A Receptor-based Switch that Regulates Anthrax Toxin Pore Formation

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Abstract

Cellular receptors can act as molecular switches, regulating the sensitivity of microbial proteins to conformational changes that promote cellular entry. The activities of these receptor-based switches are only partially understood. In this paper, we sought to understand the mechanism that underlies the activity of the ANTXR2 anthrax toxin receptor-based switch that binds to domains 2 and 4 of the protective antigen (PA) toxin subunit. Receptor-binding restricts structural changes within the heptameric PA prepore that are required for pore conversion to an acidic endosomal compartment. The transfer cross-saturation (TCS) NMR approach was used to monitor changes in the heptameric PA-receptor contacts at different steps during prepore-to-pore conversion. These studies demonstrated that receptor contact with PA domain 2 is weakened prior to pore conversion, defining a novel intermediate in this pathway. Importantly, ANTXR2 remained bound to PA domain 4 following pore conversion, suggesting that the bound receptor might influence the structure and/or function of the newly formed pore. These studies provide new insights into the function of a receptor-based molecular switch that controls anthrax toxin entry into cells.

Introduction

Cellular receptors can act as molecular switches that initiate conformational changes in microbial proteins required for cellular entry. Examples of such switches include an anthrax toxin receptor (described in detail below) as well as those for a number of viruses including HIV-1 and other retroviruses [1,2,3], measles virus [4], and herpesviruses [5]. The mechanisms by which these receptor-based switches function to promote cellular entry are only partially understood. In this report we set out to define the mechanism by which a receptor-based switch regulates anthrax toxin prepore-to-pore conversion.

Anthrax toxin, the key virulence factor secreted by Bacillus anthracis, is a bacterial AB toxin composed of three independent, plasmid-encoded polypeptide chains: the receptor-binding (B) moiety, protective antigen (PA), and two different enzymatic (A) moieties, lethal factor (LF) and edema factor (EF) [6,7,8]. The first step in cellular intoxication involves binding of an 83 kD form of PA (PA₆₅) to specific cell surface receptors. Although several PA receptors have been defined [9,10,11], anthrax toxin receptor type 2 (ANTXR2) (also known as capillary morphogenesis protein 2; CMG2), is the most physiologically relevant receptor [12,13,14]. ANTXR2 is a type 1 transmembrane protein and its extracellular von Willebrand factor type A (VWA) domain is the site of PA-binding [15,16]. Following receptor binding, PA₆₅ is cleaved to a 63kD form (PA₃₃) that spontaneously oligomerizes to form either a heptameric, or an octameric, PA₆₅ prepore structure [17,18]. Oligomeric PA₆₅-receptor complexes are then taken into cells primarily by a clathrin-dependent endocytic mechanism and delivered to an acidic endosomal compartment where low pH triggers formation of a PA₆₅ pore within an endosomal membrane [19,20]. LF and EF are then translocated through the pore and delivered to the cytosol where they promote intoxication [21].

X-ray structural analysis of monomeric and heptameric PA-ANTXR2 VWA-domain complexes revealed that the receptor acts as a molecular switch or clamp that inhibits prepore-to-pore conversion at neutral pH [15,16]. Specifically, the receptor VWA-domain interacts with the base regions of PA domains 2 and 4, thereby sterically hindering the movement of the PA 2β₃-2β₄ loop region necessary for pore formation [15,16]. Those findings led to a model in which release of the receptor contact with PA domain 2 at an acidic endosomal pH is necessary to permit the conformational changes required for PA pore formation [15,16]. Consistent with this idea, the pH threshold of the receptor-regulated toxin pore formation can be dictated by specific amino acid residues located at the PA domain 2-binding region of the ANTXR2 VWA-domain [22].

Presently, it is not clear if PA domain 2-receptor contacts are released at a step that occurs prior to, or is coincident with, prepore-to-pore conversion. Furthermore, it is not clear if the receptor remains attached following pore conversion and, if so, how it remains attached. Evidence supporting dissociation has come from co-immunoprecipitation experiments [23] and from previous NMR studies [24,25]. On the other hand, evidence in favor of receptor attachment has come from other co-immunoprecipitation studies [19,26], from NMR binding studies per-
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Author Summary

The bacterium that causes anthrax produces a toxin called anthrax toxin that is largely responsible for causing disease symptoms. The first step in anthrax intoxication involves binding of the toxin to a specific protein, called a receptor, on the cell surface. Receptor-binding acts like a switch to prevent the toxin from forming a pore in a cell membrane until the toxin-receptor complex is taken up into cells and delivered to a specific location (called an endosome) where it is exposed to an "acid bath". This acidic environment promotes structural changes in the toxin leading to pore formation in the endosomal membrane. In this report, we have studied how the receptor regulates anthrax toxin pore formation, information that could be useful for designing new therapeutic strategies to treat this disease.

formed with a fragment (Domain 4) of PA [27], and from the finding that the presence of a receptor seems to influence voltage-dependent inactivation and small molecule inhibition properties of the newly formed pore [28]. Based upon structural considerations, it has also been argued that the receptor may remain bound to serve as a structural support for the pore [16,29].

