

Babesia divergens and *Neospora caninum* apical membrane antigen 1 structures reveal selectivity and plasticity in apicomplexan parasite host cell invasion

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Abstract: Host cell invasion by the obligate intracellular apicomplexan parasites, including *Plasmodium* (malaria) and *Toxoplasma* (toxoplasmosis), requires a step-wise mechanism unique among known host–pathogen interactions. A key step is the formation of the moving junction (MJ) complex, a circumferential constriction between the apical tip of the parasite and the host cell membrane that traverses in a posterior direction to enclose the parasite in a protective vacuole essential for intracellular survival. The leading model of MJ assembly proposes that Rhoptry Neck Protein 2 (RON2) is secreted into the host cell and integrated into the membrane where it serves as the receptor for apical membrane antigen 1 (AMA1) on the parasite surface. We have previously demonstrated that the AMA1–RON2 interaction is an effective target for inhibiting apicomplexan invasion. To better understand the AMA1-dependant molecular recognition events that promote invasion, including the significant AMA1–RON2 interaction, we present the structural characterization of AMA1 from the apicomplexan parasites *Babesia divergens* (*BdAMA1*) and *Neospora caninum* (*NcAMA1*) by X-ray crystallography. These studies offer intriguing structural insight into the RON2-binding surface groove in the AMA1 apical domain, which shows clear evidence for receptor–ligand co-evolution, and the hyper variability of the membrane proximal domain, which in *Plasmodium* is responsible for direct binding to erythrocytes. By incorporating the structural analysis of *BdAMA1* and *NcAMA1* with existing AMA1 structures and complexes we were able to define conserved pockets in the AMA1 apical groove that could be targeted for the design of broadly reactive therapeutics.

Keywords: parasite invasion; Apicomplexa; moving junction; structural plasticity; neosporosis; babesiosis

Abbreviations: AMA1, apical membrane antigen 1; *B. divergens*, *Babesia divergens*; CSS, complexation significance score; DI, Domain I; DII, Domain II, DIII, Domain III; 5FAM, 5-carboxyfluorescein; HBS, HEPES buffered saline; MJ, moving junction; MR, molecular replacement; *P. falciparum*, *Plasmodium falciparum*; *P. vivax*, *Plasmodium vivax*; RON, rhoptry neck protein; RON2sp, RON2 synthetic peptide; *T. gondii*, *Toxoplasma gondii*; *N. caninum*, *Neospora caninum*.

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Introduction

Phylum Apicomplexa harbors more than five thousand parasitic protozoan species, many of which cause serious morbidity and mortality in humans and animals worldwide. Some of the most prevalent apicomplexans are *Plasmodium*, *Toxoplasma*, *Babesia*, and *Neospora*, the etiological agents of malaria, toxoplasmosis, babesiosis, and neosporosis, respectively. More than 250 million cases of malaria every year result in at least 1 million deaths,^{1,2} and up to a third of the world's population is chronically infected with *Toxoplasma*.^{3,4} Babesiosis is one of the most prevalent infections of free-living animals, particularly cattle,^{5,6} while *Neospora* is an important veterinary pathogen causing premature abortion in cattle and fatal neurological defects in dogs.^{7,8} Despite the diversity of hosts and disease outcomes, a unique feature of all apicomplexans is their obligate intracellular lifestyle. Consequently, these resourceful parasites have developed highly sophisticated mechanisms to invade a wide range of host cells.^{5,9–11}

Modes of host cell recognition and attachment vary across the phylum and likely play a key role in defining cellular tropism. However, the process of active invasion appears to be highly conserved and a major contributor to the virulence of the parasites.¹² Importantly, invasion is largely independent of traditional host cell uptake processes and proceeds with the formation of the moving junction (MJ),^{13–15} a ring-like protein structure formed at the interface between parasite and host cell. During invasion, the parasite propels its way through the MJ resulting in its encapsulation in a protective vacuole within the host cell.¹⁶ Thus, assembly of a functional MJ is crucial to parasite survival. It has recently been postulated that formation of the MJ is initiated when Rhoptry Neck Protein 2 (RON2) is secreted from the parasite's rhoptry organelles in a preformed complex with RONs 4 and 5, and also RON8 in coccidia (*Toxoplasma*, *Neospora*, and *Eimeria*).^{15,17–18} This complex is discharged into the host cell, where the leading MJ model suggests that RON2 integrates into the host cell membrane and serves as the receptor for apical membrane antigen 1 (AMA1) displayed on the parasite cell surface.^{19–23} Thus, it has been suggested that apicomplexan parasites are capable of providing both ligand and receptor to actively invade host cells. Although a recent study found that functional junctions could form in the absence of detectable AMA1,²⁴ other studies, such as those showing that RON2-derived peptide inhibitors of the AMA1–RON2 interaction inhibit MJ assembly and parasite invasion,^{25–27} provide strong support for the leading MJ model with AMA1 as a structural component of the junction. Recent evidence suggests an altered composition of the *Plasmodium* sporozoite MJ, as well as the potential for yet unidentified proteins to participate in host

cell invasion.²⁴ Overall, the importance of the AMA1–RON2 interaction is clear, but the specific role in the invasion process remains ambiguous—whether this interaction provides the scaffold for the MJ or initiates a signal within the parasite for productive invasion will require further investigation.

Homologues of AMA1 exist in nearly every apicomplexan parasite,²⁸ with the conserved role of RON2 binding as demonstrated in *Toxoplasma* and *Plasmodium*,^{20,26} but also suspected roles in immune system evasion, early invasion events such as host cell adherence, engagement of the parasite motor complex, and signaling.^{24,29–39} The importance of AMA1 in host cell invasion by the parasite is supported by the inability to generate *ama1* knockouts in *Plasmodium* merozoites⁴⁰ and *Toxoplasma* tachyzoites,⁴¹ as well as the severely attenuated invasion of conditional *ama1* knock-downs in these parasite lifecycle stages.^{24,42} Collectively, these studies support the classification of AMA1 as a leading malarial vaccine candidate, which initially prompted the structural characterization of the AMA1 ectodomain from *Plasmodium falciparum* (*P. falciparum*),^{43–44} *Plasmodium vivax* (*P. vivax*),⁴⁵ and *Toxoplasma gondii* (*T. gondii*).⁴⁶ Each structure revealed a stacked three-domain architecture (DI, DII, and DIII) with a hydrophobic groove framed by a network of divergent surface loops localized to the apical surface of DI. These structural features are targeted by invasion inhibitory antibodies and peptides,^{47–50} which presumably function through blocking AMA1–RON complex formation.^{51,52} Subsequent studies localized the AMA1 binding region on RON2 to its C-terminal portion^{26,53} and revealed the structural basis for complex formation.^{25,27}

From our recent structural characterization of the AMA1–RON2 complexes from *T. gondii* and *P. falciparum*,^{25,27} we proposed a binding model where the N-terminal helix of the RON2 peptide is seated at one end of the AMA1 hydrophobic groove and extends through ordered coil to a disulfide-bound beta-hairpin structure (cystine loop), forming a U-shape in the apical groove. However, intriguing differences in anchor points of the AMA1–RON2 interaction were also observed and clearly represented a co-evolution between ligand and receptor.^{25,27} These studies highlighted the complexity of defining the structural basis of cross species and strain selectively, which remains somewhat elusive. To more thoroughly define the structural features of AMA1 that promote host cell invasion, we present the crystal structures of AMA1 from *B. divergens* and *N. caninum*. In addition to addressing the contributions of structural plasticity and supporting the design of broadly reactive inhibitors to disrupt the AMA1–RON2 interaction, our study reveals intriguing insight into the highly variable DIII domain, which in the case of *Plasmodium* has been shown to directly coordinate a receptor on erythrocytes.³³

