Crotalase, a Fibrinogen-Clotting Snake Venom Enzyme: Primary Structure and Evidence for a Fibrinogen Recognition Exosite Different from Thrombin

Agnes H. Henschen-Edman¹, Ida Theodor¹,², Brian F.P. Edwards³, Hubert Pirkle²

From the Departments of Molecular Biology and Biochemistry¹ and Pathology², University of California, Irvine, CA, USA and the ³Department of Biochemistry and Molecular Biology, Wayne State University, Detroit, MI, USA

Summary

Crotalase, a fibrinogen-clotting enzyme isolated from the venom of *Crotalus adamanteus*, and its overlapping fragments were subjected to Edman degradation. The resulting amino acid sequence, VIGGDEC NINEHRFLVALDYWSQFLCGTTLINNEWLTAHCQRTHI LIYYVHDFSVCQDFKEQRFPPFKEKYFDCSNFTKWDKIM LIRLNPVSYSEHIAPLSLPPPIVGVSCRAMWGGTTSPQET LPDVPHCANLLDYEYVCRTAHPQFRLPATSRSTLCAVLEG GIDTCNRDSGPGPLICNGQFPQVFWGPDCAPAQDPKGLYT K VFDLHLDQSIAGKTVNCP, is characteristic of a serine proteinase. Comparison with thrombin, the physiological fibrinogen-clotting enzyme, showed that thrombin’s fibrinogen-recognition exosite (FRE) is poorly represented in crotalase. Hirudin, a FRE-dependent inhibitor, had no effect on crotalase. Spatial modeling of crotalase yielded a possible alternative fibrinogen-recognition site comprised of Arg 60F, Lys 85, Lys 87, and Arg 107 (underlined in the sequence above). Crotalase also lacks thrombin’s YPPW loop, as well as its functionally important ETW 146-148, and its heparin-binding site. The enzyme contains a single asparagine-linked glycosylation site, NFT, bearing neutral and amino sugars that account for 8.3% of the enzyme’s total molecular weight of 29,027. The calculated absorbance of crotalase at 280 nm, 1%, cm⁻¹ is 15.2.

Introduction

Thrombin is a serine proteinase which catalyzes the last in the series of narrowly specific proteolytic reactions that lead to the clotting of the blood protein fibrinogen. Crotalase (EC 3.4.21.30) is an enzyme contained in the venom of the Eastern diamondback rattlesnake *Crotalus adamanteus* (1) which, like thrombin, triggers the clotting of fibrinogen by hydrolytic release of its fibrinopeptide A (FPA) (2). Venom enzymes with this thrombin-like action are widely distributed, mainly within several pit viper genera (3), and afford the opportunity for comparative examination of the molecular structural requirements for thrombin’s narrow proteolytic action on fibrinogen.

In the present study, Edman degradation of the amino terminus and overlapping fragments of crotalase yielded a complete amino acid sequence with features characteristic of a serine proteinase. The fibrinogen recognition exosite (FRE) of thrombin is poorly represented in the crotalase sequence and this fact, along with the finding here that crotalase is not inactivated by hirudin (a FRE-dependent inhibitor), prompted a search for an alternative fibrinogen recognition site. The resulting spatial modeling of the crotalase molecule in the present work and in Massova et al. (4) reveals a possible alternative FRE comprised of Arg 60F, Lys 85, Lys 87, and Arg 107 [chymotrypsinogen numbering (15)].

Experimental Procedures

Materials

Lyophilized venom of the Eastern diamondback rattlesnake (*Crotalus adamanteus*) was purchased from Biotoxins, Inc. (St. Cloud, FL). The following materials were obtained from the sources given in parentheses: trypsin-TPCK (Worthington Biochemical Corp., Freehold NJ), Lysyl endopeptidase (Waco Pure Chemicals Industries, Tokyo), Sephadex G-50 and G-100 and SP Sephadex C-50 (Pharmacia), DEAE-cellulose, DE-52 (Whatman), Sepharose 4B aminocaprolic acid (New England Nuclear Co., Boston), N-Glycanase (Genzyme Corp.), hirudin (Boehringer Mannheim), human α-thrombin (gift from J.W. Fenton, II). Other reagents were the best available commercially.

