Saliva of blood-feeding arthropods carries several antihemostatic compounds whose physiological role is to facilitate successful acquisition of blood. The identification of novel natural anticoagulants and the understanding of their mechanism of action may offer opportunities for designing new antithrombotic disrupting blood clotting. We report here an in-depth structural and functional analysis of the anophelin family member cE5, a salivary protein from the major African malaria vector Anopheles gambiae that specifically, tightly, and quickly binds and inhibits thrombin. Using calorimetry, functional assays, and complementary structural techniques, we show that the central region of the protein, encompassing amino acids Asp-31–Arg-62, is the region mainly responsible for α-thrombin binding and inhibition. As previously reported for the Anopheles albimanus orthologue anophelin, cE5 binds both thrombin exosite I with segment Glu-35–Asp-47 and the catalytic site with the region Pro-49–Arg-56, which includes the highly conserved DPRG tetraepitope. Moreover, the N-terminal Ala-1–Ser-30 region of cE5 (which includes an RGD tripeptide) and the additional C-terminal serine-rich Asn-63–Glu-82 region (absent in orthologues from anopheles of the New World species A. albimanus and Anopheles darlingi) also played some functionally relevant role. Indeed, we observed decreased thrombin binding and inhibitory properties even when using the central cE5 fragment (Asp-31–Arg-62) alone. In summary, these results shed additional light on the mechanism of thrombin binding and inhibition by this family of salivary anticoagulants from anopheline mosquitoes.

The ability of hematophagous insects to use blood as a food source involves complex behavioral, morphological, and physiological adaptations to find suitable hosts, pierce their skin, and efficiently acquire and digest blood. As a result, the saliva of blood-feeding arthropods carries a complex mixture of compounds playing crucial roles in counteracting the three powerful and highly redundant responses of vertebrates to tissue injury: hemostasis, inflammation, and immunity (1). Blood feeding evolved independently several times in different taxa (2), and this resulted in a large variety of salivary anti-hemostatic factors targeting platelet aggregation, vasoconstriction, and blood clotting (1, 3). Salivary anticoagulants found in the mosquito family offer a good example of this convergent evolution; members of the Culiciniae subfamily (i.e. Aedes and Culex species) inhibit factor Xa, whereas species belonging to the anopheline subfamily adopted a thrombin-directed anticoagulant activity (4).

The anti-thrombin of anopheline mosquitoes was first identified in the South American malaria vector Anopheles albimanus as a small cysteine-less polypeptide of 61 amino acids named anophelin (5). Synthetic anophelin was shown to be a highly specific, slow, tight-binding, reversible inhibitor of α-thrombin (6). It binds both the catalytic site and the anion binding exosite 1 (TABE1) of thrombin, a region that is known to be involved in the recognition of fibrinogen (7). A cDNA encoding the anophelin orthologue in the African malaria vector Anopheles gambiae was previously identified by a selective cloning strategy and the encoded protein named cE5 (8). The two proteins, anophelin and cE5, are quite divergent (43% identity) with the slightly longer A. gambiae protein (82
amino acids) displaying an additional stretch of amino acids at the C terminus. Despite these differences, recombinant A. gambiae cE5 showed full preservation of anti-thrombin function, although the two proteins displayed slightly different binding kinetics. The A. albimanus anophelin is a slow-binding thrombin inhibitor (6), whereas the A. gambiae cE5 behaves as a fast-binding inhibitor (9), raising the possibility that the extended C-terminal may participate in thrombin binding.

The crystal structure of the A. albimanus anophelin in complex with human thrombin revealed its unique inhibition mechanism in comparison to other known thrombin inhibitors (10). In particular, the C-terminal segment (A(Asp-33–Phe-45) blocking the exosite I and the highly conserved DPGR tetrapeptide (A(Asp-50–Arg-53)) occupying the active site cleft of the enzyme and disrupting the characteristic catalytic triad of the serine proteinase.