To clarify these issues, we have employed NMR techniques to monitor changes in the PA63 heptamer-ANTXR2 VWA domain contacts as a function of pH. Initially we attempted to examine the interaction between the ANTXR2 VWA domain and PA63 using chemical shift perturbation (CSP) by saturating in substoichiometric amounts of unlabeled PA63 into a 1H,15N labeled ANTXR2 sample. Based on results with other systems [30], we anticipated that saturating in PA63 might allow us to monitor chemical shift changes as a function of receptor binding and/or cause selective broadening of specific peaks associated with residues at the PA binding interface. If so, this would allow us to monitor specific receptor residues bound to PA63 under different pH conditions. Additionally, if shift perturbation of crosspeaks were detected using saturating conditions of the binding partner (PA63), this would help to approximate the fractional population of bound species versus the free species at equilibrium [30]. However, when low stoichiometric concentrations of PA63 were titrated into the ANTXR2-VWA domain sample, extensive line broadening and the disappearance of cross-peaks in the 1H,15N TROSY-HSQC was observed at ratios of 1:0.25 ANTXR2-VWA domain to PA63. This was likely due to the large size of the PA complex, indicating a larger effective correlation time ($\tau_c$), restricted local motion, and complete binding at the concentrations used. Therefore, we hypothesized that the method of transferred cross-saturation (TCS) may be well suited to investigate these interactions since this approach has previously been used to identify contact residues of protein ligands in large protein complexes [31].

To investigate the function of the ANTXR2-based switch, TCS was employed to monitor changes that occur in PA63 heptamer-ANTXR2 VWA-domain contacts as a function of pH. In this approach, an unlabeled protein is added at substoichiometric amounts to a deuterated, 15N-labeled protein, in this case, PA63 and the ANTXR2 VWA-domain, respectively. The aliphatic proton resonances of the unlabeled protein are then saturated with a brief radiofrequency pulse and this saturation is transferred selectively to contact residues of the 2H, 15N-labeled protein by spin diffusion. Consequently, the intensity of amide cross-peaks representing labeled residues that lie at the protein-protein interaction surface are selectively reduced by cross-relaxation [32,33]. Here we have used this technique to obtain evidence for a new toxin-receptor intermediate in the pathway leading to pore formation and show that the receptor remains attached to PA domain 4 following low pH-dependent conversion. Additionally, chemical shift perturbations associated with receptor residues located near the PA domain 4 binding region revealed moderate conformational changes that occur during the attachment and detachment of PA from the receptor.

Results

Generating the deuterated, 15N-labeled ANTXR2 VWA-domain

The deuterated, 15N-labeled ANTXR2 VWA-domain was produced as a GST-fusion protein from bacterial cells. In order to limit spin diffusion in the 15N-labeled protein, it was extensively deuterated by growing the cells in 100% D2O minimal media using 2H-glucose as the sole carbon source [32]. The labeled VWA-domain was purified to homogeneity as described under Materials and Methods and was unfolded to protonate the residues within the protein core and refolded to increase the number of cross-peaks in the [15N,1H] TROSY-HSQC spectrum. The integrity of the refolded protein was confirmed by circular dichroism (CD) analysis performed at either pH 8.0, 6.0, or 5.0 and in each case the protein displayed alpha-helical properties (Supplementary Figure S1). The refolded protein also functioned as an efficient receptor decoy in a toxin neutralization assay (Supplementary Figure S2). Moreover, Transverse Relaxation Optimized Spectroscopy-Heteronuclear Single-Quantum Coherence (TROSY-HSQC) spectrum analysis indicated that the protein was correctly refolded when compared to a control [15N,1H] TROSY-HSQC spectrum of the ANTXR2 VWA-domain that had not been previously denatured (Figure 1A). Assignments for the backbone resonances of the ANTXR2 VWA-domain were obtained using data from the following experiments: [1H,15N] TROSY-HSQC, 3D TROSY-HNCO, 3D TROSY-HN(CA)CO, 3D TROSY-HNCACB, 3D TROSY-HNCA, and a 3D 15N-edited NOESY-HSQC. NMR data were processed using NMRPipe and analyzed using Sparky and CARA software packages [34,35,36]. A representative example of this data analysis is shown in Supplementary Figure S3. Using this approach 87% of the backbone residues of the ANTXR2 VWA-domain, including the PA contact residues, were assigned (Figure 1A and B).

