed by clostridial β toxin is yet more stable, resisting “boiling” in SDS (Steinthorsdottir et al. 2000). The pore formed by HlyII remains open at high transmembrane potentials, which is a useful property for applications in...
sensor technology (Bayley et al. 2000; Bayley and Cremer 2001). The results provided here also demonstrate that the tail carried by HlyII has little influence on the properties of the pore. This suggests that the tail might be replaced by polypeptide sequences that provide additional functionality such as the ability to form lattices or bind to surfaces, or a linked catalytic activity. Finally, the use of an oligoaspartate extension provides a convenient means to purify heteromeric pores. Combined with new findings about the subunit–subunit interface, revealed by the ability of the related binary pore-forming toxin, leukocidin, to form octamers (Miles et al. 2002), the present work suggests that it should be possible to extend the range of heteromeric pores that can be formed and purified.

Materials and methods

Isolation of genomic DNA from Bacillus cereus

Bacillus cereus strain 6A5 (Bacillus Genetic Stock Center, Ohio State University) (equivalent to ATCC #14579) was grown to saturation from log-phase inocula and stored frozen at −80°C in...
50% glycerol. To isolate genomic DNA, a portion of a culture (5 mL) grown overnight at 30°C was centrifuged at 3000 g. The pellet was resuspended in buffer B1 (Qiagen #19060; 1 mL, 50 mM Tris.HCl, pH 8.0, 0.5% Tween-20, 0.5% Triton X-100), supplemented with 125 U of mutanolysin (Sigma. #M4782; 25 mL stock made at 5000 U mL⁻¹ in deionized water and stored in frozen aliquots), 0.4 mg proteinase K (GibcoBRL; 20 mL stock made at 20 mg mL⁻¹ in 1× TE [10 mM Tris.HCl, 1 mM EDTA, pH 8.0, supplemented with 10 mM NaCl]) and 0.2 mg RNase A (Qiagen; 100 mg mL⁻¹ stock), and incubated at 37°C overnight. Genomic DNA was purified on an anion-exchange resin using a Qiagen Genomic tip 20/G (#10223) and redissolved in 1× TE. The DNA was then sheared by two passes through an 18-gauge needle.

Amplification of Bacillus hemolysin-II genes

The coding sequences for HlyII were amplified from the genomic DNA by using the following primers: HlyII (sense) 5’-ACATATGGCATCTAAAAAGGAAACTGTAGAAAATC-3’; HlyII (antisense) 5’-CAAGCTTATCAGATCTTTTTTAATCTCAATATAAGG-3’. The sense primer generated an NdeI site (CATATG), containing a new initiation codon, immediately before the first codon of the mature polypeptide, predicted by sequence alignment (Fig. 1A). The antisense primer encodes two stop codons and a HindIII site immediately following the last codon of the gene. PCR was carried out in a 50-µL mixture containing sheared genomic DNA (50 ng), primers (50 pmole of each), 200 µM dNTPs (Stratagene), and 1.5 U of Taq/Pwo DNA polymerase mixture in PCR buffer 1 (Expand Long Template PCR System) with the following program: 95°C for 2 min, 25 cycles of 94°C (60 sec), 45 to 56°C gradient (60 sec), 72°C (80 sec), followed by a final extension at 72°C for 7 min. The PCR product was then cloned into the TOPO-TA plasmid (#K4500–01, Invitrogen) to yield pHlyII-TA.

pHlyII-TA was digested with NdeI and HindIII, and the liberated DNA insert was ligated into the pT7 expression vector pT7-SC1 (Miles et al. 2001). Upon sequencing, two independently amplified clones from Bacillus cereus strain 6A5; the following differences were noted from the published sequence (B. cereus VKM-B771) (Baida et al. 1999): G66S (GGG→AGC), S236T

**Fig. 8.** Representative channel insertion events and single channel conductance histograms for [HlyII]₇, [HlyII(ΔCT)]₇, [HlyII(ΔCT)-D8]₇, and the heteroheptamer [HlyII(ΔCT)]₆[HlyII(ΔCT)-D8]₁. (A) HlyII homoheptamer. (B) HlyII(ΔCT) homoheptamer. (C) [HlyII(ΔCT)]₆[HlyII(ΔCT)-D8]₁ heteroheptamer. After a preparative IVTT reaction in the presence of rRBCM and SDS-polyacrylamide gel electrophoresis, the desired band was excised from the gel and processed as described in Materials and Methods. Current traces were recorded in 1 M KCl, 5 mM HEPES (pH 7.4). The scale for the current trace in (D), applies to the entire figure. The histograms of the current steps represent compilations from at least 15 independent recordings. Conductance values, rectification properties, and ion selectivities are summarized in Table 1.
(TCT→ACT), N276H (AAC→CACC), P294L (CCT→CCTT), I299N (ATT→AAT), G300S (GGT→AGT), N306S (AAC→AGC), N306D (AAC→GAT), Q307K (CAG→AAA), F317L (TTT→CCTT), T358S (ACA→TCA). With the exception of three residues (in bold), the variations noted occurred in the C-terminal extension. In addition, there were 35 silent changes throughout the sequence. The DNA sequence from strain 6A5 has been deposited in GenBank with the accession number AF448485.

