Ion selectivity of the anthrax toxin channel and its effect on protein translocation

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Anthrax toxin consists of three ~85-kD proteins: lethal factor (LF), edema factor (EF), and protective antigen (PA). PA63 (the 63-kD, C-terminal portion of PA) forms heptameric channels ((PA63)7) in planar phospholipid bilayer membranes that enable the translocation of LF and EF across the membrane. These mushroom-shaped channels consist of a globular cap domain and a 14-stranded β-barrel stem domain, with six anionic residues lining the interior of the stem to form rings of negative charges. (PA63)7 channels are highly cation selective, and, here, we investigate the effects on both cation selectivity and protein translocation of mutating each of these anionic residues to a serine. We find that although some of these mutations reduce cation selectivity, selectivity alone does not directly predict the rate of protein translocation; local changes in electrostatic forces must be considered as well.

INTRODUCTION

The tripartite anthrax toxin consists of lethal factor (LF; 90 kD), edema factor (EF; 89 kD), and protective antigen (PA; 83 kD). PA forms channels in cell membranes that allow for the passage of LF (a zinc metalloprotease; Duesbery et al., 1998; Vitale et al., 1998) and EF (an adenylate cyclase; Leppla, 1982) into the cytosol, where they interfere with cellular homeostasis and ultimately lead to cell death (for a general review of anthrax toxin, see Young and Collier, 2007). Channel self-assembly begins when monomeric PA binds to either of two known receptors (ANTXR1 and ANTXR2) and is subsequently activated by proteolytic cleavage, which removes the N-terminal 20-kD portion of PA (PA20). The remaining C-terminal 63-kD portion of PA (PA63) then heptamimerizes (Young and Collier, 2007) or octamerizes (Kintzer et al., 2009) to form the ring-shaped prepore, which may bind up to three molecules of EF and/or LF. The entire ligand–receptor complex is then endocytosed; in the low pH environment of the endosome, the prepore undergoes conversion to the active (PA63)7 pore form. Finally, (PA63)7 inserts into the endosomal membrane and facilitates the transfer of EF/LF into the cytosol (Young and Collier, 2007). Although there is no crystal structure of (PA63)7, a computer model depicts these channels as mushroom shaped, with a globular cap domain and a cylindrical stem domain (Nguyen, 2004). The ~100-Å stem is a 14-stranded β-barrel that spans the membrane and extends out of the cell membrane (Benson et al., 1998; Nassi et al., 2002). A recent, high resolution cryo-EM structure of the (PA63)7 pore (Fig. 1 B) (Jiang et al., 2015) confirms the structure predicted by the computer model and low resolution EM structures (Katayama et al., 2008; Gogol et al., 2013).

(PA63)7 can be reconstituted in planar phospholipid bilayer membranes, forming channels that are capable of translocating LFN (the N-terminal, 263-residue portion of LF), as well as whole LF and EF, when the appropriate voltage (Zhang et al., 2004b) or pH conditions (Krantz et al., 2006) are applied. Under such conditions, LF enters the channel from the cis side (the side to which (PA63)7 has been added), N terminus first (Zhang et al., 2004a), passes through the channel cap and stem, and then exits into the trans solution (Finkelstein, 2009). In the absence of translocating protein, these channels have a conductance of ~55 pS (in 100 mM KCl, pH 5.5) (Krantz et al., 2005) and are highly (although not ideally) selective for cations (Blaustein and Finkelstein, 1990). The mechanistic basis behind both pH- and voltage-driven translocation through the (PA63)7 channel is critically dependent on the channel’s cation selectivity. At pH 5.5, LFN bears a net negative charge of ~6. Because LF4 translocation through the channel occurs when positive voltages are applied to the cis solution, the carboxyl groups on LF4 must be neutralized for voltage-driven translocation to occur. However, at pH 5.5, the carboxyl groups on aspartate and glutamate residues are in the ionized form (~97% of the time). Nonetheless, it is the minority protonated form that is favored for entry into the channel, largely because of the cation...
selectivity of the channel (Finkelstein, 2009). Similarly, a pH gradient in which the pH of the cis solution is lower than that of the trans solution will also cause LFN translocation to occur, as the aspartate and glutamate residues on LFN are more likely to exist in their protonated form on the cis side than on the trans side. Therefore, the rate of entry of these residues into the channel is greater from the cis side than from the trans side because of the cation selectivity of the channel, biasing the Brownian motion of LFN toward the trans side (Krantz et al., 2006).