ANTXR2 residues that contact PA domains 2 and 4 are saturated by transferred cross saturation (TCS)

The principle of the TCS approach used to monitor PA63-ANTXR2 VWA-domain interactions is outlined in Figure 2A. In order to observe saturation transfer, the concentration of the binding partner must be kept sufficiently low to effectively allow for fast exchange so that amide cross-peaks are not broadened following its addition. The efficiency of TCS depends on the sample conditions as well as the binding constants between the receptor and PA. According to Shimada et al, TCS is applicable for a system where a large p_B, or fraction of bound ligands is preferred for high saturation efficiency, if $k_{off} > 0.1 \text{s}^{-1}$, or if $k_{off} \geq 10 \text{s}^{-1}$, a $p_B \geq 0.1$ is preferred [37]. Therefore, for the TCS experiments, the concentrations of the two protein partners were optimized by performing titration experiments at pH 8.0, and a
(10:1) molar ratio of the ANTXR2 VWA-domain to PA63 was chosen for all cross saturation experiments, because at this concentration there were no signs of peak broadening. Three separate sets of interleaved experiments were subsequently performed on the ANTXR2 VWA: (PA63)7 complex in buffers of pH 8.0, 6.0, and 3.0. Saturation transfer was achieved by applying a selective radiofrequency pulse at 0.8 ppm, prior to the [15N,1H] TROSY-HSQC. A pH of 8.0 was chosen for the initial analysis because it closely approximated those used previously for X-ray structural analysis of PA-receptor complexes, i.e. pH 7.5 [16] and pH 8.25, [15]. These studies revealed that the majority of the labeled residues in the ANTXR2 VWA-domain were not saturated by a radiofrequency pulse, i.e. those with similar signal intensities under conditions of no saturation (black peaks) or saturation (red peaks) (Figure 2B). However, a subset of the amide cross-peaks were saturated (black-only peaks) in the overlayed spectra (Figure 2B) and a number of those cross-peaks corresponded to contact residues with PA domains 2 or 4. For simplicity the saturation data was represented as 1D cross sections of the corresponding cross-peaks in the HSQC spectra (Figure 3A). The degree of saturation of each residue was calculated by dividing the observed peak intensity of the saturated spectrum (I_s) by the observed peak intensity of the control spectrum (I_o) (unsaturated).

In these studies an (I_s/I_o) value of 0.75 is considered significant and one of 0.5 highly significant, as in [37]. Based upon these criteria all of the PA domain 2 and 4 contact residues that could be unambiguously assigned were saturated under these conditions (Figure 3B). Taken together, this study verified that the TCS method can be used to specifically monitor contacts between the ANTXR2 VWA-domain and PA domains 2 and 4 in the heptameric toxin-receptor complex.

Identification of a new toxin-receptor complex intermediate at pH 6.0

To characterize the changes in PA-receptor contacts that occur after incubating the complex under mildly acidic
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**A**

\[ ^{2}\text{H}^{15}\text{N} \text{ labeled} \]
\[
\text{ANTXR2 VWA Domain}
\]

\[ ^{1}\text{H}^{14}\text{N} \text{ unlabeled} \]
\[
\text{PA}_{63} \text{ heptamer}
\]

\[ \text{PA}_{63} \text{ heptamer complexed to} \]
\[
\text{ANTXR2 VWA domain}
\]

\[ \text{Rotate 90°} \]

\[ \text{RF irradiation of the aliphatic sites of the unlabeled PA}_{63} \text{ heptamer} \]

\[ \text{ANTXR2} \]

Saturation is transferred via spin diffusion to the labeled ANTXR2 VWA domain.

**B**

![Data plot](https://example.com/data.png)

- X-axis: \( ^{1}\text{H} \text{ ppm} \)
- Y-axis: \( ^{15}\text{N} \text{ ppm} \)
Figure 2. The principle of Transferred Cross Saturation (TCS) applied to the (PA63)7-ANTXR2 complex. A) Schematic of TCS between ANTXR2 VWA domain and the PA63 pore. [1H,15N]-labeled ANTXR2 VWA-domain was mixed at a ratio of (10:1) with PA63 heptamer. Radiofrequency pulses were applied to the sample, in order to saturate the aliphatic protons of the PA63 heptamer. Saturation is then transferred to the contact residues of the labeled ANTXR2 VWA-domain, reducing the intensity of the corresponding cross peaks in the spectrum. B) Saturated and unsaturated spectra of the (PA63)7-ANTXR2 complex at pH 8.0. The [1H,15N] TROSY-HSQC spectra of the [1H,15N]-labeled ANTXR2-VWA domain complex with the PA63 heptameric prepore are shown overlayed under saturating or non-saturating (black) conditions.

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conditions, the TCS experiment was repeated at pH 6.0. That condition is approximately 1.0 pH unit above that needed to trigger toxin prepore to pore conversion when the PA63 heptamer is bound to the ANTXR2 VWA-domain [23,28] (Supplementary Figure S4). However, not all of the PA-contact residues that were observed at pH 8 were visible in the saturated and unsaturated spectra obtained at pH 6. This finding is probably due to a structural change upon loss of contact with the PA yielding to an increased H/D exchange or/and slow conformational exchange dynamics.

