Anthrax Toxin Receptor 2
Determinants that Dictate the pH Threshold of Toxin Pore Formation

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INTRODUCTION

Anthrax toxin is a major virulence factor of *Bacillus anthracis* that is thought to cause many of the symptoms observed in anthrax disease. The toxin consists of a single receptor-binding subunit, protective antigen (PA), and two catalytic subunits: lethal factor (LF), a metalloprotease that cleaves and inactivates MKKs [1–3], and edema factor (EF), a calmodulin-dependent adenylate cyclase that converts ATP into cAMP [4,5]. PA binds to either of two host cell receptors, ANTXR1 (anthrax toxin receptor, ATR/TEM8) and ANTXR2 (capillary morphogenesis gene 2, CMG2) [6,7]. After binding the cellular receptor, the 83 kD form of PA, (PA83), is cleaved by cell-surface furin into a 63 kD form, (PA63), which goes on to heptamerize into a ring, or pre-pore [8]. Alternatively, cell surface receptors may engage the cleaved PA63, which exists as monomers or multimers in the blood of infected animals [9,10].

The toxin-receptor complex is internalized by a mechanism involving LRP6 [11], and trafficked to a low pH endocytic compartment where acid pH triggers pore formation and translocation of the catalytic moieties into the cytoplasm [12–14].

PA is comprised of four protein domains with different functions: following PA20 removal and PA63 oligomerization, domain 1 binds EF and/or LF; domain 2 is involved in pore formation, EF/LF translocation, PA63 oligomerization, and receptor binding; domain 3 in PA63 oligomerization; and domain 4 in receptor binding [15]. PA binds to a von Willebrand factor A (vWA) domain that is common to both receptors and most similar to the inserted (I) domains of z-integrins [16]. Similar to z-integrin-ligand interactions, a carboxyate side-chain from PA domain 4 (residue D685) directly coordinates the divergent cation bound at the receptor metal ion dependent adhesion site (MIDAS) [17,18]. The co-crystal structure of the ANTXR2 I domain bound to PA has shown that the contact surface between these two proteins is much larger (~2000 Å2) than typical z-integrin-ligand binding (~1900 Å2) [17,18]. The extensive contact of ANTXR2 with PA domains 2 and 4 most likely accounts for the very high affinity of the ANTXR2 I domain-PA interaction (Kd = 170 pM) [19]. By contrast, the ANTXR1 I domain-PA interaction exhibits a much lower binding affinity (Kd = 130 nM) [20].

In the absence of receptor, PA63 forms pores at neutral pH, while low pH is required for pore formation when the toxin subunit is bound to receptor. Thus it has been proposed that each receptor acts as a molecular clamp to restrict pore formation until the complex encounters an acidic endosomal compartment where low pH induces structural changes in the PA heptameric pre-pore leading to pore formation. In this model, receptor contact with the domain 1 (residues 340–340) of the PA domain 2 membrane insertion loop (pH2–pH3; residues 285–340) restrains pore formation until protonation of PA and/or ANTXR2 residues loosens this interaction to allow domain 2 to undergo a large conformational change and form an extended β-barrel pore [17,18,21,22]. The pH required for pore formation when PA is bound to ANTXR2 (pH ~5.0) is fully one unit lower than when it is bound to ANTXR1 (pH ~6.0), suggesting that the receptor might dictate the subcellular location of pore formation [14,23]. Based upon the difference in pH threshold for pore formation, ANTXR1 is considered to be the weaker molecular clamp, being released more easily from PA than is ANTXR2. Consistent with these pH requirements, ANTXR2, but not ANTXR1, mediated cellular
intoxication is blocked by ammonium chloride (NH₄Cl) treatment, which raises endosomal pH [14].

Prior to this report it was not known which receptor determinants are responsible for dictating the receptor-specific pH thresholds of anthrax toxin pore formation. Here we identify these determinants of ANTXR2 and show that they are involved with binding PA domain 2 and the neighboring edge of PA domain 4.