Each of the seven monomers of (PA63)7 contributes a phenylalanine residue, F427, to form a ring known as the phenylalanine clamp (Φ-clamp), located at the junction between the cap and stem domains of the channel. The Φ-clamp creates a tight seal around the translocating peptide, facilitating the unfolding of the protein as well as creating a near-complete block in ion conductance that is important for efficient translocation (Krantz et al., 2005). It has also been identified as an important site of anion exclusion (Basilio et al., 2009): adding a single non-titratable sulfonate group at most positions on LFN largely abolishes translocation, as the channel strongly disfavors the entry of negatively charged groups, but mutating the residues of the Φ-clamp to alanine residues ((PA63F427A)7) restores translocation (Basilio et al., 2009). These results are consistent with the observation that mutating the Φ-clamp residues to alanine residues reduces the K+/Cl− selectivity of the channel (Basilio et al., 2009).

In addition to the Φ-clamp residues, it is also expected that the six rings of negatively charged residues that line the channel stem (D276, E302, E308, D315, D335, and E343; see Fig. 1A) may contribute to the cation selectivity of the channel. In this paper, we explore the effect on both cation selectivity and protein translocation when each of these anionic residues is mutated to a serine residue. Although we find that specific anionic mutations may reduce macroscopic selectivity and/or impede translocation, the local distribution of the anionic residues in the channel must be considered as well.

**MATERIALS AND METHODS**

**Molecular biology and protein purification**

The pET-22b (EMD Millipore) vectors encoding PA63D276S, PA63E302S, PA63E308S, PA63D315S, PA63D335S, PA63E343S, and PA63E302S/E308S/D315S were constructed using a pET-22b vector encoding WT PA63 (provided by J. Collier, Harvard Medical School, Boston, MA) and the QuikChange Site-Directed Mutagenesis kit (Agilent Technologies). PA63D276S, PA63E308S, PA63D315S, and PA63D335S were recombinantly expressed, purified, and nicked with trypsin (to detach PA20 and leave behind PA63) as described previously (Benson et al., 1998). PA63E302S, PA63E343S, and PA63E302S/E308S/D315S were expressed, purified, and then nicked with trypsin according to a slightly different protocol as described previously (Anderson and Blaustein, 2008).

WT (PA63)7 (trypsin-treated PA63 from which PA20 was detached; Blaustein et al., 1989) was provided by S. Zhang (Harvard Medical School, Boston, MA). (PA63F427A)7 was the same sample reported previously (Krantz et al., 2005). LFN (the N-terminal, 263 residues of LF, containing the EMD Millipore His6-tag at the N terminus) and LFN A59C 2-sulfonatoethyl MTS (MTS-ES; LFN with an N-terminal His6-tag and a sulfonate group attached at residue 59, which has been mutated to a cysteine) were also the same samples reported previously (Basilio et al., 2009). The His6-tags on LFN and LFN A59C MTS-ES were cleaved with thrombin according to EMD Millipore’s instructions.

**Figure 1.** (A) Residues that line the channel lumen and exterior of the (PA63)7 stem. Anionic residues are highlighted in red; cationic residues are highlighted in blue. The six anionic residues lining the lumen are circled in green. Horizontal dashed blue lines represent the lipid bilayer (adapted from Nassi et al., 2002). (B) Cryo-EM structure (Jiang et al., 2015) of the (PA63)7 channel (resolution of 2.9 Å). Horizontal dashed lines represent the lipid bilayer.
Planar lipid bilayers

Planar lipid bilayers were formed using the brush technique (Mueller et al., 1963) across a 0.5-mm hole in a Teflon partition. The partition divided a Lucite chamber into two compartments that each held 3 ml of 250 mM KCl (or 100 mM KCl), 5 mM potassium-succinate, and 1 mM EDTA, pH 5.5; each compartment was stirred using tiny magnetic stir bars. Agar salt bridges (3 M KCl; 3% agar) connected Ag/AgCl electrodes in saturated KCl solutions to the two compartments. The lipid solution was 3% diphtyanoyl-phosphatidylcholine (Avanti Polar Lipids, Inc.) in n-decane. Membrane formation was observed visually. Experiments were all performed under voltage-clamp conditions; voltages were taken as those of the cis solution with respect to the trans solution (held at virtual ground). Current responses were filtered at 1 Hz by a low-pass eight-pole Bessel filter (Warner Instruments), recorded on a DMP-4B Physiograph chart recorder (Narco Bio Systems), and digitally stored using NI USB-6211 Data Acquisition Board (National Instruments) (Basilio et al., 2009; Schiffmiller and Finkelstein, 2015).