Purification of Enzyme

Crotalase was purified from *Crotalus adamanteus* venom according to Bajwa and Markland (5). This procedure entails gel filtration on Sephadex G-100 followed by ion exchange chromatography on DEAE-cellulose, affinity chromatography on Sepharose 4B aminocaprolic acid (New England Nuclear Co., Boston), N-Glycanase (Genzyme Corp.), hirudin (Boehringer Mannheim), human α-thrombin (gift from J.W. Fenton, II). Other reagents were the best available commercially.

Carbohydrate Removal

Crotalase (5 mg/ml in 0.2 M phosphate buffer, 0.01 M EDTA, pH 8.6) was incubated with N-Glycanase (2 units/mg protein) at 37°C for 48 h and the deglycosylated protein was isolated by gel filtration on Sephadex G-50.

Protein Modification

Crotalase was denatured and reduced in 0.1 M Tris, pH 8.5 containing 6 M guanidine hydrochloride and 5% final concentration of 2-mercaptoethanol for 4 h under nitrogen at 37°C in the dark. Sulphydryl groups were carbamylated with [⁵⁷⁷] iodideoacetic acid (7) or were pyridylethylated with 4-ethylpyridine in a 1:1 molar ratio to all sulphydryl groups. Free sulphydryl groups were determined using 5,5’-dithiobis(2-nitrobenzoic acid) (8). Succinylation of free amino groups was carried out according to Habeeb et al. (9).
Chemical Cleavage

Methionyl bonds of reduced and alkylated crotalase were cleaved in 70% formic acid containing 10% (w/v) cyanogen bromide for 2 h at room temperature. Crotalase was cleaved at tryptophanyl bonds using o-iodosobenzoic acid (10), a 2-fold (w/w) excess of which was added to the reduced and alkylated protein dissolved in 80% acetic acid, 4 M guanidine hydrochloride.

Enzymatic Cleavage

Tryptic cleavage of reduced and alkylated crotalase was performed with or without prior succinylation in 0.1 M ammonium bicarbonate pH 8.2 with a 1:100 (w/w) enzyme to substrate ratio for 18 h at 37°C. Cyanogen bromide fragments of crotalase were digested with Lys-C endoproteinase at a 1:40 (w/w) enzyme to substrate ratio for 8 h at 37°C in 0.1 M Tris pH 8.6. Cyanogen bromide fragments were cleaved at Glu residues using Staphylococcus aureus V8 protease at a 1:40 (w/w) enzyme to substrate ratio for 3 h at 37°C in 0.1 M ammonium bicarbonate pH 8.0.

Peptide Isolation

Peptides generated by the chemical and enzymatic cleavage procedures were separated by open column chromatography on Sephadex G-50, eluted with a 0.2 M ammonium bicarbonate buffer pH 8.0, and on DE-52 using a linear NaCl gradient (0 to 0.2 M) in Tris buffer pH 7.8 or by high performance liquid chromatography using a Beckman model 332 instrument with a Vydac C18 RP column. The gradient was linear from a starting eluant of 0.1% trifluoroacetic acid to a limiting eluant of 0.1% trifluoroacetic acid in 80% or 50% acetonitrile.

Amino Acid Sequence Analysis

Automated Edman degradation was performed with an Applied Biosystems Model 477A sequencer equipped with an on-line Model 120A phenylthiohydantoin amino acid analyzer or, earlier, with a Beckman 890B sequencer, in which case the PTH amino acids were identified by high performance liquid chromatography and by thin layer chromatography. The analyzer of the Applied Biosystems sequencer allowed direct identification of pyridylethylated PTH cysteine; radio labeled, carboxymethylated PTH cysteine in the Beckman sequencer fractions was detected in a scintillation counter.