**METHODS**

**DNA constructs, cell transfections and protein production**

QuickChange mutagenesis (Stratagene) was performed on ANTXR2-EGFP (CMG2-A56-EGFP) [7] or ANTXR1-EGFP (ATR/TEM8-A56-EGFP) [6] plasmids with oligonucleotide primers described in Table S1. For transient receptor expression, ~5×10⁶ CHO-R1.1 cells were transfected with 1 or 3 µg plasmid DNA ANTXR1- or ANTXR2-EGFP and 3 or 3 µg pBSII KS(−) carrier DNA using Lipofectamine 2000 (Invitrogen). All constructs were confirmed by DNA sequencing. WT PA protein was isolated from the periplasm of *E. coli* [24], and purified as previously described [25]. LF₅₆-DTA (the N-terminal, PA-binding domain of LF fused to the catalytic A chain of diphtheria toxin, which kills cells when internalized) was produced as previously described [26].

**Cell intoxication assays**

At 24 hours post-transfection, triplicate samples of ~7×10⁵ cells were plated in 12-well culture dishes and pre-incubated for 1 hour with 30 mM NH₄Cl, or media only. Then cells were incubated with 30 mM ammonium chloride changed every 6 hours, for ~18 hours. Cells were analyzed by flow cytometry and the percentage of live, EGFP-positive cells in the presence of PA and LF₅₆-DTA was determined by the loss of EGFP-positive cells in the presence of PA and LF₅₆-DTA. In all cases, cells were incubated with 30 mM ammonium chloride for 10–20 M PA and 10–10 M LF₅₆-DTA. The pH of the medium was adjusted using Mes buffer (pH 5.2, 5.4, 5.6, 6.0, 6.5, 7.0, 7.2, 7.5, 8.0).

**PA pore formation on cell surfaces**

At 48 hours post-transfection, ~2×10⁶ cells were pre-incubated in supplemented F12 media with dGAB cocktail [50 mM 2-deoxyglucose (Sigma), 10 mM sodium azide (Sigma), 200 mM bafilomycin A1 (Alexis Biochemicals)] for 45 minutes at 37°C. Cells were then washed with PBS buffer with Tris pH 6.8, Tris pH 6.5, MES pH 6.0, MES pH 5.6, MES pH 5.4 and Sodium Acetate pH 5.1 or alternatively MES pH 6.0, MES pH 5.8, MES pH 5.6, MES pH 5.4, Sodium Acetate pH 5.2, and Sodium Acetate pH 5.0 respectively (final pH). Samples were analyzed as previously described [14].

**RESULTS**

**Homolog-scanning mutagenesis of ANTXR2**

We reasoned that non-conserved amino acid residues located at the PA-binding interface were likely to be responsible, at least in part, for the distinct receptor-specific pH thresholds of anthrax toxin pore formation. There are eight such residues in ANTXR2 (A56, N57, Q88, S113, V115, D152, G153, and L154), which correspond to Leu, His, Arg, Leu, Gly, His, Gln, and Asp, respectively, in ANTXR1 [17,18] (Fig. 1A). To test their involvement, each ANTXR2 residue was independently replaced by the corresponding ANTXR1 residue in the context of an ANTXR2-EGFP fusion protein [7]. The altered receptors were expressed in transiently-transfected CHO-R1.1 cells, which lack PA receptors [6]. The pH threshold of PA pore formation was then measured in each case using a previously described assay [14] (Fig. 1B).

Among the mutations in PA domain 2 contact residues, the D152H mutation had almost no effect on the pH threshold of pore formation (data not shown). By contrast, the G153E and L154D amino acid substitutions weakened the ANTXR2 molecular clamp, allowing PA pore formation at a pH value that was 0.6 units higher than that seen with wild-type ANTXR2 (Fig. 1B).

Indeed, when these two mutations were combined (construct ANTXR2-153–154), ANTXR2 was fully converted into a receptor with the pore-inducing properties of ANTXR1 (Fig. 1B). Consistently, the G153E, L154D and 153–154 amino acid substitutions rendered cellular intoxication via ANTXR2, which is normally sensitive to inhibition by NH₄Cl treatment, resistant to that treatment (Fig. 1C).

Among the mutations tested in PA-domain 4 contact residues all had almost no effect on the pH of pore formation (data not shown) with the exception of the Q88R mutation that weakened the molecular clamp, allowing PA pore formation at pH 5.4 (Fig. 1B).

These data indicated that residues G153 and L154, located in the β4-4C loop of the receptor I domain, which contacts PA domain 2, and residue Q88 which contacts PA domain 4, are major determinants of the lower pH threshold requirement associated with ANTXR2.