Selectivity profiles

After a lipid bilayer was formed, (PA63)7 prepore heptamer (or one of the mutant prepore heptamers) was added to the cis solution (250 or 100 mM KCl, 5 mM potassium-succinate, and 1 mM EDTA, pH 5.5), causing the formation of hundreds to thousands of channels in the lipid bilayer. 3 M KCl was then added in incremental amounts to the cis solution (an incremental amount of solution was withdrawn before each 3-M KCl incremental addition), and for each increment, the reversal potential (Erev) was recorded. Reversal potentials were plotted against KCl activity ratios (a_{cis}/a_{trans}) using activity coefficients recorded in the literature (Robinson and Stokes, 1965). The K+ concentration was taken as equal to the KCl concentration; it was estimated that 10 mM K+ was contributed by potassium-succinate and potassium-EDTA.

Translocation experiments

After a lipid bilayer was formed, (PA63)7 prepore heptamer (or one of the mutant prepore heptamers) was added to the cis solution (250 mM KCl, 5 mM potassium-succinate, and 1 mM EDTA, pH 5.5). Once conductance leveled off at 20 mV, LFN (or LFN_A59C_MTS-ES) was added to the cis compartment to a final concentration of \( \approx 3 \) nM. Excess ligand was subsequently removed by perfusing the cis solution; the exchange of five volumes was performed using a manual pump while the voltage was held at 20 mV. The voltage was then switched to 3 mV, and a small amount of 1 M potassium phosphate (dibasic) was added to the trans solution to raise the pH to either 6.15 or 6.5. After 15 s, the voltage was switched back to 20 mV to allow translocation to take place. To determine the rate of translocation, we normalized the final conductance to the conductance observed at 20 mV after the channels were unblocked at -50 or -55 mV for 30 s (Basilio et al., 2009).
Ion selectivity of the anthrax toxin channel

RESULTS

Ion selectivity profiles of the asp/glu mutant channels

We generated ion selectivity profiles for each of the six asp/glu mutant channels: (PA63D276S)7, (PA63E302S)7, (PA63E308S)7, (PA63D315S)7, (PA63D335S)7, and (PA63E343S)7. After inserting channels of a given type into a membrane separating symmetric solutions of 100 mM KCl, pH 5.5, we raised the KCl concentration of the cis compartment by adding incremental amounts of concentrated KCl; for each increment, we recorded the reversal potential (E_{rev}). We then plotted E_{rev} versus the activity ratio (a_{cis}/a_{trans}) of KCl (Fig. 2) and compared these profiles to those of WT (PA63)7 and (PA63F427A)7 channels. We also repeated these experiments beginning with symmetric solutions of 250 mM KCl, pH 5.5 (Fig. 3).

Applying the Goldman–Hodgkin–Katz (GHK) equation (Hodgkin and Katz, 1949), we were able to roughly calculate the permeability ratio (P_K/P_Cl) for each channel. Table 1 lists the permeability ratios (P_K/P_Cl) of each channel.

Table 1: Permeability ratios (P_K/P_Cl) of channels

<table>
<thead>
<tr>
<th>Channel</th>
<th>100 mM KCl</th>
<th>250 mM KCl</th>
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</thead>
<tbody>
<tr>
<td>WT (PA63)7</td>
<td>22.2</td>
<td>9.1</td>
</tr>
<tr>
<td>(PA63E302S)7</td>
<td>17.5</td>
<td>8.9</td>
</tr>
<tr>
<td>(PA63E343S)7</td>
<td>20.8</td>
<td>8.7</td>
</tr>
<tr>
<td>(PA63D276S)7</td>
<td>20.8</td>
<td>8.5</td>
</tr>
<tr>
<td>(PA63D315S)7</td>
<td>15.4</td>
<td>8.2</td>
</tr>
<tr>
<td>(PA63D335S)7</td>
<td>14.7</td>
<td>7.7</td>
</tr>
<tr>
<td>(PA63D315S)7</td>
<td>12.2</td>
<td>6.5</td>
</tr>
<tr>
<td>(PA63F427A)7</td>
<td>9.5</td>
<td>4.6</td>
</tr>
</tbody>
</table>