The release of the genomes of 16 Anopheles mosquitoes (11) allowed identifying anophelin/cE5 orthologues from several additional species, offering further insights into the evolution of this unique family of thrombin inhibitors. Multiple alignment of cE5/anophelin family members showed that, in comparison to the New World species A. albimanus and Anopheles darlingi, orthologues from the Old World Anopheles species display an extended, serine-rich, C-terminal region (10, 12). In order to investigate the possible involvement of the C-terminal region (cE5(Asn-63–Glu-82)) in thrombin-binding and inhibition as well as the role of the conserved N-terminal portion (cE5(Ala-1–Ser-30)), we performed a structural and functional analysis using both the cE5 protein and a set of cE5-derived peptides. Moreover, the three-dimensional structure of the human α-thrombin–A. gambiae cE5 complex unveiled the details of thrombin recognition and inhibition by an anophelin orthologue from an Old World Anopheles species.

Results

Structural properties of cE5 as determined by circular dichroism and NMR spectroscopy

Prediction analysis carried out by DISOPRED (VLXT predictor; www.pondr.com)3 (50) suggested that the cE5 protein, as previously reported for the A. albimanus anophelin (10), was intrinsically disordered in solution. However, in contrast to A. albimanus anophelin, cE5 displayed a slight propensity to be structured in the region encompassing amino acids cE5(Pro-54) to cE5(Ser-64) (Fig. 1). Similar results were obtained using the database of intrinsically disordered proteins, MobiDB (not shown) (13). Furthermore, cE5 displayed a typical circular dichroism (CD) spectrum of an unstructured protein, with a minimum at 202 nm (Fig. 2). After the addition of 2,2,2-trifluoroethanol (TFE)4 as co-solvent, which may help exploring the intrinsic tendency of a polypeptide to assume secondary structure elements (14–16), we found a very low propensity of cE5 to adopt an α-helical conformation. Indeed, even with high concentrations of TFE (60%), an increase of only ~10% in the α-helical content was observed (inset in Fig. 2).

The conformational properties of cE5 were also investigated using NMR spectroscopy. The 1H,15N HSQC spectrum of 15N-labeled cE5 showed a limited chemical shift range (Fig. 3A), indicative of an unfolded molecule having few secondary structure elements, consistent with the random-coil state suggested by CD data. In addition, the 1H,15N HSQC spectra of uniformly 15N-labeled cE5 (40 μM) in the absence and the presence of different amounts of TFE (10, 30, and 60% (v/v)) (Fig. 3, B–D) were recorded in the same experimental conditions used for CD analyses. In complete agreement with the CD data, these NMR experiments indicated a continuous increase in signal dispersion (i.e. in structuration) with increasing TFE concentration. However, even at relatively high TFE concentrations, HSQC spectra still resemble those of canonical intrinsically disordered proteins (Fig. 3, B–D).

Alignment of anophelin family members from different Anopheles species

The sequencing of the genomes of 16 anopheline species (11) spanning ~100 million years of evolution allowed identification of several additional members of the cE5/anophelin family (12). Alignment of the 18 full-length orthologues highlighted their considerable divergence: 18–77% amino acid identity among the different Anopheles species (excluding comparisons within the A. gambiae species complex). Overall, a block of 16 amino acids, including the highly conserved tetrapeptide APQY, can be recognized at the N-terminal region, and another highly conserved tetrapeptide, DPGR, is found toward the C terminus (Fig. 4). Between these two conserved blocks there is a more divergent central region enriched in acidic residues (Asp+Glu = 19–33%). A careful examination of the aligned proteins disclosed three features of potential functional relevance: (i) the N-terminal RGD tripeptide in A. gambiae and in a few other species of the complex; (ii) the conserved DPGR tetrapeptide, with proline to alanine replacement in Anopheles atroparvus and Anopheles epiroticus; (iii) the presence of a longer C-terminal in Old World anophelines.

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4 The abbreviations used are: TFE, 2,2,2-trifluoroethanol; HSQC, heteronuclear single quantum correlation; ITC, isothermal titration calorimetry; PCTP, sodium propionate, sodium cacodylate trihydrate, bis-Tris propane; bis-Tris, 2-(bis[2-hydroxyethyl]amino)-2-(hydroxymethyl)propane-1,3-diol.
Figure 2. Far UV CD spectra of the cE5 protein. CD spectra of cE5 (10 μM in 10 mM phosphate buffer pH 7.5) alone (solid line) and after the addition of 60% (v/v) TFE (dotted line). The inset shows the molar ellipticity at 220 nm in the presence of increasing concentrations of TFE.