**Homolog-scanning mutagenesis of ANTXR1**

To determine their sufficiency for the low pH threshold of pore formation associated with ANTXR2, residues G153, L154, and Q88 were introduced at the corresponding positions of ANTXR1. The ANTXR1 155–156 protein, with ANTXR2 residues G153 and L154 replacing the corresponding residues E153 and D156 of ANTXR1, displayed only a modest (0.4 pH unit) shift in threshold of PA pore formation (Fig. 2A and Fig. 2B). Consistently, intoxication via this mutant receptor was still inhibited by NH₄Cl (Fig. 2C). Therefore, these two residues conferred only partial ANTXR2-like properties on the ANTXR1 receptor.

To test whether additional β4-4C loop residues are required to convert ANTXR1 more fully into an ANTXR2-like receptor, ANTXR1 residues 154–159 were replaced with the corresponding amino acids of ANTXR2 (Fig. 2A). The ANTXR1 154–159 protein exhibited an additional 0.2 pH unit shift in the pH threshold of PA pore formation beyond that observed with ANTXR1 155–156 (Fig. 2B). Furthermore, in contrast to ANTXR1 155–156, PA pore formation associated with ANTXR1 154–159 protein was sensitive to ammonium chloride inhibition (Fig. 2C).

These data show that additional residues located in the β4-4C loop of ANTXR2 can further convert ANTXR1 into a receptor, which acts more like ANTXR2.

To test the additional requirement for ANTXR2 residue Q88, this residue was substituted in the ANTXR1 154–159 protein, generating the ANTXR1 R88Q/154–159 protein. This additional change fully converted ANTXR1 154–159 into an ANTXR2-like receptor that restricted pore formation to pH 5.0 or below (Fig. 2B). These results provide an independent line of evidence that β4-4C loop region residues including G153 and L154, as well as Q88, of ANTXR2 are largely responsible for dictating the 1.0 pH unit difference in the receptor-specific pH threshold of toxin pore formation.
Figure 1. Mutagenesis of ANTXR2. (A) Ribbon model of ANTXR2-PA binding interface (UCSF Chimera; PDB # IT6B) [17]. PA domain 2 (PA-D2) and PA domain 4 (PA-D4) are colored light and dark blue, respectively, and the ANTXR2 I domain is colored dark green. The Mn$^{2+}$ cation bound by the ANTXR2 MIDAS is shown in magenta. ANTXR2 residues that are not conserved in ANTXR1 and are involved in PA-D2 and PA-D4 contacts are depicted in orange and red stick representation, respectively. (B) Cells transiently expressing ANTXR2-EGFP or mutant ANTXR2-EGFP receptors treated with dGAB for 45 mins at 37°C were incubated with PA$_{63}$ at 4°C in the presence of dGAB to prevent PA internalization, and pore formation was induced by exposure to low pH buffers before lysing cells. PA pores, which correspond an SDS-resistant high molecular weight species, were detected by SDS-PAGE and immunoblotting with an anti-PA serum followed by an HRP-conjugated secondary antibody. The high molecular weight, SDS-resistant species on the blot representing PA pore are shown. These data are representative examples of at least three similarly performed independent experiments. (C) Triplicate samples of cells expressing WT ANTXR2-EGFP, and ANTXR2-EGFP with G153E, L154D, and 153–154 mutations were treated with PA and LF$_{N}$-DTA in the presence or absence of 30 mM NH$_4$Cl and assayed for cell viability as described in materials and methods.

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Figure 2. Mutagenesis of ANTXR1. (A) Structural diagram of the unique ANTXR2 residues involved in PA domain 2 contact. Residues 152–154 are depicted in orange stick representation with a mesh space-fill overlay, and those residues (155–157) that may affect display of the upstream contact residues are depicted in yellow sticks. The numbered residues of ANTXR2 are shown in white type. PA is depicted in space-fill representation, where C is grey, N is blue and O is red and the numbered residues are shown in black type. Below is an alignment of the primary sequence for this region in ANTXR2 and ANTXR1. The residues depicted in orange and yellow above are underlined; non-conserved changes in ANTXR1 are highlighted in red; and the ANTXR2 G153 and L154 residues shown to be important in Fig. 1 are boxed. (B) PA pore formation was assayed at different pH values on the surfaces of cells transiently expressing ANTXR1-EGFP and mutant ANTXR1-EGFP receptors, as described in Fig. 1B. These data are representative examples of at least three similarly performed independent experiments. (C) Triplicate samples of cells were assayed for intoxication with PA and LF