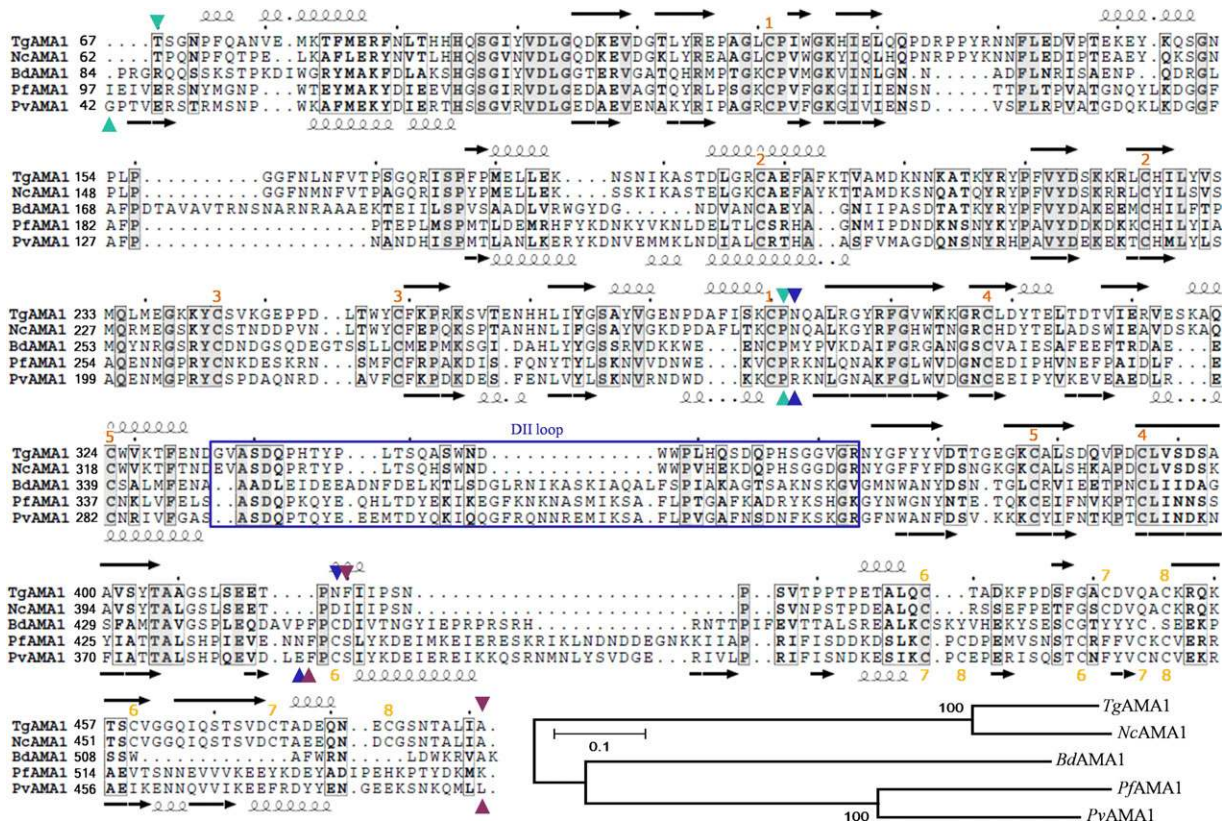


Figure 1. Sequence analysis of *BdAMA1* and *NcAMA1* compared with AMA1 proteins for which structures have been determined. Sequences are numbered from the initiation methionine in the signal sequence. Top secondary structure elements refer to *TgAMA1*, bottom elements refer to *PvAMA1*. Cyan arrows indicate DI limits; blue—DII, purple—DIII. Orange numbers indicate disulfide connectivity, with top DIII numbers referring to *Tg/NcAMA1* and bottom DIII numbers referring to *Pf/PvAMA1*. The highly variable DII loop is indicated by a blue box. Bottom—phylogenetic tree of the five structurally characterized AMA1 proteins generated in MEGA4 with 500 bootstrap replicates from a MUSCLE alignment of the fully processed ectodomains. Domain boundaries of the fully processed *Bd/NcAMA1* ectodomains were defined based on the paradigm established for previously characterized AMA1s.^{37,43,45,46,56} *BdAMA1* DI spans residues from Pro84 to Pro305 (numbering based on the initiation methionine in the signal sequence), DII from Met307 to Pro445 and DIII from Phe446 to Lys523, while *NcAMA1* DI spans residues from Thr62 to Pro281, DII from Asn282 to Ile410 and DIII from Ile411 to Ala481.

Results

Sequence analysis reveals distinct features of *Bd/NcAMA1*

To establish domain boundaries and regions of divergence in *B. divergens* AMA1 (*BdAMA1*) and *N. caninum* AMA1 (*NcAMA1*) we generated sequence alignments comparing the fully processed ectodomains of *BdAMA1* and *NcAMA1* with the structurally characterized AMA1s from *P. falciparum* (*PfAMA1*), *P. vivax* (*PvAMA1*) and *T. gondii* (*TgAMA1*; Fig. 1). The alignment reveals that *NcAMA1* is closely related to *TgAMA1*, with sequence identity of 75% over the entire ectodomain, while *BdAMA1* appears intermediate between *Tg/NcAMA1* (25% sequence identity) and *Pf/PvAMA1* (30% sequence identity). Phylogenetic analysis supports both the close relationship between *TgAMA1* and *NcAMA1* and the distant clustering of *BdAMA1* with *Pf/PvAMA1* [Fig. 1 (bottom)]. For both *BdAMA1* and *NcAMA1*, the DI and DII cystine network is conserved, suggesting a conserved structural core (Fig. 1).

However, *BdAMA1* displays unique features including large insertions and deletions corresponding to apical surface loops functionally important in related AMA1s and two fewer conserved cysteine residues in DIII, implying that it will not be able to form the ultra stable cystine knot observed in all AMA1 DIII structures to date^{43,45,46,54,55} (Fig. 1). It is noteworthy that, while the overall sequence identity between *TgAMA1* and *NcAMA1* is high, *NcAMA1* displays divergence in residues shown to be critical in the interaction of *Tg* AMA1 with a *Tg*RON2 synthetic peptide (*Tg*RON2sp)²⁵ (Fig. 1). To investigate the structural consequences of these unique sequence features, *BdAMA1* and *NcAMA1* were characterized using X-ray crystallography.

Overall structure of *Bd/NcAMA1* ectodomains

Soluble forms of the fully processed three domain (DI, DII, and DIII) ectoplasmic region of *BdAMA1* and *NcAMA1* were produced recombinantly in insect cells, purified to homogeneity using nickel affinity,

size exclusion, and ion exchange chromatography, and set in crystallization screens. Refinement of preliminary crystals required recombinant protein produced in the presence of tunicamycin to reduce N-linked glycosylations. *BdAMA1* crystallized as thin clusters of sheets in space group $P2_12_12$ with one molecule in the asymmetric unit, while *NcAMA1* crystallized as thin sheets in $C2$ with four molecules in the asymmetric unit. The final model of *BdAMA1* incorporates residue 93 through 510, with two apical surface loops and a portion of the DII loop disordered. *NcAMA1* is modeled between residues 62(chain A)/64(chains B, D, and E) and 473, with localized regions of disorder near the tip of the DII loop and in the DII-DIII linker. Four-fold non-crystallographic symmetry averaging with *NcAMA1* was required to define the structure of several surface loops. The overall structure of each chain is largely conserved with root-mean-square deviations (r.m.s.d) relative to chain A of 0.50 Å over 390 C α atoms (chain B), 0.47 Å over 394 C α atoms (chain D) and 0.56 Å over 389 C α atoms (chain E). Chain A was the most extensively modeled (398 of 431 total residues), and used for all subsequent structural analyses unless otherwise noted.