Figure 3. NMR spectroscopy of the A. gambiae cE5 protein. A, 1H,15N HSQC spectrum of uniformly 15N-labeled cE5 (20 μM in 50 mM sodium phosphate, pH 7.8; sample total volume: 550 μL with 10% D2O). B–D, monitoring the effect of TFE on the intrinsically disordered protein cE5. Comparison of HSQC spectra of 15N-labeled cE5 in the absence (red) and in the presence of TFE at increasing concentrations: B, 10% (green); C, 30% (cyan); D, 60% (blue). NaP stands for sodium phosphate buffer.
The tripeptide RGD is known for its involvement in binding to integrins (17). RGD-containing proteins with anti-platelet activity (disintegrins) have been found in snake venoms and in the salivary gland of blood-feeding arthropods (ticks and the saliva of blood-feeding arthropods (ticks and horseflies), leeches, and worms but never in mosquitoes. Disintegrins act as antagonists of integrin αIIbβ3 and inhibit fibrinogen binding to platelets and subsequent platelet cross-linking (18). Typically, integrin-binding RGD motifs are positioned in the loop of peptide hairpins formed by disulfide bonds (19), which is not the case for the cE5 RGD. We tested cE5 and the cE5-derived P1 peptide (Fig. 4) in platelet aggregation assays. We found no or very little effect on platelet aggregation (supplemental Fig. S1), which seems to rule out interaction of cE5 RGD with integrin αIIbβ3 (even though we cannot exclude binding to other integrins).

The DPGR tetrapeptide was previously found to play a crucial role in in vitro binding to α-thrombin (20) and shown to occupy the active-site cleft of the enzyme in the anophelin/thrombin crystal structure (10). Substitutions of the catalytic triad-disrupting aspartate and of the S1-targeting arginine residues in the A. albimanus protein were detrimental for the inhibitory activity (10). We do not know the effect of replacing the proline in this tetrapeptide with an alanine. However, it is conceivable that given the compact size of alanine, it may replace the pivotal proline residue without much impact in inhibitor activity.

Intriguingly, Old World anophelines (subgenera Cellia and Anopheles) carry an additional serine-enriched (29–56%) C-terminal stretch of 7–21 amino acids that is absent in the New World species A. albimanus and A. darlingi (subgenus Nyssorhynchus). It has been previously shown that the anophelin C-terminal fragment A(Glu–32–Pro–61) is mainly responsible for both binding to and inhibiting thrombin. Fragment A(Ala–1–Asp–31) did not appear to play any functional role in thrombin targeting and inhibition (10). However, A. gambiae cE5 appeared to possess higher affinity for thrombin (Table 1 and Fig. 5, A and B). In addition, binding assays with the cE5-based peptides P1, P2, P3, and P5 suggested that the region involved in the interaction with thrombin resides almost completely on the fragment cE5(Ala–1–Ser–30), P2 (cE5(Asp–31–Arg–62)), P3 (cE5(Asp–63–Glu–82)), and P5 (cE5(Asp–31–Glu–82)) (Fig. 4) and assessed their interaction with thrombin by isothermal titration calorimetry (ITC) and enzymatic assays.

**Binding studies**

As determined by ITC, cE5 formed a very stable complex with thrombin ($K_a = 1\ \text{nm}$), with an apparent affinity slightly higher than anophelin ($K_d = 2\ \text{nm}$; Table 1, Fig. 5, A and B). In addition, binding assays with the cE5-based peptides P1, P2, P3, and P5 suggested that the region involved in the interaction with thrombin resides almost completely on the fragment cE5(Asp–31–Arg–62), corresponding to the P2 peptide ($K_d = 12\ \text{nm}$). This was underscored by the comparable ΔH values obtained for cE5 ($−9.85\ \text{kcal/mol}$) and P2 ($−9.66\ \text{kcal/mol}$) (Table 1 and Fig. 5, C–E). Surprisingly, binding experiments with a P5 yielded larger $K_d = (25\ \text{nm})$ and smaller ΔH ($−4.89\ \text{kcal/mol}$) values than those obtained for P2, pointing to a negative contribution in thrombin-binding of the region corresponding to P3 in the absence of the N-terminal portion encompassed by P1 (Table 1, Fig. 5F).

