Purification and Characterization of Multisquamase, the Prothrombin Activator Present in Echis Multisquamatus Venom

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Abstract

The venom of Echis multisquamatus (Central Asian sand viper) contains a single prothrombin activator, designated multisquamase, which is structurally and functionally different from ecarin, the prothrombin activator from the venom of Echis carinatus (saw-scaled viper). Multisquamase is comprised of a 58000 Mr and a 23000 Mr subunit that consists of two disulfide-linked chains of 12000 Mr and 10000 Mr, respectively. In contrast to ecarin, which activates prothrombin and prethrombin 1 at comparable rates, and whose activity is hardly affected by Ca\(^2+\) or by changes in ionic strength, multisquamase hardly activates prethrombin 1; prothrombin activation requires Ca\(^2+\) and is strongly inhibited at high ionic strength. The most favourable kinetic parameters are observed at 1 mM Ca\(^2+\) and at low ionic strength (K_m=0.085\,\mu\text{M} and k_{cat}=0.68\,s^{-1} at 1\sim0.04). An increase in ionic strength considerably reduces the rate of prothrombin activation, due to an increase of the K_m (K_m=0.8\,\mu\text{M} and k_{cat}=1.03\,s^{-1} at 1\sim0.2). Studies in plasmas from patients on oral anticoagulant therapy show that E. Multisquamatus venom only activates carboxylated prothrombin, whereas E. carinatus activates both prothrombin and descarboxyprothrombin. Thus, multisquamase-dependent prothrombin activation appears to require post-translational modification of the gla-domain. This venom prothrombin activator may, therefore, become a useful tool to quantitate prothrombin and descarboxyprothrombin in cases where vitamin K-dependent carboxylation of prothrombin is impaired. © 1998 Elsevier Science Ltd.

Key Words: Snake venom; Prothrombin activation; Echis multisquamatus; Echis carinatus

Thrombin formation, one of the central reactions of blood coagulation, is the result of limited proteolysis of the zymogen prothrombin by blood coagulation factor Xa. Under physiological conditions, prothrombin activation is stimulated in a multiplicative way by the accessory components: Ca\(^2+\), negatively charged phospholipids and the protein cofactor, factor Va which together with factor Xa assemble in the so-called prothrombinase complex.

Prothrombin can also be activated by proteases present in snake venoms [1] from, amongst others, Australian Elapidae, Crotalidae, Viperidae, and Colubridae. Compared with factor Xa, venom activators can exhibit different mechanisms of prothrombin activation and requirements for cofac-

Abbreviations: PPACK, phenylalanyl-prolyl-arginine chloromethyl ketone; BCA, bicinchoninic acid; FPLC, fast protein liquid chromatography; INR, international normalized ratio; USP, U.S. Pharmacopoeia; BSA, bovine serum albumin.

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A well characterized and widely used prothrombin activator is ecarin [2,3], the metalloprotease present in Echis carinatus venom, that converts prothrombin exclusively into meizothrombin [4,5]. Ecarin not only activates prothrombin, but also prethrombin 1 and the descarboxy form of prothrombin [6-9]. Recently, Yamada et al. published the characteristics of a second prothrombin activator, called carinactivase, present in the venom from Echis Carinatus, with properties different from those reported for ecarin [7].

Here we report the purification and characterization of the prothrombin activator from the venom of Echis multisquamatus.

1. Materials and Methods

1.1. Materials

Hepes, Tris, EDTA, bovine serum albumin, and ovalbumin were purchased from Sigma Chemical Co. St. Louis, USA. The chromogenic substrates S2238 and S2366 were supplied by Chromogenix, Målndal, Sweden. Ecarin, the prothrombin activator from Echis carinatus venom was from Pentapharm, Basel, Switzerland. Crude Echis multisquamatus venom was from Latoxan, Rosans, France. Heparin (USP activity 175 units/mg) was from Leo Pharmaceuticals, Weesp, The Netherlands. Micro BCA protein assay kits were from Pierce, Rockford, USA. Immobilon-P membranes were from Millipore, Bedford MA 01730. Antibodies were from Kordia, Leiden, The Netherlands and Sigma Chemical Co. St. Louis, USA. Materials used for protein purification and the FPLC-system were from Pharmacia, Uppsala Sweden.