Analysis of the data clearly showed that the PA domain 4 contact residues that were resolved remained strongly saturated at pH 6 (residues G53, S54, N57, V115, E117, T118, H121, E122, and G123; Figure 4A and 4B). By striking contrast, PA domain 2 contact residues were much less saturated at pH 6.0 (Figure 4B). Taken together, these data are consistent with a model in which the receptor remains bound to PA domain 4 but its interactions with PA domain 2 are significantly weakened or are lost prior to prepore-to-pore conversion.

The ANTXR2 VWA-domain remains attached to PA domain 4 following prepore-to-pore conversion

When bound to ANTXR2, the PA63 prepore is triggered to form a pore species at pH values that are less than or equal to pH 5.2 [23,28] (Supplementary Figure S4). Therefore, to determine if the receptor remains attached to PA following pore formation, the TCS experiment was performed under both saturating and non-saturating conditions at pH 5.1. Consistent with the results obtained at pH 6.0, the receptor residues that contact PA domain 2 were not saturated at pH 5.1, with the possible exception of residue A159 (Figure 5A and 5B). More importantly however, virtually all of the domain 4 contact residues that could be resolved at this pH value were saturated at pH 5.1 (Figure 5B). These data are consistent with a model in which the receptor contacts with PA domain 2 are lost during anthrax toxin prepore to pore conversion but the receptor remains bound to PA domain 4.

The [1H,15N] TROSY-HSQC data also revealed chemical shift perturbations of certain receptor residues, including PA domain 4 contact-residues that were associated with (PA63)7 binding at pH 8.0 (Figure 6A). Small shift changes due to the isotope effect of being a highly deuterated protein were also taken into consideration as well as the pH effects [38]. Specifically, peaks associated with residues in and around helix 1 of the ANTXR2 VWA domain, that were involved in binding PA domain 4, significantly changed their position in the presence of PA63. These residues include G53, W59, and N57 (Figure 6B). A similar observation was made with residues Y46 and F47, which lie within the hydrophobic core of the ANTXR2 VWA domain, as well as with G135, which lies on the opposite face of the receptor VWA domain (Figure 6B). These latter effects are likely due to an allosteric or structural change in the receptor domain following PA binding. Strikingly, the peaks associated with all of these residues reverted back to their “unbound” configuration when the ANTXR2 VWA/PA63 complex was incubated at pH 5.1 (Figure 6A), even though the receptor remains bound to PA domain 4 under this condition. The only exception was residue Y46 which was not resolved at pH 5.1 but moved back towards its “unbound” configuration at pH 6. (Figure 6A).

Discussion

In this study we have used the TCS NMR approach to monitor how the ANTXR2-based receptor switch regulates anthrax toxin prepore-to-pore conversion. We showed that this is a robust method for identifying the receptor contacts with PA domains 2 and 4, in the prepore configuration at pH 8.0. We also obtained evidence at pH 6.0 for a new toxin-receptor intermediate in the pathway leading to pore formation, one in which the receptor remains bound to PA domain 4 but contacts with PA domain 2 have been significantly weakened. That intermediate would presumably exist within a mildly acidic early endosomal compartment during endocytic trafficking of toxin-receptor complexes [39]. Furthermore, we demonstrated that the ANTXR2 VWA-domain remains attached to PA domain 4 after triggering PA pore formation at pH 5.1, consistent with a more strongly acidic late endosomal pH [39]. Subtle structural changes, associated with reversion back to an unbound configuration, were also detected in residues located near the PA domain 4-binding site following pore conversion. This effect was also seen in the opposite face of the protein with residue G135, and with two hydrophobic residues within the core, Y46 and F47. It is known that chemical shifts of those nuclei that lie within close proximity of the binding partner can be substantially perturbed in the presence of that partner. However chemical shift perturbations (CSP) can also arise from allosteric effects as well as extended conformational changes that may occur in the target protein upon protein partner binding [40,41]. These latter effects most likely account for the chemical shift perturbations seen with residues Y46, F47, and G135, which lie distal from the PA63 binding site of the receptor (Figure 6B). Taken together, these studies have led to a revised model of the changes in toxin-receptor contacts during pore formation (Figure 7) and support the idea that the bound receptor may influence the structural and/or functional properties of the toxin pore.

It is unlikely that the TCS effects that were observed could be attributed to non-specific aggregation of the PA63 heptamer-receptor complex at the different pH values tested since these effects were almost exclusively restricted to the toxin-binding face of the receptor. Indeed, inspection of 1D slices of the TCS experiments for selected crosspeaks did not indicate a broadening of lineshapes at several residues at pH 8 or pH 5, as would have been expected if there was aggregation (data not shown). Additionally, to further assess protein aggregation at pH 8 and pH 5, a wavelength scan (from 280–360 nm) was performed on the ANTXR2 VWA domain in complex with PA63 at the same 10:1 ratio used for the TCS NMR experiments, since protein aggregation can be monitored at 340 nm [42,43]. These studies revealed no substantial increase of absorbance at 340 nm between pH 8 and pH 5 (Supplementary Table S1), and visual inspection of the sample yielded no noticeable turbidity in the supernatant, under any of the conditions tested. Furthermore, there was no
substantial difference in the absorbance at 280 nm seen with either the pH 8.0 or pH 5.0 samples before or after a 24 hour incubation at 37°C (Supplementary Table S2). Therefore, we concluded that the TCS NMR studies were not compromised by any non-specific aggregation of the PA63 heptamer-receptor complex at pH values ranging from pH 5–8.