It is worth noting that the binding affinity ($K_a$) of cE5 to thrombin is beyond the limit of direct calorimetric determination. Under these conditions, ITC titration still provides an accurate measurement of the binding enthalpy (Table 1). However, ITC can be used to determine the complete binding thermodynamics of ligands with affinities down to the picomolar range using displacement titration (21), which was performed for this system in the presence of a weak thrombin inhibitor, the synthetic tetrapeptide d-Phe–Pro–d-Arg–Ala (22). Under these conditions, cE5 displayed a binding constant of 0.3 nM (Table 1, supplemental Fig. S2, and supplemental Equation 1),
A. gambiae cE5–thrombin interaction

Table 1

| Ligand        | $K_d$  | $\Delta H$  | n  |
|---------------|--------|-------------|----|
| cE5          | 0.3 nm | −8,272      | 1  |
| cE5*         | 0.3 nm | −8,272      | ND |
| P1 (cE5(1–30)) | ND   | ND          | ND |
| P2 (cE5(31–62)) | ≤12 ± 1 nm | −9,663 ± 34 | 1  |
| P3 (cE5(63–82)) | ND   | ND          | ND |
| P5 (cE5(31–82)) | ≤25 ± 1 nm | −4,887 ± 75 | 1  |
| Anophelin    | ≤2 ± 0.2 nm | −14,120 ± 90 | 1  |

* From displacement titration experiments.

in line with the value previously reported (9, 10) and with that obtained from the thrombin amidolytic activity inhibition assays in this study (Table 2).

We also used NMR titration experiments and analysis of 2D $^1$H,$^15$N HSQC spectra (23) to study protein-protein interactions. HSQC spectra of $^{15}$N-labeled cE5 were recorded in the absence and presence of increasing amounts of unlabeled thrombin. At cE5/thrombin ratios >1/5, numerous spectral changes could be observed, pointing to interactions taking place between the two proteins (supplemental Fig. S3). However, it was not possible to reach saturating conditions, which would have allowed gaining more detailed structural information. This is most likely due to the relatively large size of the complex, which caused severe line broadening and loss of sensitivity in the NMR spectra (supplemental Fig. S3). In addition, 2D $^1$H, $^1$H NMR spectra collected for the P2 peptide showed a similar lack of relevant chemical shift dispersion and the absence of a consistent number of nuclear Overhauser effect (NOE) contacts, confirming a disordered state for the P2 peptide in aqueous solution (supplemental Fig. S4, right panel). Moreover, no significant change in either chemical shift and/or NOE intensity could be observed when comparing 2D $^1$H, $^1$H NOESY (Nuclear Overhauser Enhancement Spectroscopy) experiments with the P2 peptide (470 $\mu$m) in the absence and the presence of a substoichiometric amount of thrombin (25 $\mu$m) (supplemental Fig. S4). These observations suggest that under our experimental conditions, the peptide “dynamically” binds to thrombin by preserving disorder and flexibility.

Thrombin inhibition assays

The inhibitory effect of cE5 and of peptides P1, P2, P3, and P5 on the in vitro amidolytic activity of thrombin was measured after the hydrolysis of a fluorogenic substrate. A thrombin inhibition of ~80% was found for cE5, in line with previous measurements using a chromogenic substrate (9). The P2 peptide still retained inhibitory properties even though it was significantly less effective than cE5 (~39% inhibition), whereas neither P1 nor P3 affected thrombin activity (Fig. 6). The inhibitory effect of P5 was not significantly different from those observed using P2 alone or any combination including either P2 or P5.

Overall, thrombin inhibition results fit quite well with ITC data, confirming that the region corresponding to the P2 peptide (cE5(Asp-31−Arg-62)) is the main segment responsible for both binding to thrombin and for the inhibitory activity. The P1 peptide (cE5(Ala-1−Ser-30)) did not bind thrombin and did not affect its activity, and these results are fully consistent with previous observations on A. albimanus anophelin (10). Moreover, our results seem to rule out any direct functional role of the serine-rich C-terminal region in thrombin inhibition. In fact, the P3 peptide alone did not bind to or inhibit thrombin, and the P5 peptide, which encompasses both P2 and P3, was not a better inhibitor or a more efficient binder than P2.