Permeability ratios (P_K/P_Cl) of channels, calculated by applying the GHK equation to the cation selectivity profiles of each channel with [KCl]_{trans} = 250 mM KCl. Numbers represent P_K/P_Cl.
mutant channel (Table 1). As expected (Basilio et al., 2009), the value of $P_K/P_Cl$ for the $(PA_{32}F427A)_7$ channel was significantly smaller than that for the WT $(PA_{32})_7$ channel at both salt concentrations by a factor of $\sim 2$. In terms of the asp/glut mutant channels, at both salt concentrations, the $(PA_{32}D315S)_7$ channel was the least cation selective (although not as poor as that of $(PA_{32}F427A)_7$), followed by $(PA_{32}E308S)_7$. At 250 mM KCl, the other asp/glut mutant channels did not have cation selectivities that were very different from that of the WT channel, whereas at 100 mM KCl, the $(PA_{32}D335S)_7$ and $(PA_{32}E302S)_7$ channels were somewhat less cation selective than the WT $(PA_{32})_7$ channel.

An interesting aspect of the preceding data is that for each channel type, the reversal potential for a given KCl activity ratio is smaller if the ratio is taken with respect to 250 mM KCl rather than to 100 mM KCl (Figs. 2 and 3). In terms of the GHK permeability coefficients, this is expressed by the fact that $P_K/P_Cl$ is smaller (Table 1). This phenomenon arises naturally if the basis for the cation selectivity of the channel is that it bears a net negative charge. In a simplistic model of the channel in which we assume that it is a fixed charge membrane of uniform fixed charge density, then the reversal potential $E_{rev}$ across the membrane is given by the algebraic sum of the Donnan potentials at the two membrane–solution interfaces (Teorell, 1953):

$$E_{rev} = \frac{RT}{F} \ln \frac{[N]_{cis}}{[N]_{trans}},$$

where

<table>
<thead>
<tr>
<th>Channel</th>
<th>100 mM KCl</th>
<th>250 mM KCl</th>
<th>Mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT $(PA_{32})_7$</td>
<td>0.70</td>
<td>0.85</td>
<td>0.78</td>
</tr>
<tr>
<td>$(PA_{32}E302S)_7$</td>
<td>0.60</td>
<td>0.83</td>
<td>0.72</td>
</tr>
<tr>
<td>$(PA_{32}E343S)_7$</td>
<td>0.68</td>
<td>0.80</td>
<td>0.74</td>
</tr>
<tr>
<td>$(PA_{32}D276S)_7$</td>
<td>0.70</td>
<td>0.74</td>
<td>0.72</td>
</tr>
<tr>
<td>$(PA_{32}D335S)_7$</td>
<td>0.58</td>
<td>0.77</td>
<td>0.68</td>
</tr>
<tr>
<td>$(PA_{32}E308S)_7$</td>
<td>0.55</td>
<td>0.72</td>
<td>0.64</td>
</tr>
<tr>
<td>$(PA_{32}D315S)_7$</td>
<td>0.50</td>
<td>0.63</td>
<td>0.57</td>
</tr>
<tr>
<td>$(PA_{32}F427A)_7$</td>
<td>0.38</td>
<td>0.48</td>
<td>0.43</td>
</tr>
</tbody>
</table>

Table 2: Negative fixed charge density (molarity) of channels (calculated from Eq. 1) with $[KCl]_{trans} = 100$ and 250 mM KCl. Each entry is the value of $[N]$ that fits the data in Figs. 2 and 3.