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Structure-based mutagenesis of conserved receptor residues

At the PA binding interface, ANTXR2 amino acid residues K51, Y119, H121, E122 and Y158 are absolutely conserved with ANTXR1, and residues S87, R111 and E117 are residues are highly similar (Thr, Lys and Asp, respectively, in ANTXR1) [17,18] (Fig. 3A). We reasoned that some of these residues are likely to constitute a conserved core of the molecular clamp mechanism that is shared by both anthrax toxin receptors. To test their involvement in the context of ANTXR2, each of these residues was independently changed to an Ala, and in some cases also to other amino acid side chains.

Among the PA domain 2-contact residues, the most important was Y119: changing this residue either to an Ala or Phe significantly weakened the molecular clamp allowing some toxin pore formation at pH 5.6–6.0 (Fig. 3B). Also, mutant receptors bearing either H121A, H121N, H121E, E122Q, E122R, Y158A, or Y158F amino acid substitutions displayed a 0.2–0.4 pH unit shift in this threshold (Fig. 3B). Among the PA domain 4-contact residues, changing residue E117, either to an Ala, or to the structurally-related Gln side chain, led to a weaker molecular clamp that allowed PA pore formation at pH 5.2–5.4 (Fig. 3B).

Taken together these results indicate that the conserved residues E117, Y119, H121, E122, and Y158, which contact PA domain 2 and the neighboring edge of domain 4, influence the low pH threshold of ANTXR2-associated anthrax toxin pore formation.

DISCUSSION

In this report, receptor determinants that influence the strength of the ANTXR2 clamp which acts to restrict PA pore formation at neutral pH have been defined. Residues G153 and L154, located on the β4-α4 loop of the receptor I domain which interacts with the flexible 340–380 loop of PA domain 2 [17,18] were shown to be the major determinants underlying the more acidic pH requirement for PA pore formation when the toxin is bound to ANTXR2. Substituting both of these residues in ANTXR2 for the corresponding ANTXR1 amino acids gave rise to a much weaker molecular clamp with the pH threshold properties of ANTXR1. Moreover, the ANTXR2 residues G153 and L154 strengthened the ANTXR1 molecular clamp when they were introduced together, especially in combination with residue Q88 and additional β4-α4 loop residues (ANTXR2 residues 152–157), at corresponding positions of the ANTXR1 receptor.

Residue G153 permits a tight turn in the β4-α4 loop and L154 participates in hydrophobic contacts with the side-chains of PA.
residues L340 and T349 (Fig. 2A). The sequence downstream of
the β4-α4 loop of ANTXR2 contains a Pro residue (P155) that
induces a backbone kink and is aliphatic and solvent-exposed,
along with a neighboring Ser residue (S156). However, the
corresponding residues of ANTXR1 are very hydrophobic (F158
and F159, respectively) (Fig. 2A). Thus, it seems likely that this
region of ANTXR1 will adopt a very different conformation from
that of ANTXR2, one that affects the surface presentation of PA
domain 2-contact residues, potentially weakening the toxin
interaction. Indeed, the contacts between the ANTXR1 I domain
and PA domain 2 have yet to be characterized structurally.

Among the conserved receptor residues, the key regulator of
acid pH-dependent pore formation was Y119 of ANTXR2, which
occupies a planar position between domains 2 and 4 of PA (Fig. 4A)
and appears to make hydrophobic and H-bond interactions with
both domains. In particular, the hydroxyl moiety of this residue is
critical since replacement by Phe also gave rise to a dramatic 0.6–
0.8 unit shift in the pH threshold of toxin pore formation (Fig. 4A).