Overall structure of the A. gambiae cE5–human α-thrombin complex

The recombinant P5 fragment (cE5(Asp-310−Glu-82)) of A. gambiae cE5 was crystallized in complex with human α-thrombin, and its structure was determined at 1.45 Å resolution (supplemental Table 1 and Fig. 7). There are three copies (A, B, and C) of the complex in the asymmetric unit: complex A comprising thrombin residues T(Ser-1E) to T(Arg-15) and T(Ile-16) to T(Gly-246) and inhibitor residues cE5(Asn-63) (the chymotrypsinogen-based numbering will be used for thrombin); complex B encompasses thrombin residues T(Thr-1H) to T(Arg-15) and T(Ile-16) to T(Gly-246) and inhibitor residues cE5(Glu-36) to cE5(Leu-42) and cE5(Ala-46) to cE5(Arg-62); finally, in complex C thrombin residues T(Gly-1D) to T(Ile-14K), T(Ile-16) to T(Gln-244), and inhibitor residues cE5(Arg-52) to cE5(Leu-61) could be modeled. In all complexes, thrombin’s 149 insertion loop was disordered (from T(Thr-148) to T(Lys-149E), T(Thr-148) to T(Gly-149D), and T(Thr-147) to T(Gly-149D), for complexes A to C, respectively) and could not be modeled. Although the three complexes are structurally very similar (root mean square deviations of 0.44 Å, 0.82 Å, and 0.93 Å for the superposition of the protease moieties of complexes A-B, A-C, and B-C, respectively), complex C displays an unusual flexibility of segments of the heavy chain of thrombin, encompassing residues T(Gln-38)–T(Lys-70), T(Arg-101)–T(Glu-127), T(Met-210)–T(Cys-220), and T(Phe-227)–T(Gln-244). Therefore, only complex A will be described hereafter (Fig. 7). No significant density was present for the N- and C-terminal tetrapeptide cE5(Asp-31) to cE5(Glu-34) or for the C-terminal 21 amino acids (cE5(Ser-64) to cE5(Glu-82)) of the P5 fragment, and these segments were, therefore, excluded from the model. Superposition of the structures of free thrombin (PDB entry 3U69; Ref. 22) and the protease moiety of the A complex revealed that only minor rearrangements occur in the protease upon A. gambiae cE5 binding (root mean square deviation of 0.36 Å for 277 aligned Ca atoms).

Unsurprisingly, A. gambiae cE5 displays the same unique mode of thrombin inhibition observed for A. albimanus anophelin (10), further confirming it as a general mechanism for this class of inhibitors (MEROPS family I77; Ref. 24; Fig. 7B). In agreement with the results of CD (Fig. 2) and NMR measurements (Fig. 3), the inhibitor adopts a mostly extended conformation, running in a reverse orientation to substrates on the surface of thrombin and interacting with both the exosite I and the active-site region of the proteinate (Fig. 7, A, C, and D). Further supporting the observed binding mode, both full-length A. gambiae cE5 and its truncated P5 fragment display 3 orders of magnitude larger inhibition constants toward the exosite I-disrupted γ-thrombin than toward α-thrombin (Table 2).
Bidentate binding of cE5 to thrombin

Similar to the A. albimanus anophelin-thrombin complex (10), the cE5 fragment spanning residues cE5(Glu-35) to cE5(Asp-47) blocks the exosite I of the proteinase (Figs. 4 and 7C). However, despite the overall resemblance, the first half of this cE5 segment runs closer to the proteinase surface, with its main-chain atoms deviating significantly from the path of the equivalent region of A. albimanus anophelin (Fig. 7B). The inhibitor’s tripeptide cE5(Glu-35)–cE5(Glu-36)–cE5(Phe-37)

Table 2

| Inhibitor       | α-Thrombin | γ-Thrombin |
|-----------------|------------|------------|
| cE5             | 5.50 ± 1.26| 3.79 ± 0.16|
| P5 fragment     | 20.80 ± 0.99| 10.99 ± 0.69|