The similarity of the *BdAMA1* and *NcAMA1* structures to previously characterized AMA1 proteins was probed using a DALI search,⁵⁷ which revealed that while *BdAMA1* is clearly structurally homologous with *Pf/PvAMA1* (Z-scores of 34–36), it is unexpectedly also closely related to *TgAMA1*, with a Z-score of 32. As expected, *NcAMA1* shows a high degree of structural similarity with *TgAMA1* (Z-score 60), and lower, but still significant, similarity to *Pf/PvAMA1* (Z-scores of 32–34). These observations confirm that *BdAMA1* is a “hybrid” between *TgAMA1* and *Pf/PvAMA1*, consistent with the distant phylogenetic clustering with *Plasmodium* AMA1s (Fig. 1). Intriguingly, *NcAMA1* DI and DII are not classified as PAN module containing domains, similar to *TgAMA1*,⁴⁶ while DI and DII of *BdAMA1* are recognized as PAN domains, as was identified for *PfAMA1* and *PvAMA1*^{43,45} (highest Z-score with PAN-containing protein—*Tg/Nc/Bd/Pf/PvAMA1*: 3.9/3.9/8.0/7.1/4.1). However, both *NcAMA1* and *BdAMA1* contain similar arrangements of secondary structure in DI and DII with a central alpha helix in each surrounded by a curved beta sheet of four anti-parallel strands, with the majority of insertions, deletions, and structural variations localized to surface loops (Fig. 1).

Divergence in the apical groove reveals structural consequences for the DII loop

The *BdAMA1* and *NcAMA1* ecto domains are approximately equivalent in size and predicted to extend more than 80 Å from the parasite cell membrane [Fig. 2(A)]. Both AMA1 proteins are struc-

tured with DI assembled on top of DII, a loop of DII extending up the side of DI to form an integral part of the apical surface, and DIII forming the membrane proximal base [Fig. 2(A)], analogous to the other structurally characterized AMA1 proteins.^{43,45,46} Both *BdAMA1* and *NcAMA1* present a hydrophobic groove that extends roughly 40 Å across the length of the apical surface [Fig. 2(B)]. The tip of the *BdAMA1* DII loop is disordered [Fig. 2(B top)], which is not surprising given that the corresponding region is completely disordered in *PvAMA1*,⁴⁵ only partially modeled in *PfAMA1*,⁴³ and displaced from the apical groove in *Tg/PfAMA1* upon *Tg/PfRON2* binding.^{25,27} In contrast, the DII loop is relatively well modeled in *NcAMA1*, likely due to its truncated size (15 residues shorter than *BdAMA1*) [Figs. 1 and 2(B bottom)]. However, despite the truncated structure, the *NcAMA1* DII loop is markedly more mobile than observed in *TgAMA1* [Fig. 2(C)] indicating that loop length is only partially responsible for order; the interactions formed by the tip of the DII loop also appear to play an influential role in its organization. The tip of the *NcAMA1* DII loop is anchored in the apical groove by a pair of tryptophan residues, Trp347 and Trp348, which bury into pockets on either side of a central serine, Ser224 [Fig. 2(C left, top middle)]. This serine does not appear to serve as an effective anchor for the DII loop, as the two pockets are not well separated, leading to two observed conformations of Trp348 [Fig. 2(C left, top middle)]. The observed organization is in contrast with apo*TgAMA1*, where the DII loop is well ordered and modeled in its entirety, with the two Trp residues (Trp353, Trp354) anchored in a hydrophobic pocket bifurcated by Tyr230 [Fig. 2(C bottom middle, right)].⁴⁶ The Ser-Tyr substitution in *NcAMA1* may be advantageous as it could reduce the energy barrier for DII loop rearrangement upon RON2 binding.

DIII displays the most structural divergence of the three ectoplasmic AMA1 domains

Although the apical surface of AMA1 has a defined role in RON2 coordination,^{25,27} the membrane proximal DIII has multiple potential roles, including functioning as an adhesin in *Plasmodium* attachment to erythrocytes³³ and as a conduit for any signal passed from the ectoplasmic region through to the intracellular domain.³⁵ The divergence of DIII indicated by sequence alignments (Fig. 1) is clearly retained at the structural level. *NcAMA1* DIII consists of two beta-strands layering across the bottom of DI/DII and a short C-terminal helix, which is stabilized by a cystine knot [Fig. 3(A left)]. *BdAMA1* DIII is comprised entirely of beta strands and extended connecting loops that form a single sheet beneath DI/DII [Fig. 3(A middle)]. The *NcAMA1* and *TgAMA1* minimalist DIII [Fig. 3(A left)] are highly

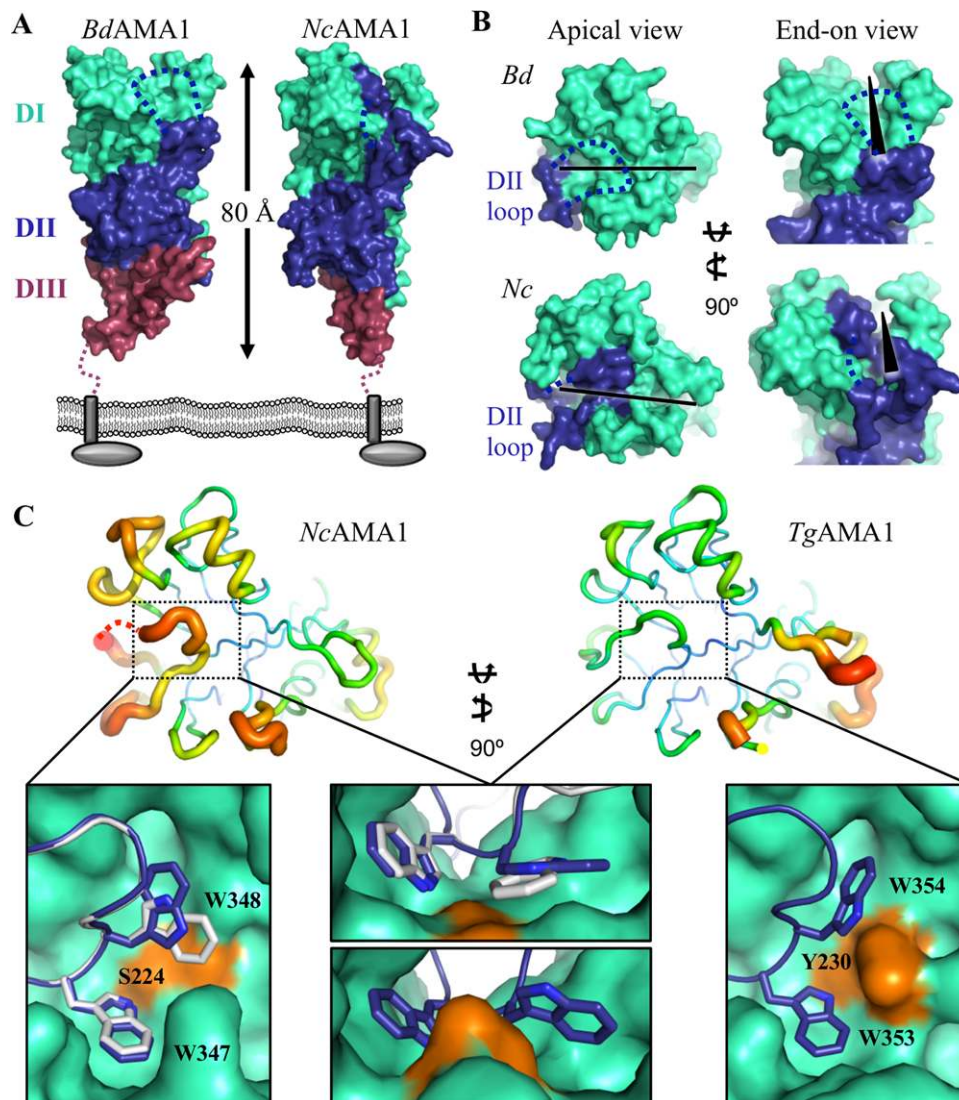


Figure 2. Initial structural characterization of *Bd*AMA1 and *Nc*AMA1. (A) Three-domain architecture of *Bd*AMA1 and *Nc*AMA1 shown in the predicted orientation to the parasite membrane, with DI cyan, DII blue, and DIII purple. Dotted blue line indicates the un-modeled portion of the DII loop. (B) Apical (left) and end-on (right) surface views of *Bd*AMA1 (top) and *Nc*AMA1 (bottom), colored as in (A). The DII loop extends up the side of DI to form part of the apical surface. Among the apical surface loops, a deep groove is present (black bar/wedge). (C) Top—B-factor analysis of the *Nc*AMA1 and *Tg*AMA1 apical surfaces, showing a relatively higher mobility in the DII loop (black box) of *Nc*AMA1 (left) compared with *Tg*AMA1 (right). Tubes colored blue to red and with thin to thick lines, indicating the scale of order to disorder within each structure. Bottom—the DII loop of *Nc*AMA1 is held into the apical groove (cyan) by a pair of tryptophans on either side of a central serine (orange); overlay of *Nc*AMA1 chains A (blue) and E (white) illustrates the two orientations of Trp348 (left, top middle). The DII loop of *Tg*AMA1 (blue) is anchored into the apical groove (cyan) by a pair of Trp residues burying into pockets on either side of a central tyrosine (orange) (bottom middle, right; PDB ID: 2X2Z). All molecular figures generated in PyMol. An interactive view is available in the electronic version of the article.