Figure 4. The effect of mutating the anionic residues in the anthrax toxin channel stem on the rate of pH-driven LF$_N$ translocation (top, $\Delta pH = 0.65$; bottom, $\Delta pH = 1.0$). Each graph shows the normalized (see Materials and methods) increase in conductance that occurs (after perfusion of excess LF$_N$ out of the cis compartment) upon raising the pH of the trans compartment from pH 5.5 to pH 6.15 (top) or pH 6.5 (bottom) by adding a small amount of potassium phosphate (1 M, dibasic). Note the difference in the timescale between the two graphs; as is expected, translocation occurs at a much faster rate at $\Delta pH = 1.0$. Experiments were performed in 250 mM KCl.
fit at both $[KCl]_{trans} = 100$ mM and $[KCl]_{trans} = 250$ mM by the same $[N]$. The advantages of this approach compared with the GHK equation will be addressed in the Discussion.

pH-driven $L_{F_N}$ translocation through the asp/glu mutant channels

Although we wished to observe the effects of mutating the anionic residues in the anthrax toxin stem on both voltage- and pH-driven translocation, we unfortunately encountered significant voltage-dependent gating at high positive voltages (e.g., 50 mV) in several of the asp/glu mutant channels ($\text{PA}_63\text{E}302\text{S}$), $\text{PA}_63\text{D}315\text{S}$, and $\text{PA}_63\text{D}335\text{S}$), thereby making it very difficult to interpret the kinetics of $L_{F_N}$ translocation. Consequently, we were limited to the study of pH-driven translocation. All experiments were performed in 250 mM KCl to minimize gating that occurred at even small positive voltages (e.g., 20 mV).

At $\Delta p H = 0.65$ (Fig. 4, top, and Table 3), the rate of $L_{F_N}$ translocation through (PA63E302S)7 and (PA63E308S)7 channels ($t_{1/2}$: 20 and 22 s, respectively) was basically the same as that for the WT channel ($t_{1/2}$: 23 s). On the other hand, (PA63D335S)7 and (PA63E343S)7 channels exhibited significantly slower translocation rates ($t_{1/2}$: 50 and 43 s, respectively), and the (PA63D315S)7 channel exhibited the slowest translocation rate of all (the halftime was too slow to be measured).

At $\Delta p H = 1.0$ (Fig. 4, bottom, and Table 3), where $L_{F_N}$ translocation occurs on a much faster timescale, the (PA63D315S)7 channel still exhibited slow translocation (once again, the halftime was too slow to be measured). Meanwhile, all of the other asp/glu mutant channels...
exhibited very similar translocation rates to that of the WT channel (t1/2: 2.5 s), except for (PA63D335S)7 (t1/2: 4.9 s) and the (PA63E308S)7 channel (t1/2: 4.0 s).

Unfortunately, we were not able to observe translocation for the (PA63D276S)7 channel, because LFN exhibited poor binding to this channel at this salt concentration (250 mM KCl). Because residue D276 is located near residue D59, which exhibits the poorest cation selectivity of all the mutant channels (Table 1), has significantly slower translocation rates than those of the WT channel at both pH gradients (Fig. 4). However, although the (PA63E308S)7 channel is also noticeably less cation selective than WT (PA63)7, the E308S mutation does not seem to impede pH-driven translocation at ΔpH = 0.65 (Fig. 4, top). Conversely, (PA63E343S)7 is nearly as cation selective as the WT channel (Table 1), yet it exhibits slower LFN translocation at ΔpH = 0.65 (Fig. 4, top).

pH-driven translocation of LFN with a sulfonate group attached

As mentioned above, the Φ-clamp has been identified as a site of anion exclusion during protein translocation through the anthrax toxin channel, and mutating the phenylalanine residues near the Φ-clamp to alanines allows for the passage of a nontitratable sulfonate group (SO3-). Thus, we attempted to pass through the triple mutant channel (PA63E302S/E308S/D315S)7, whose cation selectivity is even poorer than that of (PA63F427A)7 (Fig. 5 B). As an aside, preliminary data have shown that the channels formed by the triple mutant channel have a single-channel conductance that is slightly smaller than that of the WT channel, but tend to open and close more frequently than does the WT channel (Fig. S2).

We observed pH-driven translocation of LFN A59C MTS-ES (LFN in which residue 59 is mutated from an alanine to a cysteine, followed by incubation with MTS-ES to attach a sulfonate group; Basilio et al., 2009) in both WT (PA63)7 and (PA63E302S/E308S/D315S)7 channels at a ΔpH of 1 (Fig. 6). As a control, we also observed the translocation of WT LFN through both channels. Whereas the translocation of LFN A59C MTS-ES through the WT channel was mostly abolished, there was virtually no translocation at all through the (PA63E302S/E308S/D315S)7 channel either. Thus, despite the poor cation selectivity of the triple mutant channel, it does not allow the passage of SO3-. It should be emphasized that this effect was not caused by steric hindrance of the bulky sulfonate group, as the similarly sized trimethylammonium group (-N(CH3)3+), attached to LFN at the same location as the sulfonate group (residue 59), was able to pass through the triple mutant channel unhindered (unpublished data).