Figure 5. ITC analysis of thrombin binding to A. gambiae cE5, A. albimanus anophelin, and cE5-derived peptides. Isothermal titration calorimetry of thrombin interaction with A. gambiae cE5 (A), A. albimanus anophelin (B), P1 (C), P2 (D), P3 (E), and P5 (F) is shown. The top and bottom panels report raw and integrated data, respectively.
establishes water-mediated main chain–main chain contacts with thrombin’s T(Ile-82)–T(Ser-83)–T(Met-84) tetrapeptide. This region is also stabilized by an intramolecular hydrogen bond between the carbonyl oxygen of cE5(Asp-38) and the amide nitrogen of cE5(Leu-41), further reinforced by two interactions with the side chain of T(Gln-38), involving the main chain nitrogen of cE5(Glu-43) and the carbonyl oxygen of cE5(Leu-41). This conformation results in the placement of the side chain of cE5(Phe-37) snugly inside the hydrophobic depression bordered by the side chains of T(Leu-65), T(Ile-82), and T(Met-84), to which it establishes van der Waals interactions. Additionally, the aromatic ring of cE5(Phe-37) also contacts the side chain of the well-conserved cE5(Leu-42), in the same position of the homologous A(Leu-41) of A. albimanus anophelin, and in turn contacting T(Phe-34) and T(Tyr-76). The side chains of cE5(Pro-39) and cE5(Leu-41) pack on top of that cE5(Phe-37), completing this strong hydrophobic cluster.

Furthermore, toward the active site cleft of thrombin, cE5(Asp-47) forms a salt bridge with T(Arg-73). Notably, the replacement of A. albimanus anophelin A(Asp-45) by the much shorter cE5(Asp-48) impairs the highly conserved stacking interaction with T(Phe-34) observed in numerous thrombin ligands, from protease-activated receptor 1 (PAR1) (25) to hirudin (26) and from boophilin (27) to anophelin (10). Instead, two polar interactions are observed between the carbonyl oxygen of cE5(Ala-48) and the side chains of T(Leu-65) and T(Asp-151). From cE5(Pro-49) all the way to cE5(Arg-56), the interactions established between thrombin and the homologous A(Thr-47–Arg-53) segment of A. albimanus anophelin are strictly conserved (10), in agreement with the remarkable amino acid sequence conservation observed in all members of the family (Fig. 4). Therefore, cE5(Arg-53) interacts with the catalytic T(His-57) and T(Ser-195), whereas cE5(Arg-56) occupies the S1 specificity pocket of the protease (Fig. 7D). The downstream cE5(Asn-57) interacts with the side chain of T(Thr-192), resembling the contact established by A. albimanus anophelin A(Arg-54) (10), but the main chain of the remaining ordered cE5(Pro-58)–cE5(Leu-61) tetrapeptide deviates considerably from the homologous segment of anophelin (Fig. 7B). Despite that, the position of the side chain of cE5(Phe-60) closely resembles that of A. albimanus anophelin A(Leu-55), therefore occupying the aryl binding site of thrombin.

Discussion

Overall, the cE5 central region represented by the P2 peptide appeared as the main responsible for both thrombin binding and inhibition. The fact that only the cE5(Glu-35–Asn-63) segment of peptide P5 could be modeled in the crystal structure and no significant electron density was detectable for the serine-rich C-terminal portion reinforces this main role of the region corresponding to P2. Moreover, the C-terminal region was not constrained by crystal packing, explaining its observed disorder (supplemental Fig. S5). The binding of cE5 to thrombin was to a large extent similar to that previously described for A. albimanus anophelin. Nonetheless, the cE5 N-terminal and C-terminal regions still appear to play some role in both binding and inhibition. Indeed, when these two disordered segments were missing, we observed a decrease in binding affinity (≤1 nM for cE5 versus ≤12 nM for P2) as well as in thrombin inhibition (80% for cE5 versus 39% for P2). These more efficient binding and inhibitory properties of the full-length protein are in agreement with the flanking model, already proposed for interactions involving intrinsically disordered proteins (28, 29).

Unexpectedly, ITC studies indicated a negative contribution of the C-terminal region encompassed by P3 (higher $K_d$ and lower $ΔH$ values for P5 than for either P2 or the entire cE5 protein). This conflicts with the conservation of the C-terminal serine-rich extension among Old World anophelines, suggesting a potential functional role for this region. However, specific experimental conditions and/or the in vitro nature of our study should be taken into account. The situation may be different in vivo, where the regions encompassed by peptides P1 and P3 may provide interaction sites for other binding partners (e.g. the RGD tripeptide may be involved in binding to integrins) and/or undergo post-translational modifications (as suggested from the different mobility between recombinant and native cE5 protein in SDS-PAGE) (9). In conclusion, through structural, functional, and binding studies we shed additional light on the unique mechanism of thrombin binding and inhibition by this family of salivary anticoagulants from anopheline mosquitoes. It is anticipated here that future proteomic and structural studies on salivary proteins of hematophagous animals are expected to provide insights into novel antihemostatic molecules and novel mechanisms of disrupting blood clotting that may be of great help in the design of novel antithrombotics. This appears especially true considering (i) the fast evolutionary rate of salivary proteins (11, 12), (ii) the convergent evolutionary nature of hematophagy, and (iii) the fact that so far we acquired some knowledge on the salivary repertoires of just a few of the >14,000 arthropod species estimated to feed on blood (30).