conserved, overlaying with an r.m.s.d. of 0.69 Å over 51 C α atoms (>80% sequence identity). Importantly, both the small *Tg*/*Nc*AMA1 DIII and the large *Pv*AMA1 DIII [Fig. 3(A right)] adopt the structurally ultra-stable cystine knot motif, with a single disulfide bond threading through two others within DIII. In contrast, *Bd*AMA1 DIII contains only two disulfide bonds and is noticeably less compact, forming a

more extended layer across the base of DI and DII [Fig. 3(A middle)]. This is evident from a comparison of the DIII—DI/DII interfaces. *Bd*AMA1 DIII buries 1958 Å² of surface area with DI/DII, with a complexation significance score (CSS)⁵⁸ of 0.44, while *Nc*/*Tg*AMA1 DIII, which is roughly the same size, buries 2236 Å² with DI/DII, with the highest CSS of 1.0 indicating a more intimate association between

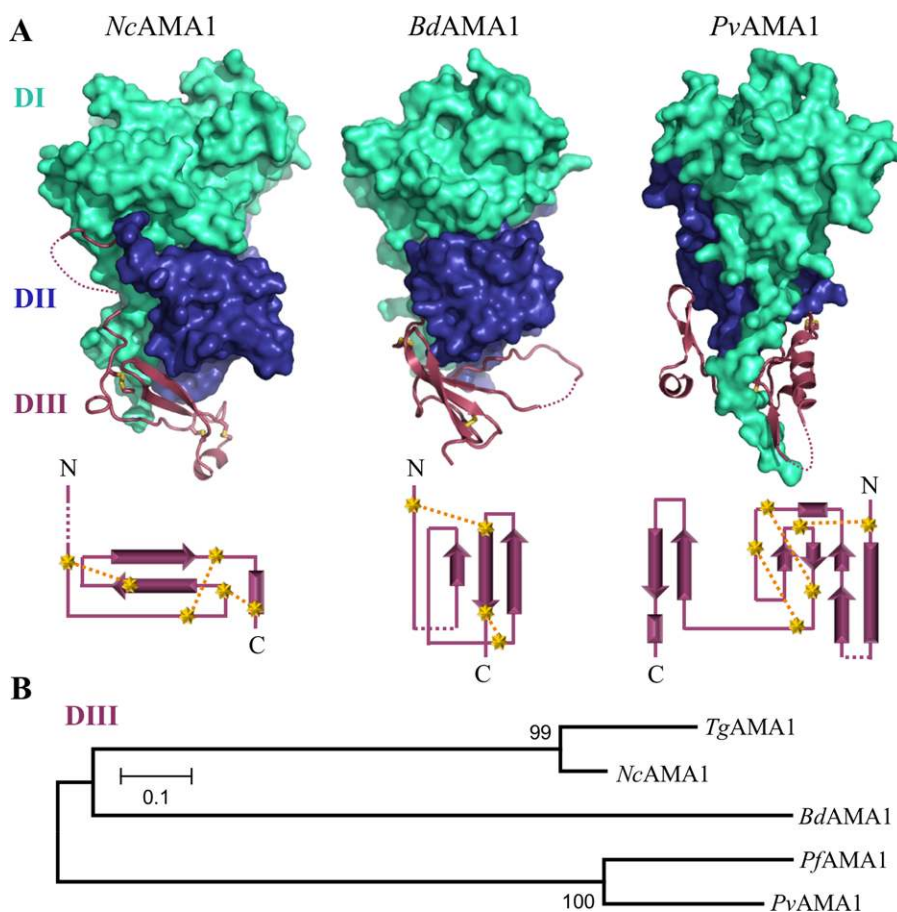


Figure 3. Extreme divergence in DIII of AMA1. (A) Top—solid surfaces of *NcAMA1*, *BdAMA1*, and *PvAMA1* (PDB ID: 1W8K) DI (cyan) and DII (blue), with secondary structure of DIII (purple). Disulfide bonds in DIII shown as yellow sticks. Bottom—corresponding DIII topology diagrams with cysteines depicted as yellow stars, disulfide bonds as yellow dotted lines, and disordered regions of DIII as purple dotted lines. (B) Phylogenetic analysis in MEGA4 with 500 bootstrap replicates based off of MUSCLE alignments of DIII domains (as indicated by purple arrows in Fig. 1) supports divergence of the DIII domains and distant clustering of *BdAMA1* DIII with *Nc/TgAMA1* DIII.

the domains. By comparison, *PvAMA1* DIII, which is 1.5 times the size, buries 3325 Å² with DI/DII with a CSS of 1.0 and appears to have an additional role in stabilizing the DI N-terminal extension. Construction of a phylogenetic tree of the DIII domains for the five structurally characterized AMA1s supports the observation that *BdAMA1* DIII is more closely related to *Nc/TgAMA1* DIII [Fig. 3(B)].

NcAMA1* displays cross reactivity with a C-terminal region of *TgRON2

With the flexible DII loop removed, the apical surfaces of *BdAMA1* and *NcAMA1* maintain a deep groove, similar to the mature receptor binding grooves of *TgAMA1*²⁵ and *PfAMA1*,²⁷ that is likely capable of accommodating RON2 [Fig. 4(A, black bar)]. The similarity of *NcAMA1* and *TgAMA1* coupled with the divergence between *BdAMA1* and *TgAMA1* provides an excellent opportunity to investigate the roles of divergent substructures in the RON2 binding region. Consequently, we tested the capacity of the *TgRON2* peptide to bind *BdAMA1*

and *NcAMA1* in a native gel electrophoresis assay [Fig. 4(B)—multiple AMA1 bands are due glycosylation heterogeneity]. Although no interaction between *BdAMA1* and the *TgRON2* peptide was observed [Fig. 4(B left)], *NcAMA1* and *TgRON2sp* formed a stable complex [Fig. 4(B middle)] at the same molar ratio observed for *TgAMA1* [Fig. 4(B right)].

To probe cross-reactivity at a more detailed level, a fluorescence polarization assay was developed to assess binding between a fluorescently labeled *TgRON2* peptide [5-carboxyfluorescein (5FAM)—*TgRON2sp*] and *TgAMA1*, *NcAMA1*, and *BdAMA1*. Similar to previous studies, the EC₅₀ value of the *TgAMA1*—5FAM-*TgRON2sp* interaction was determined to be in the low nanomolar range (51.7 ± 5.9 nM; [Fig. 4(C)]). As expected from our structural predictions, the *NcAMA1*—5FAM-*TgRON2sp* binding event occurs with an EC₅₀ value similar to *TgAMA1* (57.4 ± 3.8 nM; [Fig. 4(C)]). In contrast, no binding could be detected between *BdAMA1* and 5FAM-*TgRON2sp* (ND; [Fig. 4(C)]) consistent with the native gel shift assay.