Figure 3. The receptor contact residues with PA domain 2 and 4 that are saturated at pH 8.0. A) A subset of the 1D slices of the [15N,1H] TROSY-HSQC spectra obtained at pH 8.0 highlighting several cross-peaks without saturation at pH 8 (left panels) or with saturation at pH 8 (right panels). Cross-peaks representing PA domain 2 and 4 contact residues are indicated with red and blue labels, respectively. B) A plot of the intensity ratio (I/I_0) of the transferred cross saturation of (PA63)_7 and the interacting residues on the ANTXR2 VWA domain. Significant cross saturation (I/I_0 ≤ 0.75) is indicated with a single asterisk, and highly significant (I/I_0 ≤ 0.5) is indicated with a double asterisk. The errors were calculated by propagating the base-plane noise, which was derived from the signal-to-noise ratios of both control and the saturated spectra and this value was averaged from two duplicate experiments.

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Figure 4. The receptor contact with PA domain 2 is weakened at pH 6.0. **A** A subset of the 1D slices of the $[^{15}N, ^{1}H]$ TROSY-HSQC spectra obtained at pH 6.0 highlighting several cross-peaks without saturation at pH 6 (left panels) or with saturation at pH 6 (right panels). Cross-peaks representing PA domain 2 and 4 contact residues are indicated with red and blue labels, respectively. **B** A plot of the intensity ratio ($I_s/I_0$) from the transferred cross saturation of (PA63)$_7$ heptamer to interacting residues on the ANTXR2 VWA-domain. Significant cross saturation ($I_s/I_0\leq0.75$) is indicated with a single asterisk, and highly significant ($I_s/I_0\leq0.5$) is indicated with a double asterisk. The errors were calculated by propagating the base-plane noise, which was derived from the signal-to-noise ratios of both control and the saturated spectra and this value was averaged from two duplicate experiments.

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Figure 5. Receptor remains bound to PA domain 4 at pH 5.1. A) A subset of the 1D slices of the \([^{15}N,^{1}H]\) TROSY-HSQC spectra highlighting several cross-peaks without saturation at pH 5.1 (left panels) or with saturation at pH 5.1 (right panels). Cross-peaks representing PA domain 2 and 4 contact residues are indicated with red and blue labels, respectively. B) A plot of the intensity ratio \((I/I_o)\) from the transferred cross saturation of \((PA_{63})_7\) to interacting residues on the ANTXR2 VWA domain. Significant cross saturation \((I/I_o \leq 0.75)\) is indicated with a single asterisk, and highly significant \((I/I_o \leq 0.5)\) is indicated with a double asterisk. For all graphs the errors were calculated by propagating the base-plane noise, which was derived from the signal-to-noise ratios of both interleaved experiments. The data was taken from two separate experiments performed at pH 5.1 and pH 5.15 and the average was derived from these experiments.
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A

|     | Unbound pH 8.0 | Bound pH 8.0 | Bound pH 6.0 | Bound pH 5.1 |
|-----|----------------|--------------|--------------|--------------|
| Y46 | ![Image](image1) | ![Image](image2) | ![Image](image3) | N.D.         |
| F47 | ![Image](image4) | ![Image](image5) | ![Image](image6) | ![Image](image7) |
| G53 | ![Image](image8) | ![Image](image9) | ![Image](image10) | ![Image](image11) |
| N57 | ![Image](image12) | ![Image](image13) | ![Image](image14) | ![Image](image15) |
| W59 indole | ![Image](image16) | ![Image](image17) | ![Image](image18) | ![Image](image19) |
| G135 | ![Image](image20) | ![Image](image21) | ![Image](image22) | ![Image](image23) |

B

![Diagram](image24)

Domain 2

GS3

N57

Domain 4

W59

G135

Y46

C

N
Figure 6. Chemical shift changes of the ANTXR2 spectra due to PA binding at several pH values. A) Chemical shift changes are shown for residues Y46, F47, G53, N57, the W59 indole, and G135, by comparing the unbound VWA domain at pH 8 (left panel) with the bound VWA-(PA63)7 complex at different pH values (right panels). N.D: Not Determined; Residue Y46 was not resolved at pH 5.1. B) Chemical shift perturbation of the ANTXR2 VWA domain upon PA63 heptamer binding. Chemical shift perturbations were seen for residues Y46, F47, G53, N57, the W59 indole, and G135 upon PA63 heptamer binding. These residues are modeled onto the crystal structure of the ANTXR2 VWA domain and highlighted in red.