DISCUSSION

Ion selectivity

In this paper, we observed the effects of mutating each of the six anionic residues lining the interior of the anthrax toxin channel to serines. Before discussing the effects of these mutations on cation selectivity, we must first address the possibility that streaming potentials and/or polarization effects (dilution potentials) may have skewed our results. By polarization effects, we mean that as a consequence of osmotic water flow across the membrane from the cis to the cis solution caused by the higher KCl concentration in the cis compartment, the KCl concentration at the cis membrane–solution interface is reduced, and at the trans membrane–solution interface it is elevated. Consequently, the actual auni/auni.
across the membrane is less than the bulk $a_{\text{ion}}/a_{\text{trans}}$, thereby artificially reducing the magnitude of $E_{\text{rev}}$. To check for polarization effects, we performed cation selectivity experiments (not depicted) with valinomycin (which is ideally selective for potassium) at both $[\text{KCl}]_{\text{trans}} = 100 \, \text{mM}$ and $[\text{KCl}]_{\text{trans}} = 250 \, \text{mM}$. At both salt concentrations, the reversal potential profiles for valinomycin (at the same $a_{\text{ion}}/a_{\text{trans}}$ as in the (PA63)$_7$ experiments) did not deviate from ideal selectivity, indicating that no polarization is taking place.

As far as streaming potentials are concerned, note that if streaming potentials were an issue, the greatest “bending” of the selectivity profiles should be observed with the WT channel, as it is more cation selective than any of the mutant channels (Tables 1 and 2). On the contrary, the reverse is the case (Figs. 2 and 3). Furthermore, any bending of the selectivity profiles of the mutant channels caused by streaming potentials is expected to be even less significant, as the mutant channels are less cation selective than the WT channel (Tables 1 and 2). Thus, we believe that the cation selectivity profiles that we have generated for the mutant channels have not been significantly affected by streaming potentials.

Using the GHK equation, we calculated the permeability ratio ($P_k/P_C$) for the WT channel, the (PA63F427A)$_7$ mutant channel, and the six asp/glutamate mutant channels (Table 1) at both $[\text{KCl}]_{\text{trans}} = 100 \, \text{mM}$ and $[\text{KCl}]_{\text{trans}} = 250 \, \text{mM}$. To reiterate our findings, $P_k/P_C$ was much smaller for the (PA63F427A)$_7$ channel than for the WT (PA63)$_7$ channel at both salt concentrations, whereas the (PA63D315S)$_7$ channel exhibited the poorest cation selectivity, followed by (PA63E308S)$_7$. At 250 mM $[\text{KCl}]_{\text{trans}}$, the other asp/glutamate mutant channels had similar cation selectivities to that of the WT channel, whereas at 100 mM $[\text{KCl}]_{\text{trans}}$ the (PA63D335S)$_7$ and (PA63E302S)$_7$ channels were a little less selective than the WT (PA63)$_7$ channel.

Although widely used, the GHK equation has certain deficiencies. The particular one that concerns us here is that it assumes that the partition coefficients of K$^+$ and Cl$^-$ between the solution and the channel are constant, independent of salt concentration. In practice, we see that the permeability ratios calculated at $[\text{KCl}]_{\text{trans}} = 250 \, \text{mM}$ are much smaller than those calculated at $[\text{KCl}]_{\text{trans}} = 100 \, \text{mM}$ (Table 1), whereas the GHK equation implies, at face value, that the permeability ratio is independent of salt concentration.