Experimental procedures

cE5 protein and peptides

Recombinant cE5 and 15N-labeled cE5 proteins were expressed and purified as described (9). The ORF encoding P5 (cE5(Asp-310–Glu-82)) was chemically synthesized (Life Technologies) and cloned into pETM11 (Novagen). Expression was

Figure 6. Thrombin inhibition assays. Residual thrombin activity in the absence and the presence of A. gambiae cE5, or of the different peptides as indicated, was measured after the hydrolysis of the fluorogenic substrate Boc-Val-Pro-Arg-7-amido-4-methylcoumarin hydrochloride. Dots represent data from three independent experiments, each in triplicate. Thick lines show the mean values, and bars indicate S.D.
performed as above. Protein integrity and purity was confirmed by mass spectrometry (Agilent Q-TOF LC/MS). Peptides P1 (cE5(Ala-1–Ser-30), APQYARGDVPTYDEEDFDEESLK-PHSSSS), P2 (cE5(Asp-31–Arg-62), DDGEEEFDPSLLEEH-ADAPTARDPGRNPEFLR), and P3 (cE5(Asn-63–Glu-82), NSNTDEQASAPASSESE) were purchased from INBIOS S.r.l. (Naples, Italy). The *A. albimanus* anophelin was synthesized at the Peptide Synthesis Laboratory, Research Technologies Branch, NIAID, National Institutes of Health (Rockville, MD) by Dr. Jan Lukszo.

**Circular dichroism analyses**

CD spectra were recorded at 20 °C in the far UV (190–260 nm) on a Jasco J-710 spectropolarimeter (31). Each spectrum was obtained averaging three scans, subtracting contributions from corresponding blanks and converting the signal to mean residue ellipticity in units of degree × cm² × dmol⁻¹. Spectra were recorded using 10 μM cE5 in 10 mm phosphate buffer at pH 7.5 in the absence and in the presence of increasing concentrations (up to 60% (v/v)) of 2,2,2-trifluoroethanol (TFE, 99.5% isotopic purity, Sigma). Prediction analysis of secondary structure content was performed with CDPro (31, 32).

**Isothermal titration calorimetry**

ITC studies were performed at 22 °C with an ITC200 calorimeter (MicroCal/GE Healthcare). Anophelin and cE5 (100 μM) or the peptides P1, P2, P3, and P5 (150 μM) were titrated into a solution of human α-thrombin (20 μM; Hematologic...
**A. gambiae cE5–thrombin interaction**

Technologies Inc., HCT-0020). Before all titration experiments proteins and peptides were dialyzed against 20 mM HEPES, pH 7.5, 50 mM NaCl. All data were analyzed and fitted using the Microcal Origin version 7.0 software package. Binding enthalpy, dissociation constants, and stoichiometry were determined by fitting the data using a one set-of-site-binding model. ITC runs were repeated twice to evaluate the reproducibility of the results.

**Nuclear magnetic resonance spectroscopy**

NMR spectra were recorded at 298 K on a Varian Unity Inova 600 MHz spectrometer provided with a cold probe. 2D $^1$H, $^{15}$N HSQC spectra of $^{15}$N-labeled cE5 were acquired by implementing the following set of experimental conditions: 50 mM sodium phosphate buffer, pH 7.8; 550-μl sample total volume; H$_2$O/D$_2$O (98% deuterium, Armar Scientific, Switzerland) 90/10, 0.1%(w/v) NaN$_3$;5 –4 0

expansion were performed using GraphPad Prism 6.0 (GraphPad Software, Inc. La Jolla, CA). Analysis of titration experiments and overlays of 2D spectra were performed with Sparky (36). In addition, $^1$H, $^1$H NOESY 300 (37) and TOCSY 70 (total correlation spectroscopy) (38) experiments were recorded for the P2 peptide (470 μM in 50 mM sodium phosphate pH 7.8) in its unbound form and complexed to thrombin (25 μM).