1.2. Proteins

Normal human plasma was obtained by collecting nine parts of blood from healthy volunteers in one part of 0.13 M trisodium citrate (pH 7.8). The blood was centrifuged twice for 15 minutes at 3,000Xg at room temperature. Plasma from orally anticoagulated patients was a kind gift of Dr. Hamulyak (Dept. of Hematology, University Hospital, Maastricht). Human prothrombin was purified from fresh frozen plasma according to DiScipio et al. [10]. Prethrombin 1 was purified from a reaction mixture of human prothrombin with thrombin in 50 mM Tris (pH 7.9), 175 mM NaCl. After inhibition of thrombin with PPACK, the reaction mixture was applied to a column (XK 5/10) with Q Sepharose fast flow. Prethrombin 1 did not adhere to the resin and fragment 1, and non-cleaved prothrombin were eluted with a gradient of 250 to 600 mM NaCl. Human plasma, prothrombin, prethrombin 1, fragment 1, and purified venom activators were stored at -80°C. Protein preparations were homogeneous and >95% pure as judged by SDS-PAGE according to Laemmli [11].

1.2. Protein Concentrations

Protein concentrations were determined with the micro BCA protein assay [12]. Prothrombin concentrations were determined after complete activation of prothrombin with Echis carinatus venom as described earlier [13].

1.3. Methods

1.3.1. Prothrombin Activation

Prothrombin activation was determined by measuring the generation of amidolytic activity towards chromogenic substrates (S2238 or S2366). Human prothrombin or prethrombin 1 was preincubated in a buffer containing 25 mM Hepes (pH 7.7), 175 mM NaCl at 37°C. When divalent cations were present, these were included in the preincubation mixture. Activation was started by addition of appropriate dilutions of crude venom or purified venom activator in the same buffer. After different time intervals samples were withdrawn from the activation mixtures and transferred to cuvettes containing a final volume of 1 ml of 50 mM Tris-HCl (pH 7.9), 175 mM NaCl, 20 mM EDTA, 0.5 mg/ml ovalbumin, and 235 μM S2238 at 37°C, and the amidolytic activity was quantitated as described earlier [14]. The amounts of activated prothrombin were calculated from a calibration curve made with known amounts of active site-titrated thrombin.

During purification, the activity of the venom activator was routinely assayed for prothrombin activation in reaction mixtures that contained 0.1 μM prothrombin, 3 mM CaCl₂, and appropriate dilutions of the venom activator.

When plasma prothrombin was activated, the plasma was diluted 2000-fold in a buffer containing 25 mM Hepes (pH 7.7), 175 mM NaCl, 1 mM CaCl₂ and preincubated at 37°C. Activation was started by addition of appropriate dilutions of crude venom in the same buffer. After different time intervals the amount of prothrombin activated was quanti-
tated by measuring the generation of amidolytic activity towards S2366 on a SLT platereader set in the dual wavelength mode at 405–492 nm. Since crude Echis multisquamatus venom shows a remarkably high amidolytic activity towards S2238, the activation of plasma prothrombin was quantitated with S2366 in these experiments.

1.3.2. Kinetic Analysis
Initial rates (v) of prothrombin or prethrombin 1 activation were determined at varying substrate concentrations (S) and the kinetic parameters $K_m$ and $V_{max}$ were obtained by fitting the v vs. S curves to the Michaelis-Menten equation ($V = V_{max}S/(K_m + S)$) using nonlinear least squares regression analysis.

1.3.3. Gel Electrophoretic and Immunoblot Techniques
Gel electrophoresis of the purified activator was carried out in the presence of SDS on 13% slabs [11] in a MiniProtean II cell from BioRad and the gels were stained with Coomassie Brilliant Blue R-250. Products of prothrombin proteolysis in plasma were identified after SDS-PAGE on 10% slab gels, electrophoretic transfer of proteins from the gel to Immobilon-P membranes, and visualization with a monoclonal mouse-antibody directed against human prothrombin and anti-mouse IgG conjugated with alkaline phosphatase [15].