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Figure 7. Model of changes in the PA-receptor contacts that accompany toxin prepore-to-pore conversion. For clarity, only domains 2 and 4 of a single PA monomer are shown with the receptor. 1. The unbound receptor with the PA Domain 4 binding site highlighted in yellow to indicate its “unbound” configuration. 2. The receptor binds to PA domains 2 and 4 forming a molecular clamp that blocks pore formation and inducing a conformational change in PA domain 4 contact residues (indicated with cyan shading). 3. At pH 6 which is similar to the conditions in an early endosomal compartment, the receptor contacts with PA domain 2 are weakened and PA domain 4 contact residues begin to revert back to their “unbound” configuration (indicated with green shading). Additional allosteric effects are also detected at this pH value. 4. At pH 5 which is similar to the conditions in a late endosomal compartment, PA domain 2 is no longer bound to receptor, presumably permitting movement of the 2|2-2|3 loop of PA to mediate pore formation. The receptor remains bound to PA domain 4 after pore formation although certain PA domain 4 binding residues of the receptor revert back to their “unbound” configuration (indicated with yellow shading). The bound receptor may stabilize the structure and/or modify the function of the newly formed pore.

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hemolysin [44,45]. However, the dimensions of these structures are drastically different. Crystallographic studies of the α-hemolysin pore revealed a mushroom structure with a 100 Å diameter cap and a stem region of 32 Å [44]. By contrast, electron micrograph studies of the PA pore stabilized with GroEL revealed a 125 Å diameter cap and a stem region that was almost as long (100 Å) [45]. Santelli and colleagues hypothesized that the receptor might occupy the predicted 75 Å gap between the pore cap structure and the membrane, thereby stabilizing the pore [16]. The results of the present study, which demonstrates that receptor remains bound to PA domain 4 after pore formation is triggered, provide direct support for a possible pore-stabilizing role for the receptor.

A pore-stabilizing role for the receptor is also consistent with results from a previous voltage patch clamp study of ion conductance by the PA pore in whole cells versus artificial membranes. That study indicated that the receptor might influence pore structure since it was associated with altered voltage-dependent inactivation properties of the pore and with altered sensitivity to inhibition by the small molecule inhibitor, TBA [28]. Also consistent with such a role, it has been reported that disulfide bond formation in the extracellular immunoglobulin-like region of ANTXR2, which lies between the membrane and the VWA-domain of the receptor, can influence anthrax toxin pore function [46]. Future studies will aim to uncover how the receptor-PA domain 4 contacts influence the structure and/or function of the anthrax toxin pore.

Materials and Methods

Protein expression and purification

The VWA-domain of ANTXR2 (residues Ser38 to Cys218) was produced as a GST fusion protein [47] in Escherichia coli C43 (DE3) cells (OverExpress). The RIL plasmid of BL21-GolonPlus-RIL cells (Stratagene) was co-expressed in the C43 (DE3) cells due to rare codons within the VWA-domain protein-encoding region. Isotopically enriched 15N, 15N/13C/2H, 15N/13C, and 15N/1H ANTXR2 VWA-domain samples were prepared for NMR studies from 4 liters of E. coli culture grown in standard M9 minimal media with 15NH4Cl at 0.1% (wt/vol), with and without 13C6-glucose or 2H/13C-glucose (0.4% wt/vol). Unlabeled samples of the ANTXR2 VWA-domain were produced in standard Terrific Broth. The cell cultures were grown with carbenicillin (50 µg/ml), chloramphenicol (34 µg/ml), and spectinomycin (50 µg/ml) for plasmid selection.

For the transfer cross saturation (TCS) experiments, the ANTXR2 VWA-domain was produced in 100% D2O-based M9 minimal media supplemented with 15NH4Cl (0.1% wt/vol), 2H/13C6-glucose (0.4% wt/vol) and MEM Vitamin B solution (Sigma). Growth of the C43 (DE3) cells in 100% D2O required acclimating the cells in 5 ml of standard M9 minimal media and slowly acclimating the cells to a 20% increase in D2O levels every 12–24 hours until growth was sustained in 100% D2O-containing medium. A 5 ml sample of cells grown in 100% M9 media was then used to inoculate 1L of 100% D2O M9 media, which was then used for standard isotopic labeling procedures. Once the cell populations had reached an OD600 of 0.75, ANTXR2 VWA-domain expression was induced with 0.5 mM isopropyl β-D-thiogalactopyranoside (IPTG) for 6–8 hours at 37°C. The bacterial cells were then harvested by centrifugation at 8000 × g for 15 minutes at 4°C in a JA-10 rotor and the pellet was resuspended with stirring in a cold 5 mM Tris-HCl pH 8.0, 150 mM NaCl. It was then added drop-wise with stirring into refolding buffer (50 mM Tris-HCl; 150 mM NaCl; 10% vol/vol glycerol) at 4°C, and kept under agitation for one hour. The refolded protein sample was then dialyzed against NMR buffer (50 mM Tris-HCl pH 8.0; 150 mM NaCl). PA83 was purified to 90% homogeneity, as judged by a Coomassie stained SDS-PAGE gel and concentrated using centrifugal filter devices (Vivaspin 15R, Sartorius). The integrity of the refolded protein was demonstrated by a [15N,1H] TROSY-HSQC which was comparable to a control spectrum of a non-denatured 1H,15N ANTXR2 VWA-domain, and through an in vitro toxin neutralization assay as described elsewhere [48].