Molecular Modeling

The space-filling and secondary structure images of the crotalase molecule were generated by INSIGHT II (Kabsch-Sander algorithm for secondary structure; Molecular Simulations Inc.), using the coordinates of the model reported by Massova et al. (4).

Hirudin Inhibition

Crotalase (6.3 NIH thrombin-equivalent units/ml) was incubated with hirudin (30 antithrombin units/ml) in 0.15 M NaCl 0.02 M Tris pH 7.4 at 22°C. Hirudin was in 1.4-fold molar excess over crotalase. The mixture was periodically tested for fibrinogen-clotting activity (6) for 30 min. A comparable mixture of thrombin and hirudin showed immediate loss of fibrinogen-clotting activity.

Results and Discussion

Amino Acid Sequence Determination

The strategy for Edman degradation of crotalase and its overlapping fragments and the resulting amino acid sequence of the enzyme is given in Fig. 1. Here, the sequence analysis relied both on peptides generated by enzymatic digestion and on chemical cleavage of fragments produced by cyanogen bromide cleavage of the pyridylethylated or carboxymethylated protein.

Primary Structural Features of Crotalase

Comparison of the amino acid sequence of crotalase to those of chymotrypsin, kallikrein, trypsin, and thrombin [Fig. 2, chymotrypsin numbering (15)] reveals typical characteristics of a serine protease. The residues of the catalytic triad, His 57, Asp 102, and Ser 195, are in their expected positions, flanked by sequences that are
largely conserved. The only nonconserved neighboring residue is Arg
193, a position which turns out to be variable also in other thrombin-
like enzymes (TLEs) (Fig. 2). As in all other serine proteinases position
194 of crotalase is occupied by an Asp residue which makes an impor-
tant contribution to stabilizing the active center in a catalytically active
configuration. In general this is accomplished by forming a salt bridge
with the enzyme’s N-terminal α-amino group (11) and such a salt
bridge is a feature of the modeled three-dimensional structure of

crotalase (4).

Since crotalase has no free sulfhydryl groups (1) it is likely that all of
its cysteines are disulfide bonded to one another. Based on homology
with trypsin (12) and with another thrombin-like venom enzyme (13),
the disulfide bridges of crotalase are predicted to link the following
sequence positions: 22-157, 42-58, 91-250, 136-201, 168-182, and
191-220.

The overall rates of sequence identity between crotalase and the
mammalian serine proteinases in Fig. 2 are kallikrein 33%, trypsin
32%, thrombin 30%, and chymotrypsin 24%. As might be expected, the
TLEs exhibit a much higher degree of similarity among themselves
with a mean rate of identity of 64%. The primary structures of crotalase
and the other TLE are also strikingly similar to venom serine proteinases
that do not clot fibrinogen (62% mean rate of identity) (14).

Modeling of the spatial structure of the crotalase molecule shows its
core to be comprised, like thrombin (15), of two six-stranded
β-barrels between which lies the active center (Fig. 3). Other secondary structu-
ral features are three
α-helices formed from segments Tyr 165-Ala-171,
Val 231-Leu 235, and Trp 237-Ala 243, homologous to three of
thrombin’s five B-chain helices.

A single consensus site for asparagine-linked glycosylation, Asn 94,
was first inferred from a sequencing blank two steps before Thr 96 and
later confirmed after enzymatic deglycosylation of the crotalase
molecule. The total neutral and amino sugar content of crotalase is
8.34% with molar ratios of mannose 3, galactose 6.1,
N-acetylglucosamine 10.5,
N-acetylgalactosamine 0.6, and fucose 1.5, a composition
consistent with a complex type of asparagine-linked chain (16) and not
with
O-linkage. Analysis for sialic acid was not done since earlier