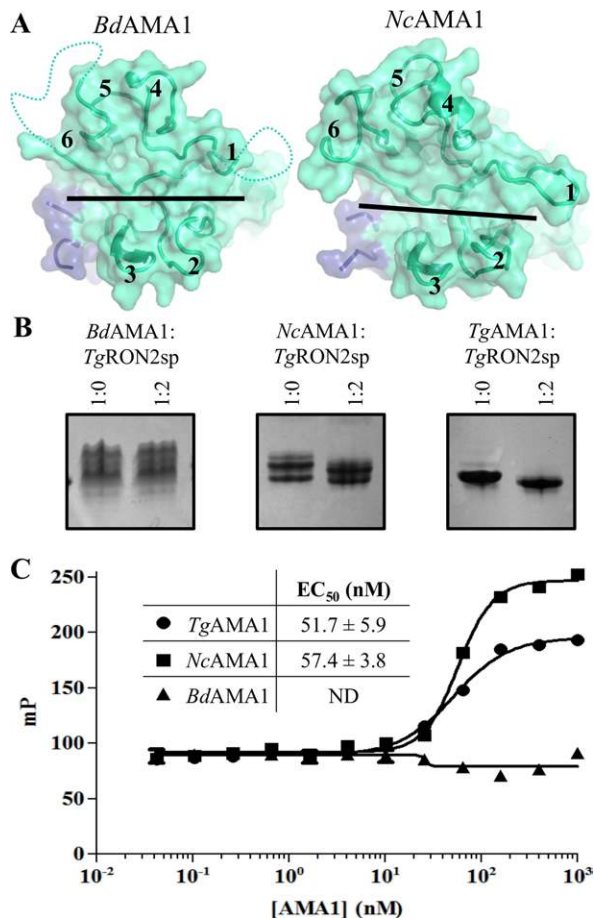


Figure 4. Cross-genera reactivity in the AMA1-RON2 interaction. (A) Apical surface views of *BdAMA1* (left) and *NcAMA1* (right) with the DII loop removed to the point of corresponding disorder in the *TgAMA1*-*TgRON2sp* co-structure. DI/DII are shown as semi-transparent cyan/blue surfaces, with secondary structure of loops visible, and un-modeled loops indicated by dotted lines. Black bars indicate the apical groove and black numbers represent the apical surface loops as defined by previous AMA1 structures. (B) Native gel electrophoresis assays reveal a level of structural plasticity in the AMA1-RON2 interaction. *BdAMA1* does not interact with *TgRON2sp* (left), but *NcAMA1* is able to bind *TgRON2sp* (middle) at the same molar ratio as *TgAMA1* (right). (C) Fluorescence polarization assay with recombinant *TgAMA1*, *NcAMA1*, and *BdAMA1*, and 5FAM-*TgRON2sp*. Error bars indicate SEM for triplicate samples. EC₅₀ values calculated from a sigmoidal nonlinear regression model in GraphPad Prism 5.0. mP—millipolarization units; ND—not determined.

Discussion

Hyper-variability of DIII implies divergent functions

Based on sequence alignments (Fig. 1) and structural analyses (Fig. 3), DIII displays the most extensive divergence of the three AMA1 ectoplasmic domains. *NcAMA1* and *TgAMA1* DIII are notably compact and stabilized by the cystine knot, and may serve a simple function such as orienting the apical

surface toward the host cell, or a more complex function in signal transduction. However, DIII of *P. falciparum* AMA1 has been implicated in attachment to the erythrocyte surface through the membrane protein Kx.³³ In contrast, the attachment of *BdAMA1* to erythrocyte surfaces is Kx-independent and has a similar adhesion profile to *Plasmodium yoelii* AMA1 DI/II: neuraminidase resistant, but trypsin and chymotrypsin sensitive.^{37,59} The divergence between DIII of *BdAMA1* and *PvAMA1* (which likely has a similar structure to *PfAMA1* DIII) is therefore not surprising as *Plasmodium* and *Babesia* have similar cellular tropisms, but AMA1-mediated attachment appears to occur through variable AMA1 domains and distinct erythrocyte receptors.^{33,37,60,61} The loss of the cystine knot in *BdAMA1* DIII leading to a more extended structure [Fig. 3(A middle)] suggests it does not have a major role in orienting DI/DII. Ultimately, additional studies are required to define the role of the DIII domains, though the structural variability suggests the potential for a broad range of functions.

Structural plasticity in the AMA1-RON2 interaction

Prior to this study cross reactivity between AMA1 from one species and RON2 from a different species had not been identified, which led us to perform a detailed comparative structural analysis to understand the basis for both plasticity and selectivity in this important interaction.

In the *T. gondii* and *P. falciparum* AMA1-RON2 interactions, when the AMA1 DII loop is displaced a clear pocket is revealed formed by a trio of tyrosine residues (Tyr110/142, Tyr213/234, and Tyr215/236) into which a conserved RON2 proline (Pro1309/2033) docks [Fig. 5(A,B)]. Although *BdAMA1* has a histidine (His131) substituted for Tyr110/142, the remaining two tyrosine residues (Tyr233 and Tyr235) are conserved, and together these three residues retain the generally conserved architecture of the surface pocket [Fig. 5(B 1 top)]. The aromatic imidazole ring of the histidine residue aligns with the corresponding benzene ring of Tyr110/142 and likely maintains a stacking interaction with the analogous *BdRON2* proline (Pro953). In *NcAMA1*, as for *TgAMA1*, this pocket was previously occupied by the DII loop Trp353 [Fig. 2(C)]. The *TgAMA1* tyrosine trio is conserved in both sequence and structure with *NcAMA1* (Tyr104, Tyr207, and Tyr209) and the *TgRON2* proline is conserved in sequence with *NcRON2* [Fig. 5(A,B 1 bottom)], suggesting a similar anchor point is exploited during complex formation. Notably, in the co-structure of *PfAMA1* with the invasion inhibitory phage display peptide R1, a Leu residue docks into this same pocket,²⁷ suggesting that occupation of the Tyr trio defined pocket is an important component of complex formation,