A simple explanation for the difference in cation selectivity at the two salt concentrations ($[\text{KCl}]_{\text{trans}} = 100 \, \text{mM}$ and $250 \, \text{mM}$) is the existence of fixed negative charges in the channels. For this reason, we simplistically modeled the channels as a membrane with a uniform fixed negative charge density, using Eq. 1 (Teorell, 1953). Although this is a gross oversimplification of the distribution of negative charges in the (PA63)$_7$ channels, it has the advantage that the poorer cation selectivity we observe when $[\text{KCl}]_{\text{trans}} = 250 \, \text{mM}$ arises as a natural consequence of the model (see Eq. 1). We see, in fact, that a given $[\text{N}]$ adequately fits the selectivity data for both 100 and 250 mM KCl for each channel type (Fig. S1). We emphasize that this model cannot be taken literally; if the distribution of negative fixed charge density in the channel was indeed uniform, then selectivity should be affected in the same manner regardless of which asp/glutamate residue was mutated. Furthermore, the (PA63F427A)$_7$ channel exhibited worse cation selectivity ($P_k/P_C$) than the WT channel by a factor of $\sim 2$ (Table 1) and was thus modeled as having a smaller value of $[\text{N}]$, yet no actual charges were mutated in the $\Phi$-clamp mutant channel.

Translocation
Our results of pH-driven translocation of LF$_5$ through the WT (PA63)$_7$ channel, as well as through five of the six asp/glutamate mutant channels (Fig. 4 and Table 3), indicate that the translocation rates through these channels are not necessarily correlated with cation selectivity. At $\Delta pH = 0.65$ (in 250 mM KCl), the (PA63D315S)$_7$, (PA63D335S)$_7$, and (PA63E343S)$_7$ channels all exhibited significantly slower translocation rates than through the WT channel (Fig. 4, top, and Table 3). That the translocation rate through the (PA63D315S)$_7$ channel was the slowest is consistent with its being the least anion exclusionary of the six asp/glutamate mutant channels at this salt concentration (Table 1). However, it is not obvious why the (PA63D335S)$_7$ and (PA63E343S)$_7$ channels should have such slow translocation rates, given that the cation selectivities of these two channels are comparable to that of the WT channel (Table 1).

This observation emphasizes that to truly understand translocation through the anthrax toxin channel, we must consider not only macroscopic cation selectivity but rather the local electrostatic forces acting in the channel as well. A previous electrostatic modeling study (Wynia-Smith et al., 2012) has suggested that there exist two areas of opposite charge in the channel stem, comprised of an anion-repulsive region (the top of the stem) and a cation-repulsive region (the middle of the stem). These electrostatic forces are generated by the charged residues lining both the lumen and exterior of the channel (Fig. 1 A). D276, D335, and E343 have been identified as residues that contribute to the anion-repulsive region (Wynia-Smith et al., 2012) and may be part of an “ion selectivity filter” that includes the $\Phi$-clamp. The model that has been proposed is that once the appropriate pH gradient has been established, a polypeptide substrate can pass through the anion-repulsive feature of the upper stem after its aspartate and glutamate residues are protonated by the low pH of the cis solution (Wynia-Smith et al., 2012). As the polypeptide moves farther down the channel, the higher pH of the trans solution will deprotonate its aspartate and glutamate residues, propelling it away from the anion-repulsive region.
The cation-repulsive region will also favor deprotonation of these residues. In this way, an electrostatic “ratchet” is imposed on the substrate, biasing its movement toward the trans solution and preventing reverse translocation of the chain.

Thus, if D335 and E343 are indeed important components of this electrostatic ratchet, then it is understandable that at \( \Delta pH = 0.65 \), mutating these residues to serines would result in slow, inefficient translocation compared with what is observed in the WT channel (Fig. 5 B). On the other hand, at \( \Delta pH = 1.0 \), LF\(_N\) translocation in the (PA\(_{63}\)E343S)\(_7\) channel is comparable to that of the WT channel, whereas translocation in the (PA\(_{63}\)D335S)\(_7\) channel is only a little slower (Fig. 4, bottom). Perhaps the larger pH gradient results in more efficient deprotonation of the polypeptide chain once it moves down the channel stem (closer to the trans solution), compensating for any interference of the D335S/E343S mutations with the electrostatic ratchet.

Finally, we also sought to identify the defining feature that prevents a nontitratable sulfonate group (-SO\(_3^-\)) from passing through the (PA\(_{63}\))\(_7\) channel. Although mutating the \( \Phi \)-clamp residues to alanines allows for the translocation of a sulfonate group attached at most locations on LF\(_N\) (Basilio et al., 2009), it was simply not clear at the time whether this phenomenon was caused by reducing the macroscopic cation selectivity of the channel or by possible changes in the local electric field, caused by mutating the \( \Phi \)-clamp. (It was clearly not caused by a steric effect, as the comparably sized -N(CH\(_3\))\(_3^+\) attached at the same locations did not hinder LF\(_N\) translocation.) We have now found that the (PA\(_{63}\)E302S/E308S/D315S)\(_7\) channel, despite its being less cation selective than the (PA\(_{63}\)F427A)\(_7\) channel (Fig. 5 B), did not allow for the passage of LF\(_N\) once an SO\(_3^-\) group was attached (Fig. 6). Thus, it would appear once again that local electrostatic forces must be considered, and not simply macroscopic cation selectivity.