PA83 was expressed from a pET22b+ vector (Novagen) [49] in Rosetta 2 cells (Novagen) due to rare codon usage and grown at 37°C in Luria Broth containing carbenicillin (50 µg/ml) and chloramphenicol (34 µg/ml). Cells were grown in an OD600 of 1.0, and PA83 expression was induced by addition of 0.5 mM IPTG for 6 h at 25°C. Periplasmic proteins were obtained by osmotic shock by first resuspending pelleted cells in 1L of Buffer B (20% sucrose; 5 mM EDTA; 50 mM Tris-HCl pH 8.0) with stirring at room temperature for ten minutes. The cells were then harvested at 8000 × g for 15 minutes at 4°C in a JA-10 rotor and the pellet was resuspended with stirring in a cold 5 mM MgSO4 solution at 4°C for 15 minutes. This sample was centrifuged again with the same harvesting conditions, protease inhibitor tablets (Roche) were added and the resulting supernatant containing the desired protein was brought up to 50 mM Tris-HCl at pH 8.0 with a 1M stock solution of Tris-HCl pH 8.0. The supernatant was then circulated over an anion-exchange HiTrap QFF column (Amersham Pharmacia) and purified with a gradient of 0M to 1M NaCl in buffer A (50 mM Tris-HCl; pH 8.0) using an AKTA-FPLC system (Amersham Pharmacia). Column fractions containing PA83 were then concentrated and applied to a Hi Load Superdex 26/60 gel filtration column (Amersham Pharmacia) and eluted using gel filtration buffer (50 mM Tris-HCl; 150 mM NaCl; pH 8.0). PA83 was purified to 90% homogeneity, as judged by a Coomassie stained SDS-PAGE gel and concentrated using centrifugal filter devices (Vivaspin 15R, Sartorius). This protocol
was modified from Miller et al. 1999 in order to produce a large scale prep for NMR studies [49].

To generate PA_{83}, by trypsin cleavage [49], the purified PA_{83} sample was concentrated down to 1.5 ml [final concentration 5 mg/ml] for treatment with trypsin-conjugated magnetic beads. Prior to that incubation, 1 ml of the magnetic beads slurry (Mag-Trypsin, Clontech) was washed in gel filtration buffer and then separated from the wash using a microfuge magnetic stand (Promega). The washed beads were then mixed with the purified PA_{83} for 45 minutes at room temperature with constant agitation (Nutator). The trypsin beads were then removed using the magnetic stand and the generated PA_{83} heptamer was purified by gel filtration using a Hi Load Superdex 26/60 gel filtration column (Amersham Pharmacia) and samples were eluted with gel filtration buffer (50 mM Tris-HCl; 150 mM NaCl; pH 6.0). Fractions containing the heptamer were then concentrated by a filtered centrifugal device (Vivaspin 15R, Sartorius). The composition of the PA_{83} heptamer was confirmed by static light scattering/refractive index measurements coupled with size exclusion chromatography (data not shown).

Circular Dichroism spectroscopy of ANTXR2

The ANTXR2 VWA-domain was purified as described, and the samples were concentrated to 10 mg/mL and stored at 4°C. The concentrated samples were diluted into either Buffer A (50 mM Tris-HCl, pH 8.0; 5 mM DTT, 150 mM NaCl, 2.5 mM MgCl_{2}) or into Buffer B (50 mM sodium phosphate buffer, pH 5.0 or 6.0; 150 mM NaCl; 2.5 mM MgCl_{2}) to a final concentration of 25 μM. Supplementary Figures S1 and S2, respectively. The solution was then placed into a 0.1-cm path-length quartz cell (Hellma, Forest Hills, NY). Spectra were acquired using a BioLogic MOS-450 (Molecular Kinetics, Pullman, WA). All measurements were done at 25°C. Spectra were recorded at a wavelength range of 190-260 nm. Three independent experiments were performed with each sample. Raw data were manipulated by smoothing and subtraction of buffer spectra, according to the manufacturer’s instructions.