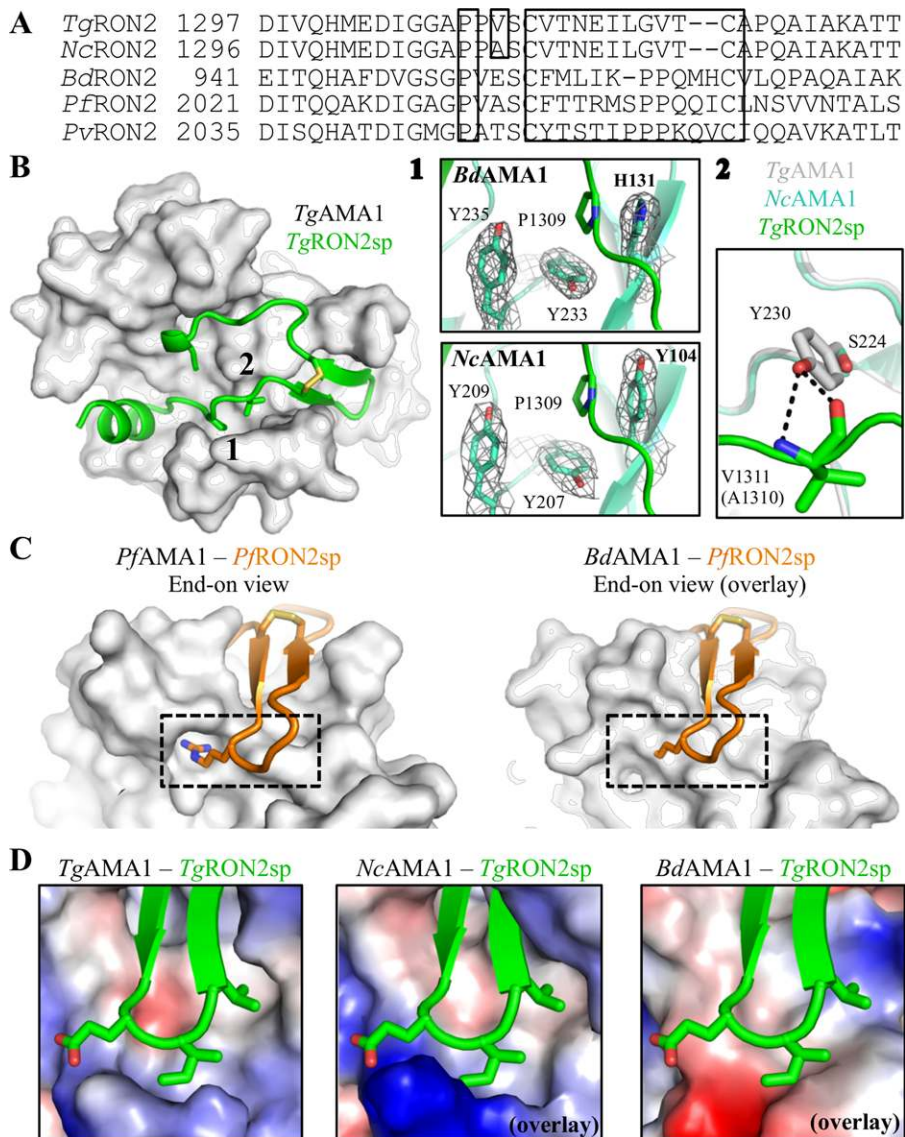


Figure 5. Structural plasticity and specificity of the AMA1–RON2 interaction. (A) Sequence alignment of RON2 sequences for the regions that align with *Tg*RON2sp, with boxes around the conserved RON2 Pro, the single difference between *Tg*RON2 and *Nc*RON2 in this region, and the cystine loop. (B) *Tg*AMA1 apical surface (gray) with *Tg*RON2sp secondary structure (green) (PDB ID: 2Y8T) and residues of interest shown as sticks; black numbers indicate conservation and divergence with *Bd*/*Nc*AMA1 under investigation. 1—Top: overlay of *Tg*RON2sp onto *Bd*AMA1 shows that the AMA1 Tyr trio that accommodates a key RON2 Pro (green sticks) is substituted in *Bd*AMA1 with a Tyr/Tyr/His (cyan sticks) that conserves the architecture of the pocket. Bottom: overlay of *Tg*RON2sp (green) onto *Nc*AMA1 shows that the complete Tyr trio is conserved in *Nc*AMA1 (cyan sticks). Gray mesh: 2Fo-Fc electron density maps around the pocket defining residues contoured at 1.0 σ . 2—The central *Tg*AMA1 Tyr230 (gray) forms a functionally important bifurcated hydrogen bond with the backbone of *Tg*RON2 Val1311 (green). Plasticity may allow similar binding between *Nc*AMA1 Ser224 (cyan) and the corresponding *Nc*RON2 Ala1310. (C) Left—End-on view of *Pf*RON2sp1 (orange secondary structure) bound to *Pf*AMA1 (gray surface) (PDB ID: 3ZWZ) shows a deep pocket on the surface of AMA1 into which a critical RON2 arginine residue anchors (orange sticks, dotted box). Right—An overlay of *Pf*RON2sp1 on *Bd*AMA1 (gray surface) shows that a similar pocket is not present (dotted box). (D) Electrostatic surface, end-on views of *Tg*AMA1 bound to *Tg*RON2sp (green secondary structure with cystine loop tip side chains shown as sticks) (left; PDB ID: 2Y8T), and *Nc*AMA1 (middle) and *Bd*AMA1 (right) overlaid on the *Tg*AMA1-*Tg*RON2sp co-structure, reveal key differences in the chemical nature of the cystine loop binding region. An interactive view is available in the electronic version of the article.

perhaps in maintaining the DII loop in its displaced conformation.

Intriguingly, only one residue differs between the AMA1-binding regions of *Tg*RON2 and *Nc*RON2

[Fig. 5(A)]. In the *T. gondii* interaction, RON2 Val1311 forms two backbone hydrogen bonds with the AMA1 apical groove central tyrosine (Tyr230). The structurally equivalent residue to *Tg*AMA1

Tyr230 in *NcAMA1* is Ser224, and the sequence-aligned residue of Val1311 is *NcRON2* Ala1310 [Fig. 5(A,B 2)]. It is possible that the *NcAMA1* serine aids in destabilizing the DII loop to facilitate displacement [Fig. 2(C left)], while the *NcRON2* alanine provides flexibility to maintain hydrogen bonding interactions important in complex formation. The ability of *NcAMA1* to bind *TgRON2sp* with similar binding affinity as that observed for *TgAMA1* reflects an intriguing level of structural plasticity [Fig. 4(C)]. The similar binding affinity fits well with the observations of *NcAMA1* DII loop instability [Fig. 2(C right)] combined with the predicted absence of backbone hydrogen bonds between *NcAMA1* Ser224 and *TgRON2sp* Val1311 [Fig. 5(B 2)], and is also consistent with the order of magnitude lower binding affinity of the *TgAMA1* Y230A mutant for *TgRON2sp*.²⁵ Collectively, these data indicate that interaction with the central tyrosine, and by extension the central serine, is important but not essential.²⁵ Aside from the Tyr–Val interaction, every residue on *TgAMA1* shown to play a critical role in binding *TgRON2sp* is conserved in *NcAMA1*. Spatially, Phe157 (*TgAMA1* Phe163) likely needs to rotate in order to accommodate RON2, while loops 1 and 2 guarding the cystine loop binding surface need to be displaced by at least 3.0 Å to enable access to this region. A similar loop reorganization is observed between the apo and receptor bound forms of *TgAMA1*.

No binding could be detected between *BdAMA1* and 5FAM-*TgRON2sp* [Fig. 4(C)]. This is not surprising based on the observation that neither *PfAMA1* nor *PvAMA1* are capable of binding *TgRON2* in SPR assays,²⁵ and is likely due to a different anchoring mechanism in the cystine loop binding region.

Selectivity in the AMA1-RON2 interaction

Based on a comparison of the *TgAMA1-TgRON2sp* and *PfAMA1-PfRON2sp* coordination events, it is clear that selectivity is governed by the cystine loop region.^{25,27} Analysis of the *TgAMA1-TgRON2sp* co-structure did not reveal any particular hot spots of interaction, aside from the disulfide bond structure.²⁵ However, a similar analysis of the co-structure from *P. falciparum* showed that a single arginine residue in the cystine loop is absolutely critical for complex formation²⁷; very little of the *PfRON2* cystine loop contacts the *PfAMA1* surface, but Arg2041 buries into a deep pocket [Fig. 5(C left)], resulting in numerous polar contacts.²⁷ The *PfAMA1* surface pocket that accommodates this Arg residue is also exploited by the invasion inhibitory, species-specific phage display peptide, R1.²⁷ From the alignment of RON2 sequences [Fig. 5(A)], and given that there are no well-defined pockets on the *BdAMA1* surface within the cystine loop binding region

[Fig. 5(C right)], *BdRON2* is unlikely to have a singular anchor as was observed for the *PfAMA1-PfRON2sp* interaction. Therefore, although *BdAMA1* is overall more similar to *PfAMA1*, the *BdAMA1-BdRON2* interaction is more likely to resemble *TgAMA1-TgRON2sp*, with numerous contacts anchoring the cystine loop.

The surface of *TgAMA1* engaged by the *TgRON2sp* cystine loop is largely hydrophobic, with a partially hydrophilic border [Fig. 5(D left)]. This surface is compatible with the Glu-Ile-Leu tip of the cystine loop, in which the Ile and Leu brace the interaction.²⁵ The *NcRON2* cystine loop region is conserved with *TgRON2sp*, yet the corresponding AMA1 region is notably more basic [Fig. 5(D middle)]. The increased alkalinity is due to an Asn-Lys substitution (*TgAMA1* Asn184–*NcAMA1* Lys178), which may result in a stabilizing salt bridge with *NcRON2* Glu1316. Not surprisingly, the *BdAMA1* surface likely capable of accommodating the *BdRON2* cystine loop has a hydrophobic core, but with an acidic border that would coordinate the Ile-Lys-Pro-Pro loop tip [Fig. 5(A, D right)]. It is noteworthy that the electrostatic charges across the remainder of the groove are well conserved despite low sequence identity, with a central hydrophobic region, and a basic patch revealed by displacement of the DII loop. Together, these observations support the previously proposed model of specificity, wherein the RON2 cystine loop governs cross-species selectivity within an otherwise highly conserved binding paradigm.^{25,27}