Figure 4. Receptor residues involved in molecular clamp function. (A) Structural diagram of the contribution of individual residues to the receptor
molecular clamp (from the monomeric ANTXR2-PA structure solved at 2.5 Å resolution; PDB # IT6B)[17]. Molecular clamp residues from panel A are
colored black if mutation of the residue caused a 0.4–0.6 pH unit shift in pore formation, or grey if the mutation caused a <0.4 pH unit shift in pore
formation. (B) Left, Two ANTXR2 I domains bound to a PA63 dimer from the heptameric PA-ANTXR2 structure (solved at 4.3 Å resolution; PDB
# ITZN)[18]. PA domains 2 and 4 are depicted in light and dark blue respectively, with the membrane insertion loop (residues 285–340) from PA
domain 2 colored red. Right, One ANTXR2 I domain-PA monomer complex. Structural diagram of the contribution of individual residues to the
receptor molecular clamp. Molecular clamp residues from panel A are colored black if mutation of the residue caused a 0.4–0.6 pH unit shift in pore
formation, or grey if the mutation caused a <0.4 pH unit shift in pore formation.
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It is not known how this hydroxyl moiety is involved, but it may make stabilizing H-bonds with the backbone of PA residue A341 and/or the side-chain of PA residue R659 in the PA prepro-receptor complex (Fig. 4A). Leppla and collaborators have also recently shown that the ANTXR2 Y119H mutation has a major effect upon the pH threshold of PA pore formation [27].

Substituting other conserved residues E117, H121, E122, and Y158 each gave rise to similar 0.2–0.4 unit changes in the pH threshold of toxin pore formation (Fig. 4A). E117 is H-bonded to the PA D683 backbone (Fig. 3A). The imidazole ring of H121 does not make any H-bonds with PA in the co-crystal structure, but this residue is in close proximity to ANTXR2 E122, the PA G342 backbone and PA R344 side-chain (Fig. 3A). A salt bridge between ANTXR2 residue E122 and PA residue R344 was recently implicated in strengthening the ANTXR2 receptor clamp [27]. The hydroxyl moiety of ANTXR2 Y158 appears to be important since a Phe substitution at this position led to a modest decrease in clamp strength. However, it is unknown how this hydroxyl participates because Y158 does not make H-bonds with the monomeric or prepore forms of PA [17,18].

It remains to be determined if the residues implicated in this study affect the affinity of PA-ANTXR2 binding at neutral pH: the long half-life of the complex (15 hours) [19] precluded the use of moderate throughput analysis of all the mutants. Indeed, even SPR analysis is not sensitive enough to accurately measure the apparent Kd of the toxin-wild-type receptor complex and a more sensitive FRET-based assay involving the use of AF488 (donor fluorophore)-labeled PA and AF546 (acceptor fluorophore)-labeled ANTXR2 I domain had to be used in that study. Consequently, previous measurements obtained by binding PA to cell surfaces for 1 hour at 4°C [27,28] likely reflect only association rate differences between different mutant ANTXR2 proteins. Since these experiments are not performed under equilibrium conditions, they also likely underestimate the actual PA-binding affinities of wild-type and mutant forms of ANTXR2.

The ANTXR2 residues that form the functional core of the receptor clamp, as well as those that dictate the receptor-specific pH thresholds of pore formation, map to the region that contacts PA domain 2 and the neighboring edge of PA domain 4 (Fig. 4A). These data therefore provide genetic evidence that receptor release of these two PA regions is likely necessary to allow the unraveling of the 285–340 region of PA domain 2 involved in pore formation (Fig. 4B). Our results also illuminate the genetic basis for the previous observation that there was a one unit difference in the pH threshold at which ANTXR2 and ANTXR1 allowed toxin pore formation, and support a model where toxin pore formation may be occurring in different endosomal compartments depending on the receptor bound [14]. Previously it was noted that the pH profile of anthrax toxin pore formation was consistent with the titration of histidines and that there are several histidines (H299, H304, H310, and H336), located in the 285–340 region of PA which forms the membrane-spanning pore [17,18]. Moreover, it was suggested that ANTXR2 residue H121 might be a key component of this triggering mechanism since this residue is located at the PA domain 2 binding interface and is conserved in ANTXR1 [17,18]. However, the current study excludes a major role for ANTXR2 residue H121 since its substitution by other amino acids had only a modest effect on the pH of pore formation. The precise molecular changes in PA-receptor complexes, which accompany anthrax toxin pore formation in response to a low pH stimulus, remain to be determined and are under investigation.

SUPPORTING INFORMATION

Table S1 Mutagenesis Primers

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Author Contributions

Conceived and designed the experiments: GR JY HS. Performed the experiments: HS JM. Analyzed the data: RC GR JY HS JM DL. Wrote the paper: GR JY HS JM.

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