**Thrombin inhibition assays**

Inhibition of human α-thrombin by cE5 and peptides P1, P2, P3, or P5 was evaluated after the hydrolysis of the fluorogenic substrate Boc-Val-Pro-Arg-7-amido-4-methylcoumarin hydrochloride (Boc, t-butoxycarbonyl; Sigma, B9385). Reactions were carried out in a 100-μl total volume in reaction buffer (50 mM Tris–HCl, pH 8.0, 50 mM NaCl, 20 μM CaCl$_2$, 0.01% (v/v) Triton X-100). Human α-thrombin (0.5 mM; Sigma, T7009) was preincubated for 10 min at 37 °C with 100 nm recombinant cE5 or cE5-derived peptides. Reactions were initiated by the addition of substrate (250 μM) and hydrolysis followed at 37 °C for 60 min at 460 nm (excitation = 360 nm) on a Synergy HT microplate reader (BioTek). Data represent three independent experiments, each one in triplicate. Graphics and statistical analysis were performed using GraphPad Prism 6.0 (GraphPad Software, Inc. La Jolla, CA).

**Platelet assays**

Inhibition of collagen-mediated platelet aggregation was evaluated using human platelet-rich plasma obtained from donors at the National Institutes of Health blood bank and diluted to a density of 2 × 10$^5$ platelets/μl in Tyrode-BSA (40) (final volume 300 μl). cE5, P1, or an equal volume of Tyrode buffer was added to each sample, which was then placed in an aggregometer (40) and stirred at 1200 rpm at 37 °C for 1 min before the addition of collagen (type-I fibrils, Chrono-log) to a concentration of 1.6 μg/ml.

**Crystallization of A. gambiae cE5 in complex with human α-thrombin**

Human α-thrombin (Hematologic Technologies) was mixed in 20 mM HEPES, pH 7.5, 125 mM NaCl with a 4-fold molar excess of P5 peptide (cE5(Asp-310–Glu-82)) and incubated on ice for 1 h. The resulting complex was concentrated by ultrafiltration using a 2-kDa cutoff centrifugal device (Sartorius). An initial crystallization screen at 20 °C on sitting drop geometry was performed at the High Throughput Crystallization Laboratory of the European Molecular Biology Laboratory (Grenoble, France). Preliminary crystallization conditions were systematically optimized in-house until single monoclinc crystals belonging to space group C2 were obtained after 4–6 days at 20 °C using the vapor diffusion sitting-drop method from drops consisting of equal volumes (1 μl) of protein complex (at 6.4 mg/ml) and precipitant solution (0.1 M PCTP, pH 5.0, 25% (w/v) PEG 1500) equilibrated against a 300-μl reservoir.

**Data collection and processing**

Crystals were cryoprotected by brief immersion in 0.1 M PCTP, pH 5.0, 20% (w/v) PEG1500, 20% (v/v) ethylene glycol or 0.1 M PCTP, pH 5.0, 35% (w/v) PEG1500 and flash-cooled in liquid nitrogen. Diffraction data were collected from two isomorphous crystals (2100 and 2400 images in 0.05° oscillation steps and 0.037-s exposure) on a Pilatus 6 M-F detector (Dectris) using a wavelength of 0.973 Å at beam line ID30B of the European Synchrotron Radiation Facility (Grenoble, France). Data were integrated with XDS (41), scaled with XSCALE (41), and reduced with utilities from the CCP4 program suite (42). Data collection statistics are summarized in supplemental Table 1. The raw experimental datasets for the three-dimensional structure reported in this manuscript (PDB 5NHU) are available in the SBGrid Data Bank (https://data.sbgrid.org/data/) (45) (Ripoll-Rozada, J., and Pereira, P. J. B. (2017) X-ray diffraction data for human thrombin-A. gambiae cE5 complex–datasets 1 and 2. Structural Biology Data Grid, V1, http://dx.doi.org/10.5281/zenodo.2581408).
Structure determination and refinement

The structure of the complex was solved by molecular replacement with Phaser (46) using the coordinates of free wild-type human α-thrombin (PDB entry 3U69; Ref. 22) as the search model. Alternating cycles of model building with COOT (47) and refinement with PHENIX (48) were performed until model completion (refinement statistics are summarized in supplemental Table 1). All crystallographic software was supported by SBGrid (49). Refined coordinates and structure factors were deposited at the PDB with accession number 5NHU.

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