Assessment of in vitro aggregation

300 μM of ANTXR2 VWA domain was incubated at 37°C for 48 hours in buffers ranging from pH 7, 8, 6, and 5. Buffer A (50 mM Tris-HCl, 150 mM NaCl, 2.5 mM MgCl_{2}) was used for the pH 8 and pH 7 samples. Buffer B (50 mM sodium phosphate buffer, pH 5.0 or 6.0; 150 mM NaCl; 2.5 mM MgCl_{2}) was used for and the pH 6 and pH 5.1 samples. The samples were subjected to a wavelength scan using a Beckman DU 530 Life Science UV/VIS spectrophotometer. The apparent optical density, which is proportional to turbidity, was then analyzed at 340 nm.

Assignment of the ANTXR2 VWA-domain

All NMR experiments were recorded at 310K on a Bruker 700-MHz spectrometer equipped with four radiofrequency channels and a triple-resonance cryoprobe with a shielded z-gradient coil. Measurements were performed on either 15N, 15N/13C, or 15N/15N/13C/1/H, 350 μM labeled ANTXR2 VWA-domain in NMR buffer (50 mM Tris-HCl (pH 8.0); 150 mM NaCl; 0.01% NaN_3; 10/90 D_2O/H_2O), if not otherwise stated. Assignments for the backbone resonances were obtained using data from the following experiments: [1H, 15N] TROSY-HSQC, 3D TROSY-HNCO, 3D TROSY-HNCACB, 3D TROSY-HNCA, and a 3D 15N-edited NOESY-HSQC. NMR data were processed using NMRPipe and analyzed using the Sparky and CARA software packages [34,35].

Transferred cross saturation experiments

Transferred cross-saturation experiments were performed with deuterated, 15N-labeled ANTXR2 VWA-domain in a buffer containing 85-90% D_2O. The final NMR sample contained 350 μM VWA-domain and 35 μM PA_{83} (ratio 10:1) in NMR buffer (50 mM deuterated Tris-HCl pH 8.0 buffer; 5 mM DTT; 150 mM NaCl; 2.5 mM MgCl_{2}; 85% D_2O/H_2O) or low pH NMR buffer (50 mM sodium phosphate buffer pH 6.0 or 5.1; 150 mM NaCl; 2.5 mM MgCl_{2}; 85% D_2O). Experiments were performed at 310K. Selective saturation of the protein was achieved by applying a train of Gaussian shaped pulses prior to the [1H,15N] TROSY-HSQC experiment with the saturation frequency set to 0.8 ppm [32,50]. The experiment was performed in an interleaved manner with a phase sensitive Echo/Antiecho gradient selection. Experiments were performed similarly with 120 scans, 0.5 sec saturation durations, and a relaxation delay of 2.0 s. The experiments were performed with 2048×256 complex points in the 1H and 15N dimensions with spectral widths of 10000 and 2270 Hz, respectively. The spectra were transformed to 2048×256 complex points using zero-filling.

Supporting Information

Figure S1 Far-UV CD spectrum of the refolded ANTXR2 VWA-domain in A) 50 mM Tris-HCl buffer, pH 8.0, B) 50 mM sodium phosphate buffer, pH 6.0, and C) 50 mM sodium phosphate buffer, pH 5.0 (TIF)

Figure S2 The refolded, double-labeled ANTXR2 VWA-domain acts as an efficient receptor decoy that protects RAW264.7 cells against intoxication by anthrax lethal toxin. (TIF)

Figure S3 Selected cross-sections of the HNCACB showing connectivity between several backbone residues of the ANTXR2 VWA-domain. Additional spectra were also obtained for backbone assignment. (TIF)

Figure S4 PA_{83} heptamer forms a SDS-resistant species that is consistent with the pore in the presence of ANTXR2 VWA-domain at an acidic pH of <5.2. PA_{83} at 15 μM was incubated with 150 μM ANTXR2 VWA domain overnight at 37°C at the various pH values shown. The samples were then subjected to SDS-PAGE. As shown, in the presence of ANTXR2 VWA-domain, the SDS resistant PA63 heptameric pore was formed only at pH<5.2. (TIF)

Table S1 Turbidity of the ANTXR2:PA complex at a 10:1 ratio. A sample containing 200 mM ANTXR2 and 20 mM PA_{83} was incubated at 37°C and either pH 8.0 or pH 5.1 for 24 hours and the absorbance values at 340 nm were measured. (TIF)

Table S2 Protein absorbance of ANTXR2:PA after 24 hours incubation at 37°C. A 250 mM PA_{83} was incubated at 37°C and either pH 8.0 or pH 5.1 for 24 hours. The samples were then centrifuged using a table top centrifuge (Eppendorf Centrifuge 5424) at 13,000 rpm for 1 minute and the protein concentrations in the supernatants were measured at A_{280} nm. SDS-PAGE analysis was used to confirm that the PA heptamer remained in solution (data not shown). (TIF)
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Author Contributions

Conceived and designed the experiments: RMP MB RR JATY. Performed the experiments: RMP JMM. Analyzed the data: RMP RR JATY. Contributed reagents/materials/analysis tools: MB RR. Wrote the paper: RMP RR JATY.