Hemolysin-II C-terminal truncation

Unless otherwise noted, the constructs used in this study were created by PCR and cloned into the TOPO-TA plasmid prior to being subcloned into pT7-SC1 by ligation. Each construct was verified by DNA sequencing.

HlyII(ΔCT), a mutant of HlyII truncated at the C-terminus, was constructed by removing the sequence encoding the last 94 amino acids of HlyII (residues 289–382). PCR was carried out on linearized pT7-HlyII(Δ6A5) using the forward primer SC001 5′-CAC TATAGGAGACCAACACCGG-3′ and the reverse primer BAC6 ASTRN1 5′-TAAAGCTTCTTAAGAGTAACTGTAGT-3′. The latter encodes two stop codons and a HindIII site immediately after the Leu-288 codon in the HlyII gene.

The C-terminal extension (TL) was cloned separately into the pT7 vector by using PCR. The sense primer BAC6A5TAILBEGIN 5′-ACATATGGAACCAAAAAGCCCTT-3′ and the reverse primer BAC6 ASTRN1 5′-TAAAGCTTCTTAAGAGTAACTGTAGT-3′. The latter encodes two stop codons and a HindIII site immediately after the Leu-288 codon in the HlyII gene.

Construction of mutants with C-terminal (oligo)–aspartic acid extensions

We sought to develop a genetic alternative to the original gel shift method for counting subunits, which was based on a mobility change brought about by site-specific chemical modification (Gouaux et al. 1994). The mutants HlyII-D8, HlyII(ΔCT)-D8, and αHL-D8 were constructed with an extension encoding eight C-terminal aspartate residues (Fig. 7A). The “D8 tail” was expected to change the electrophoretic mobility of the assembled pore based on charge. PCR was carried out with SC001 as the sense primer, using each of the following reverse primers: HlyII-D8, 5′-AAGCT TATCAATGTCATGGCTATGGCTATGCTGATTCTTAA TCTCAAA-3′; HlyII(ΔCT)-D8, 5′-AAGCTTATCAATGTCATG CTCATGTCATGGCTATGGCTATGCTGATTCTTAA TCTCAAA-3′; αHL-D8, 5′-AAGCTTATCAATGTCATGG CTATGTCATGGCTATGGCTATGCTGATTCTTAA TCTCAAA-3′. These electrophoretically purified antisense oligonucleotides incorporated two stop codons (bold) after the D8 tail codons (underscored) followed by a HindIII site (italics).

Construction of the fusion protein, αHL-TL

In vivo recombination (Howorka and Bayley 1998) was used to fuse a 3′ extension directly to the last codon of the αHL gene (Jones 1995). The extension encoded the 94 amino acids of the Bacillus hemolysin II C-terminal tail (residues 289–382). The fused gene (αHL-TL) was generated in pT7-SC1 by cotransformation with the plasmid E. coli (Jones 1995). The extension encoded the 94 amino acids of the Bacillus hemolysin II C-terminal tail (residues 289–382). The fused gene (αHL-TL) was generated in pT7-SC1 by cotransform-
Oligomerization of HlyII on liposomes

Egg yolk phosphatidylcholine, cholesterol, and phosphatidic acid in chloroform were mixed in the desired molar ratio of 55:25:20. After drying under vacuum, the lipid film was resuspended in buffer (10 mM Tris, pH 8.0, 150 mM NaCl, 1 mM EDTA) to a total lipid concentration of 5 mg mL\(^{-1}\). Liposomes were prepared by ultrasonication for 30 min on ice using a probe sonicator (Dy-xotech Sonic Dismembrator Model 150) (relative output set to 50% power), followed by a brief centrifugation (30 sec, 16,000 \(g\)) to remove titanium particles. \(^{35}\)S-Methionine-labeled HlyII or HlyII(\(\Delta CT\)) (2 \(\mu\)L of an IVTT reaction mix) was incubated with freshly prepared liposomes (8 \(\mu\)L) for 1 h at room temperature. After solubilization in Laemmli sample buffer, the mixture was subjected to SDS-polyacrylamide gel electrophoresis.

Proteinase K treatment of HlyII, HlyII(\(\Delta CT\)) and \(\alpha HL\) polypeptides on membranes

Proteinase K (Sigma, #P-0390) solutions (5.0, 0.5, and 0.05 mg mL\(^{-1}\) in water) were prepared by dilution of an enzyme stock (10 mg mL\(^{-1}\) in water) and used immediately. Limited proteolysis was performed on HlyII, HlyII(\(\Delta CT\)) and \(\alpha HL\) bound to rRBCM. The membranes were resuspended in MBSA at 0.19 mg membrane protein mL\(^{-1}\) and divided into four tubes (18 \(\mu\)L in each). Proteinase K or water (2 \(\mu\)L) was added to each tube. After 5 min at room temperature, the reactions were stopped by treatment with PMSP (9 mM final, added in 2 \(\mu\)L of isopropanol, 5 min, room temperature), followed by the addition of 2x Laemmli loading buffer. The samples were subjected to electrophoresis in 10% SDS-polyacrylamide gels (heated samples) or 12% gels (heated samples: 95°C, 5 min).