Implications for generating cross genera inhibitors of AMA1-RON2 binding

In addition to better defining the basis for both plasticity and selectivity in the AMA1-RON2 interaction, the five AMA1 structures, two AMA1-RON2 co-structures, and AMA1-R1 co-structure serve as a robust model for the design of broadly reactive inhibitors. The first step in this process is the identification of well-conserved pockets that can be exploited by small molecules. Overlaying all three structurally characterized ligands—*TgRON2sp*, *PfRON2sp*, and R1—on the five AMA1 structures—*TgAMA1*, *NcAMA1*, *BdAMA1*, *PfAMA1*, and *PvAMA1*—[Fig. 6(A left), *TgAMA1* alone shown for clarity] reveals three defined pockets on the surface of AMA1 [Fig. 6(A inset)]. Pocket 1 is very well conserved (Table I), and has been shown to be occupied by the tip of the *NcAMA1* and *TgAMA1* DII loops (Trp347 and Trp353), a conserved RON2 proline (*TgRON2* Pro1309; *PfRON2* Pro2033), and R1 Leu6.^{25,27,46} Clearly, occupation of Pocket 1 is a strategic target, but to be accessed by a small molecule the DII loop would need to be displaced. To overcome the energy barrier associated with DII loop rearrangement, additional anchor points within the

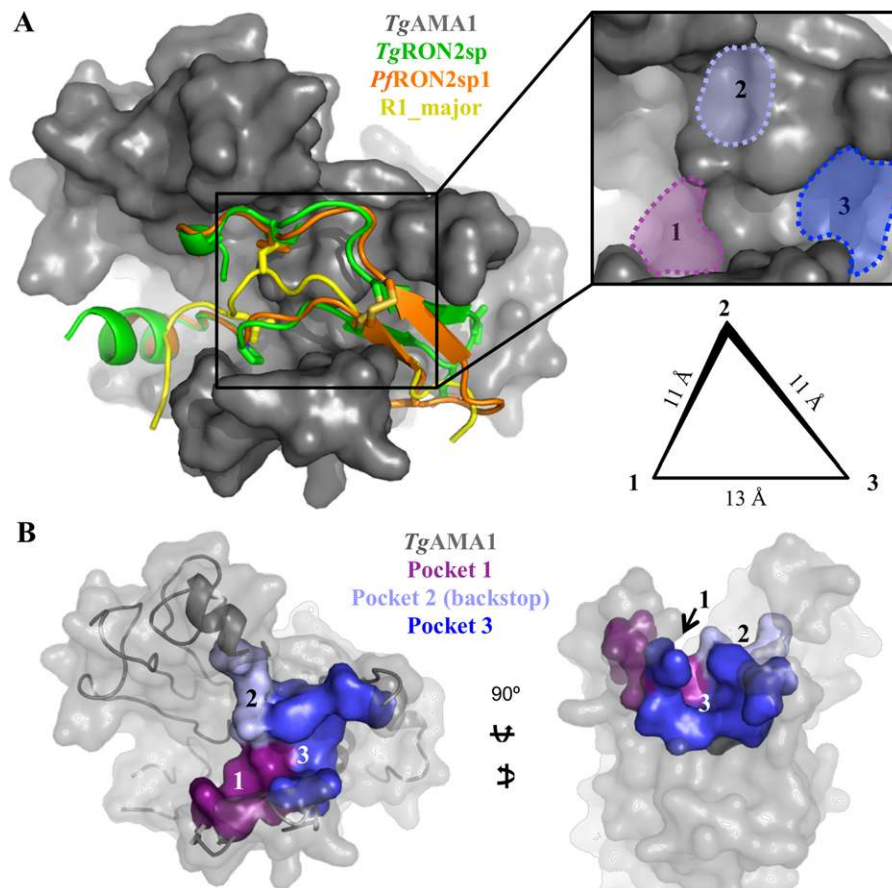


Figure 6. Identification of surface pockets ideal for targeting with therapeutics. (A) Left—apical view of an overlay of three structurally characterized ligands (shown as secondary structure, with residues occupying pockets of interest shown as sticks)—*TgRON2sp* (green; PDB ID: 2Y8T), *PfRON2sp1* (orange; PDB ID: 3ZWZ), *R1_major* (yellow; PDB ID: 3SRJ)—on the surface of *TgAMA1* (gray). Inset—enlarged apical view for the identification of three surface pockets with structural and chemical features conserved across all the structurally characterized AMA1 proteins. Right—schematic representation showing the relative positions of the three pockets in a triangular organization, with Pocket 2 in a higher plane than Pockets 1 and 3. Pocket 2 is equidistant from 1 and 3 (11 Å), while Pockets 1 and 3 are separated by roughly 13 Å. (B) Left—apical surface of *TgAMA1* (gray) in the same orientation as (A), with the residues defining Pocket 1 shown purple; 2, light blue; 3, bright blue. Right—end-on view highlighting the depth of Pocket 3 and the backstop nature of Pocket 2.

AMA1 groove would be essential. *TgRON2* Ile1328, *PfRON2* Val2054, and *R1* Leu8 all brace against the base of loop 4 in a shallow hydrophobic pocket, which constitutes Pocket 2 (backstop; Fig. 6, Table I). Binding to this pocket alone would not provide enough traction to displace the DII loop, but extending too far toward the cystine loop binding region would severely limit cross reactivity. However, the disulfide bond plays a critical and highly conserved role in complex formation,^{25,27} and thus the pocket that binds this structural feature would be an excellent anchor point, and is presented here as Pocket 3 (Fig. 6 and Table I). These three pockets have a highly conserved chemical and structural architecture (Fig. 6 and Table I), are in close proximity, and maintain a triangular geometry within the apical groove of AMA1 [Fig. 6(A right)] that could be efficiently occupied by a broadly reactive small molecule inhibitor.

Materials and Methods

Bioinformatics

Boundaries for DI, DII, and DIII were defined based on the paradigm established for previously characterized AMA1s.^{43,45,46,56} Phylogenetic analysis was performed using MEGA 4^{62,63} and multiple sequences were aligned using MUSCLE⁶⁴ and illustrated in ESPript.⁶⁵ Accession numbers for aligned AMA1 sequences are as follows: *T. gondii* (UniProt ID: B6KAM0), *N. caninum* (UniProt ID: A2A114), *B. divergens* (UniProt ID: C0IR59), *P. falciparum* (UniProt ID: Q7KQK5), and *P. vivax* (UniProt ID: A5K4Z2; sequence modified to reflect crystallized sequence,⁴⁵ with 3 N-linked glycosylation sites mutated—PDB ID: 1W8K).

RON2 sequences in the region of *Tg/PfRON2sp* were aligned using ClustalW,⁶⁶ and manually edited based on structural evidence. Accession numbers

Table I. Residues Lining the Conserved Apical Surface Pockets of AMA1

<i>Tg</i> AMA1	<i>Nc</i> AMA1	<i>Bd</i> AMA1	<i>Pf</i> AMA1	<i>Pv</i> AMA1
Pocket 1				
Val105	Val99	Val126	Val137	Val82
Tyr110	Tyr104	His131	Tyr142	Tyr87
Ala203	Ala197	Pro225	Pro226	ND ^a
Tyr213	Tyr207	Tyr233	Tyr234	Tyr179
Tyr215	Tyr209	Tyr235	Tyr236	His181
Pocket 2 (backstop)				
Val142	Ile136	Ile157	Val169	Val114
Pro143	Pro137	Ser158	Ala170	Ala115
Tyr230	Ser224	Phe250	Tyr251	Tyr196
Pocket 3				
Leu161	Met155	Phe169	Phe183	Phe128
Phe163	Phe157	Pro170	Thr186	Ala131
Thr165	Thr159	ND ^a	Phe188	His134
Ile171	Ile165	Leu195	Met190	Ile135
Thr201	Thr195	Ile223	Met224	Phe169
Met204	Met198	Ala226	Asp227	ND ^a
Tyr215	Tyr209	Tyr235	Tyr236	His181
Ile228	Ile222	Ile248	Ile249	Met194

Structurally equivalent residues are horizontally aligned.