**Comparison of voltage- and pH-driven translocation**

As mentioned earlier, because of significant voltage-dependent gating observed in several of our asp/glu mutant channels, we were unable to perform voltage-driven translocation experiments. However, we think it is useful to briefly consider here voltage-driven translocation compared with \( pH \)-driven translocation.

In both cases, reducing the cation selectivity of the channel, and thereby allowing nonprotonated aspartates and glutamates to enter the stem, should reduce the rate of translocation of LF\(_N\), but by different mechanisms. In the case of voltage-driven translocation, the force on the translocated peptide is reduced because its net positive charge is reduced. In the case of \( pH \)-driven translocation, the force is reduced because not all of the aspartates and glutamates “feel” the \( \Delta pH \), as they retain the same ionized state at both their entry to the channel at the cis end and their exit from the channel at the trans end.

Consider, in fact, three limiting scenarios, specifically in regard to the mechanism of protonation/deprotonation of negatively charged residues on LF\(_N\) that was discussed in this paper. In the first extreme scenario, if there were only positively charged and neutral residues on a peptide chain, translocation could be driven only by a \( \Delta V \), as \( pH \)-driven translocation requires that a Brownian ratchet be established through the protonation states of negatively charged residues. In the opposite scenario, if there were only negatively charged and neutral residues present on a peptide chain, translocation could occur only by establishing a \( pH \) gradient, as an applied positive voltage will cause translocation to occur only if the peptide has a net positive charge. Finally, in the third limiting scenario, if there were only neutral residues present on a peptide chain, then neither a \( \Delta V \) nor a \( \Delta pH \) would cause translocation to occur.

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Figure S1. Fitting Eq. 1 to the \( E_{\text{rev}} \) profiles of WT \((\text{PA}_63)_7\), \((\text{PA}_63\text{F427A})_7\), \((\text{PA}_63\text{D276S})_7\), and \((\text{PA}_63\text{E302S})_7\) channels with \([\text{KCl}]_{\text{trans}} = 100 \text{ mM}\) (left column) and \([\text{KCl}]_{\text{trans}} = 250 \text{ mM}\) (right column). \([N]\) is the mean of the negative fixed charge density calculated for each salt concentration (see Table 2). Dashed black lines represent ideal cation selectivity.
Figure S2. Fitting Eq. 1 to the $E_{rev}$ profiles of (PA$_{63}$E308S)$_7$, (PA$_{63}$D315S)$_7$, (PA$_{63}$D335S)$_7$, and (PA$_{63}$E343S)$_7$ channels with [KCl]$_{\text{trans}} = 100$ mM (left column) and [KCl]$_{\text{trans}} = 250$ mM (right column). [N] is the mean of the negative fixed charge density calculated for each salt concentration (see Table 2). Dashed black lines represent ideal cation selectivity.
Figure S3. Records of the single-channel conductance of both the (PA₆₃E₃₀₂S/E₃₀₈S/D₃₁₅S)₇ channel (top) as well as the WT (PA₆₃)₇ channel (bottom), in 250 mM KCl at 20 mV. WT (PA₆₃)₇ channels have a conductance of \( \sim 100 \) pS, whereas (PA₆₃E₃₀₂S/E₃₀₈S/D₃₁₅S)₇ channels have a conductance of \( \sim 80-90 \) pS. Blue dashed lines represent the open-state conductance of a single channel, whereas the red dashed lines represent the zero baseline of conductance. Note that there are three channels present in the top record, whereas there are two channels present in the bottom record. Also note that the (PA₆₃E₃₀₂S/E₃₀₈S/D₃₁₅S)₇ channels open and close more frequently than the WT (PA₆₃)₇ channels. The top record is \( \sim 10 \) s long and the bottom record is \( \sim 70 \) s long; both records are filtered at 10 Hz.