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**Fig. 2** Comparison of amino acid sequences of thrombin-like venom enzymes with bovine thrombin and with other serine proteinases. Dashes indicate gaps introduced to optimize the alignment. All residue positions correspond to the first digit of the residue number. CHY, bovine chymotrypsin (35); KAL, rat pancreatic kallikrein (36); TRY, bovine trypsin (37); THB, bovine thrombin B-chain (38); Cro, crotalase (39); Anc, anced (40); Mut, *Lachesis muta muta* (41); Bot, bothrombin (42); Bat, batroxo-
bin (12); Fla, flavoxobin (43), Bil, bilineobin (13), Cal, calobin (44). Chymotrypsinogen numbering (15)

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analyses yielded values that varied from one crotalase preparation to another (1, 5). Although N-acetylgalactosamine, as found here, does not ordinarily occur in asparagine-linked carbohydrate chains, it has now been demonstrated as subterminal to sialic acid residues of batroxobin (17, 18) and ancrod (19), partially or completely replacing galactose in that structural role. The unevenness of molar ratios among the individual sugars of crotalase most likely reflects the heterogeneity of carbohydrate chain structure often exhibited by glycoproteins.

From the amino acid sequence of crotalase its molecular weight can be computed as 26,606 without carbohydrate or 29,027 adjusted for the sugar content just given. From the enzyme’s tryptophan, tyrosine, and cystine content its molar absorption coefficient, \( \epsilon(280)(M^{-1} cm^{-1}) \), can be calculated as 44,180 (20, 21). Then, assuming a molecular weight of 29,027, the absorbance of crotalase at 280 nm, 1%, cm\(^{-1}\) is 15.2.

Comparison of Crotalase Sequence with Thrombin

An important role in fibrinogen recognition and cleavage specificity has been attributed to the 60A-60D (YPPW) insertion loop of thrombin which forms part of the rim of its active site cleft (22). This feature is absent in crotalase, leaving a more open entrance to the active site (4). This feature is also absent in the mutant des-PPW thrombin (23). But both crotalase and the mutant thrombin exhibit, nonetheless, a narrow specificity for fibrinogen as a coagulant substrate. By contrast, a component of the opposite rim of the active site cleft of thrombin, ETW 146-148, also absent in crotalase, has been shown in mutant forms of the thrombin molecule, crotalase possesses only three and none of the three is conserved among the other TLEs (Fig. 2).

The absence in the TLEs of a cationic region homologous to the FRE of thrombin raises the question of whether the corresponding region of the crotalase might contain a similar, positively charged site made up of nonhomologous residues. But, in contrast to thrombin’s net positive charge of 5 in the critical center of the FRE (positions 67-82), the same crotalase sequence has a positive charge of only 1 and most of the other TLEs have a negative net charge, usually of 1 (Fig. 2).

In addition to the foregoing structural grounds for doubting that crotalase recognizes fibrinogen through an interaction at the same region of the molecule used by thrombin, the following three function-

Specificity for the release of FPA by thrombin is also served by a hydrophobic binding site near the catalytic center comprised of Tyr 60A, Trp 60D, Leu 99, Ile 174, and Trp 215 which interacts, in particular, with Phe 8 of the fibrinopeptide (26). As crotalase lacks any part of the 60 insertion loop, this hydrophobic site is formed instead by Trp 97A, Phe 174A, Leu 176, and Trp 215 and is favorably configured to accommodate Phe 8 of FPA (4). Other residues of crotalase that are homologous to thrombin’s FPA-interacting amino acids are Lys 97 and Gly 216 (26).

Of the seven basic residues of thrombin that form its heparin-binding site (exosite 2) (15), only one is represented in crotalase, this in keeping with the insensitivity of crotalase to heparin (27).