Gel purification of HlyII oligomers

HlyII, HlyII(\(\Delta CT\)) and HlyII(\(\Delta CT\))\(\cdot D8\), and were prepared by translation in the presence of rRBCM as described above, but in preparative amounts (75 \(\mu\)L IVTT reaction). To obtain, HlyII(\(\Delta CT\))\(\cdot D8\), HlyII(\(\Delta CT\)) and HlyII(\(\Delta CT\))\(\cdot D8\) were translated using the corresponding plasmids at a ratio of 5:1. The oligomers were purified by SDS-polyacrylamide gel electrophoresis in an 8% gel in the presence of 0.1 mM sodium thioglycolate (Movileanu et al. 2001; Miles et al. 2001), stored at −80°C in 10 mM Tris.HCl, pH 7.5, and used for bilayer recordings without further treatment.

Hetero-oligomer formation for the determination of stoichiometry

Hetero-oligomers of hemolysin II subunits containing HlyII and/ or HlyII-D8, and HlyII(\(\Delta CT\)) and/or HlyII(\(\Delta CT\))\(\cdot D8\) were prepared by mixing the corresponding plasmids in the desired molar ratios (see Fig. 7 legend) prior to IVTT in the presence of rRBCM. To obtain \(\alpha HL\) heteromers, \(\alpha HL\) and/or \(\alpha HL\)-D8 were used. The washed membrane pellets were solubilized with Laemmli sample buffer and subjected, without heating, to electrophoresis in 5% SDS-polyacrylamide gels. Autoradiographs were made of the dried gels.

Planar lipid bilayer recordings

All measurements were performed at 25°C. Numerical values are given as the mean ± SD (\(\sigma_m\)). Planar lipid bilayer membranes were formed with 1,2-diphytanoyl-sn-glycero-3-phosphocholine (Avanti Polar Lipids) on a 150–160 \(\mu\)m-diameter aperture in a 25 \(\mu\)m-thick Teflon film (Goodfellow Corporation) separating the cis and trans compartments (2 mL each) of a bilayer apparatus (Mittal and Mueller 1972). Prior to forming the lipid bilayer, the orifice was pretreated with 10% (v/v) hexadecane (#29,631-7; Aldrich) in n-pentane (Burdick & Jackson) and allowed to dry thoroughly. For most measurements, the cis and trans chambers contained 1 M KCl, 5 mM HEPES, pH 7.4. Various concentrations of KCl in 5 mM HEPES, pH 7.4 were used for ion selectivity measurements. Protein samples were added to the cis chamber, which was at ground.

Currents were recorded by using a Dagan 3900A patch clamp amplifier (Dagan Corporation) with a 3910 expander and a built-in low-pass four-pole Bessel filter set at 5 kHz. Data were stored on digital audio tape with a DAS-75 data recorder (Dagan Corporation). Prior to analysis, the signal was low-pass filtered at 1 kHz with an eight-pole Bessel filter (Model 902, Frequency Devices) and acquired with a Digidata 1200A A/D board with a sampling time interval of 200 \(\mu\)sec. Data were acquired and analyzed with pClamp 8.0 software (Axon Instruments). Single-channel conductance values were determined by fitting the peaks in amplitude histograms to Gaussian functions. Current–voltage (I-V) relationships for single channels were determined by recording the currents obtained after stepwise changes in applied potential. The permeability ratios (\(P_{K+}/P_{Cl-}\)) were calculated from experimentally determined reversal potentials (\(V_{r}\)) by using the Goldman-Hodgkin-Katz (GHK) equation (Hille 2001) and the appropriate activity coefficients for KCl solutions (Zemaitis et al. 1986). In these measurements, the cis compartment contained 1000 mM KCl, while the other chamber contained 200 mM KCl. Any electrode DC offset was balanced prior to the addition of protein to the cis chamber. The applied voltage that gave zero current was noted. In addition, the reversal potential was more accurately determined by polynomial fits to current–voltage (I-V) data. Symmetrical solutions were then reestablished to evaluate whether or not any DC offset had built up during the course of the experiment. In all cases, the offset was less than 1 mV.

Acknowledgments

This work was supported by grants from the DOE and NIH, G.M. holds an MD-PhD training fellowship at The Texas A&M University System Health Science Center, College of Medicine, and was the recipient of an ASSERT (ARO) award. The authors thank Daniel Zeigler, BGSC Director, Ohio State University, for graciously supplying Bacillus strains; Orit Braha and Li-Qun Gu for their advice on ion selectivity; Brian Lauman for technical help; Michael Palmer for advice on liposomes; and Sean Conlan for guidance on using Spock.

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