^a Not determined (ND) due to flexible, unmodeled loop.

and sequence limits for aligned RON2 sequences are as follows: *T. gondii* (UniProt ID: F6KDI4, residues 1297-1333), *N. caninum* (UniProt ID: F0VQN9, residues 1296-1332), *B. divergens* (UniProt ID: G8Z7K1, residues 941-978), *P. falciparum* (UniProt ID: Q8IKV6, residues 2021-2059), and *P. vivax* (UniProt ID: A5K3N8, residues 2035-2073).

Protein expression and purification

Sequences encoding the fully processed ectoplasmic domains of *Bd*AMA1 and *Nc*AMA1 were synthesized by GenScript, codon optimized for insect cells, and subcloned into a modified pAcGP67b vector (Pharmingen) incorporating a C-terminal hexahistidine tag separated from *ama1* by a thrombin cleavage site. *Bd/Nc*AMA1 encoding viruses for insect cell protein production were generated and amplified using established protocols.^{25,46} Selected protein preparations had a final concentration of 0.3 µg/mL tunicamycin. Following a 65 h infection the supernatant was harvested, concentrated, buffer exchanged into Buffer A1 (20 mM HEPES pH 8.0, 30 mM imidazole, 1 M NaCl) and allowed to batch bind with Ni-agarose beads at 4°C for 1 h. AMA1 proteins were eluted with Buffer B1 (20 mM HEPES pH 8.0, 250 mM imidazole, 1 M NaCl), and fractions were analyzed by SDS-PAGE and pooled based on purity. The hexahistidine tag was removed by thrombin cleavage and AMA1 proteins were further purified by size exclusion chromatography (Superdex 16/60 200) in HEPES buffered saline (HBS), followed by anion exchange chromatography (Source 30Q). The final yield of both *Bd*AMA1 and *Nc*AMA1 was approximately 2 mg of purified protein per liter of insect cell culture.

Crystallization and data collection

Using the sitting drop method, crystals of *Bd*AMA1 were grown in 16% PEG8000, 300 mM calcium acetate, 100 mM sodium cacodylate pH 5.5 and 3% 2-methyl-2,4-pentanediol (MPD) and crystals of *Nc*AMA1 were grown in 20% PEG3350, 300 mM sodium citrate pH 4.2 and 10 mM zinc chloride. The final drops consisted of 1.5 µL protein (*Bd*AMA1 12 mg/mL; *Nc*AMA1 15 mg/mL) with 1.5 µL reservoir solution and were equilibrated against 100 µL of reservoir solution. Cryo protection of the *Bd/Nc*AMA1 crystals was performed in mother liquor supplemented with 12.5% glycerol for 20 s and the crystals were flash cooled at 100 K directly in the cryo stream. Diffraction data were collected on beamline 9-2 at Stanford Synchrotron Radiation Lightsource for *Bd*AMA1 and on beamline CMCF-ID at Canadian Light Source for *Nc*AMA1.

Data processing, structure solution and refinement

Diffraction data were processed using Imosflm⁶⁷ and Scala⁶⁸ in the CCP4 suite of programs.⁶⁹ Initial phases for *Bd*AMA1 were obtained by molecular replacement (MR) using PHASER⁷⁰ with the DI and DII domains of *Pf*AMA1 (PDB ID: 2Q8A) pruned with CHAINSAW.⁷¹ Despite multiple attempts, no MR solution was obtained for DIII, which was ultimately traced de novo into the electron density maps using COOT⁷² and shown to be divergent from other characterized DIIs. Initial phases for *Nc*AMA1 were obtained by MR using PHASER⁷⁰ with chain B of *Tg*AMA1 (PDB ID: 2X2Z) pruned with CHAINSAW,⁷¹ and improved with four-fold non-crystallographic symmetry averaging. Solvent molecules were selected using COOT⁷² and refinement performed using Refmac5.⁷³ Stereochemical analysis performed with PROCHECK and SFCHECK in CCP4⁶⁹ showed good stereochemistry with more than 92% of the residues in the favored conformations and no residues modeled in disallowed orientations of the Ramachandran plot. Overall 5% of the reflections were set aside for calculation of R_{free} . Data collection and refinement statistics are presented in Table II.

Native gel electrophoresis assay

Purified AMA1 protein in HBS buffer pH 8.0 was incubated for 30 min at room temperature with or without *Tg*RON2sp²⁵ in a 1:2 protein to peptide molar ratio. Reactions were run on 8–25% gradient gels using native buffer strips and the PhastGel system (GE Healthcare).

Fluorescence polarization assay

In black 96-well assay plates (Nunc), AMA1 proteins in HBS were serially diluted from 1 µM to 0.042 nM and incubated with > 90% pure 10 nM *Tg*RON2sp

Table II. Data Collection and Refinement Statistics

	<i>BdAMA1</i>	<i>NcAMA1</i>
A. Data collection		
Spacegroup	P2 ₁ 2 ₁ 2	C2
a, b, c (Å)	67.82, 139.18, 45.01	251.56, 51.00, 145.34
α, β, γ (°)	90, 90, 90	90, 90.93, 90
Wavelength (Å)	0.9791	0.9795
Resolution range (Å)	42.83–2.30 (2.42–2.30)	50.00–2.90 (3.06–2.90)
Measured reflections	90,064	161,770
Unique reflections	19,533	41,269
Redundancy	4.6 (4.5)	3.9 (3.8)
Completeness (%)	99.2 (96.4)	99.1 (96.6)
<i>I</i> / σ (<i>I</i>)	12.5 (3.7)	9.8 (4.4)
<i>R</i> _{merge} ^a	0.070 (0.389)	0.112 (0.385)
B. Refinement		
Resolution (Å)	42.83–2.30 (2.36–2.30)	49.98–2.90 (2.98–2.90)
<i>R</i> _{cryst} ^b / <i>R</i> _{free} ^c	0.190/0.251 (0.228/0.284)	0.193/0.271 (0.273/0.396)
No. of atoms		
Protein (A/B/D/E)	2624	3155/3123/3132/3094
Solvent	166	142
Calcium	1	N/A
Acetate/Glycerol/MPD	12/36/8	N/A
<i>B</i> -values (Å ²)		
Protein (A/B/D/E)	38.9	36.3/39.0/34.7/45.1
Solvent	45.7	31.0
Calcium	30.2	N/A
Acetate/Glycerol/MPD	62.3/65.4/66.7	N/A
r.m.s. deviation from ideality		
Bond lengths (Å)	0.01	0.01
Bond angles (°)	1.33	1.05
Ramachandran statistics		
Most favoured	97.0%	92.8%
Allowed	3.0%	7.2%
Disallowed	0.0%	0.0%

Values in parentheses are for the highest resolution shell.

^a $R_{\text{merge}} = \sum_{hkl} \sum_i |I_{hkl,i} - [I_{hkl}]| / \sum_{hkl} \sum_i I_{hkl,i}$, where $[I_{hkl}]$ is the average of symmetry related observations of a unique reflection.

^b $R_{\text{cryst}} = \sum |F_{\text{obs}} - F_{\text{calc}}| / \sum F_{\text{obs}}$, where F_{obs} and F_{calc} are the observed and the calculated structure factors, respectively.

^c R_{free} is R using 5% of reflections randomly chosen and omitted from refinement.

labeled at the N-terminus with 5FAM (Kinexus; Vancouver, Canada). The plate was mixed on a shaker for 15 min and incubated at room temperature for 30 min to reach equilibrium. Fluorescence polarization in millipolarization units (mP) was read using a Spectra-Max 5M plate reader (Molecular Devices) with an excitation wavelength of 485 nm and emission wavelength of 538 nm. For determination of EC₅₀ values, data were fit with a sigmoidal dose dependent nonlinear regression model (variable slope) in GraphPad Prism 5.0. Errors were calculated as the standard error of the mean (SEM) from triplicate samples.

Accession numbers

The atomic coordinates and structure factors have been deposited in the Protein Data Bank under the following codes: *BdAMA1*—PDB ID: 4APM, r4APMsf; *NcAMA1*—PDB ID: 4APL, r4APLsf.

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