Fibrinogen Recognition Exosite (FRE)

The remarkably narrow specificity of thrombin for fibrinogen and other macromolecular substrates is thought to reside largely in a surface groove that extends “southeast” from the active site cleft, using the conventional orientation of the molecule (28). The interaction of this fibrinogen recognition site (FRE) with a complementary site on the fibrinogen molecule has been, to a major extent, attributed to several positively charged residues at or immediately subjacent to this groove. Eleven basic residues of thrombin [R35, K36, R67, K70, R73, R75, R77A, K81, K109, R110 (human and bovine), K149E (human)] have been implicated in one way or another in this electrostatic interaction with fibrinogen (29, 30). However, of these 11 basic residues of the thrombin molecule, crotalase possesses only three and none of the three is conserved among the other TLEs (Fig. 2).

The absence in the TLEs of a cationic region homologous to the FRE of thrombin raises the question of whether the corresponding region of the crotalase might contain a similar, positively charged site made up of nonhomologous residues. But, in contrast to thrombin’s net positive charge of 5 in the critical center of the FRE (positions 67-82), the same crotalase sequence has a positive charge of only 1 and most of the other TLEs have a negative net charge, usually of 1 (Fig. 2).

In addition to the foregoing structural grounds for doubting that crotalase recognizes fibrinogen through an interaction at the same region of the molecule used by thrombin, the following three function-

Fig. 3 Secondary structure diagram of the model of crotalase. The program INSIGHT II (Kabsch-Sander algorithm) was used to represent beta sheets as arrows (ribbons), helices as cylinders, and turns as sleeves
al considerations point in the same direction. First, crotalase is not inactivated by the thrombin inhibitor hirudin. During 30 min of incubation (Experimental Procedures section, Hirudin Inhibition) hirudin-crotalase mixtures showed no change from control (hirudin-free) fibrinogen clotting times. In contrast, the thrombin-hirudin mixtures exhibited immediate loss of fibrinogen-clotting activity. Since it has now been shown directly that hirudin inhibits thrombin by binding to its FRE (31, 32), the failure of hirudin to affect the clotting activity of crotalase suggests a different fibrinogen recognition site. And hirudin-insensitivity is shared with other TLEs (3). A second functional consideration is that crotalase does not promote platelet aggregation (27). Triggering of platelet aggregation by thrombin entails cleavage of platelet thrombin receptor, mediated by binding of the receptor to the FRE of the enzyme (33). Thus, absence of a thrombin-like FRE in crotalase would constitute a sufficient explanation for the inability of crotalase to bring about platelet aggregation. Finally, Binnie and Lord (34) have shown that, while a fibrinogen peptide complement­ary to the FRE of thrombin competitively inhibits FPA release and clotting by thrombin, this peptide had no effect on these activities of crotalase’s close homolog batroxobin.

The evidence for lack of a thrombin-like FRE in the crotalase molecule prompted a search for an alternative cationic fibrinogen recognition site. Fig. 4, generated using the coordinates of Massova et al. (4), shows the basic residues that present at the surface of the crotalase molecule. The right-hand lower part of the model shows the region that corresponds to the FRE of thrombin, as has already been discussed. The basic residues at the upper left can also be excluded as possible components of an alternative FRE as they are not conserved among the TLEs. The remaining residues, Arg 60F, Lys 85, Lys 87, and Arg 107 line a groove that extends from the active site cleft toward the northeast. Three of these residues are absolutely conserved among the TLEs and one is largely conserved. In the thrombin molecule, three of these positions are also occupied by basic residues but a crotalase-like FRE groove is absent, as it is occupied by thrombin’s 60A-60I loop (4). Thus, the cationic groove on the surface of the crotalase molecule, favorably oriented to its active site cleft, is a plausible candidate for an alternative fibrinogen recognition exosite.

It still must be considered, however, that fibrinogen recognition by crotalase may depend on totally different forces, e.g. hydrophobic interactions at yet a different exosite. Also, it is conceivable that interactions near the active site of crotalase may be sufficiently constrained to account for the observed degree of fibrinogen specificity.

Acknowledgments

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References