2. Results

2.1. Purification of the Prothrombin Activator from Echis Multisquamatus

The prothrombin activator present in the crude venom from Echis multisquamatus was purified by ion-exchange chromatography on Resource Q and Mono S at pH 7.5. The activator adhered to Resource Q and was eluted with a linear salt gradient (Fig. 1a) resulting in a purification of about 5-fold with a 76% yield (Table 1). Subsequent chromatography on Mono Q (Fig. 1b) resulted in a further two-fold purification and a final overall yield of 13% (Table 1). SDS-PAGE on 13% slab gels (Fig. 2) showed that the purified activator contains two protein bands of 58000 and 23000 Mr, respectively, and that the 23000 band consisted of two polypeptide chains held together by disulfide bond(s) as judged from the reduced sample (Fig. 2, lane 3). The activator eluted as a single symmetrical peak during gel permeation chromatography of Superdex-200 at $Mr=67000$ (data not shown). Attempts to separate the two protein bands by incubation with 0.8 M NaSCN invariably resulted in complete loss of prothrombin converting activity. We conclude, therefore, that the purified activator is a two subunit enzyme. Furthermore, multisquamase likely is a metalloproteinase since treatment with EDTA (0.1–

![Fig. 1. Ion-exchange chromatography of the prothrombin activator from Echis multisquamatus. Panel A. 6 mg crude venom was dissolved in 22 ml of 25 mM Hepes (pH 7.5), 50 mM NaCl, and 3 mM CaCl$_2$, applied at a flow rate of 0.5 ml/min to a 1 ml Resource Q column and washed with 20 ml startbuffer. Bound protein was eluted with a linear gradient of 2×5 ml (50 mM to 500 mM NaCl) in the same buffer. Protein was monitored by $A_{280}$ (solid line) and fractions were tested for prothrombin activation (■) as described under “Material and Methods”. Panel B. Prothrombin activator-containing fractions obtained after chromatography on Resource Q were pooled, diluted in 25 mM Hepes (pH 7.5), 3 mM CaCl$_2$, buffer to a final NaCl concentration of 50 mM, and applied to a 1 ml Mono S column. The activator was eluted with a 2×5 ml linear gradient (50 mM to 500 mM NaCl) in 25 mM Hepes (pH 7.5), 3 mM CaCl$_2$.](image-url)
Table 1. Purification of the prothrombin activator from Echis multisquamatus venom

<table>
<thead>
<tr>
<th></th>
<th>Total Protein (mg)</th>
<th>Specific Activity (nmol PT-activated/ min/μg venom)</th>
<th>Recovery (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crude venom</td>
<td>5.21</td>
<td>0.22</td>
<td>100</td>
</tr>
<tr>
<td>Resource Q</td>
<td>0.78</td>
<td>1.11</td>
<td>76</td>
</tr>
<tr>
<td>Mono S</td>
<td>0.06</td>
<td>2.38</td>
<td>13</td>
</tr>
</tbody>
</table>

Protein was determined with the micro BCA assay [12]. Prothrombin activation was determined as described under “Material and Methods.” PT=prothrombin.

0.3 mM) resulted in a rapid loss of prothrombin converting activity (data not shown). From the final specific activity obtained it appears that an unusually high percentage (-9%) of the crude venom actually consists of prothrombin activator (Table 1).

2.2. Prothrombin and Prethrombin 1 Activation

Rates of prothrombin and prethrombin 1 activation with purified activator were determined and compared to those obtained with ecarin, the prothrombin activator present in Echis carinatus venom. It was verified that time courses of prothrombin and prethrombin 1 activation were linear with time and proportional to the amount of enzyme present (data not shown). Table 2 summarizes the initial rates of activation of 1 μM prothrombin or prethrombin 1 by 1 ng/ml ecarin or multisquamase. Prothrombin was activated some threefold faster by ecarin than by multisquamase. However, whereas ecarin activated prethrombin 1 and prothrombin at comparable rates, multisquamase was almost unable to activate prethrombin 1 whose rate of activation was approximately 500-fold slower than that of prothrombin (Table 2).

Figure 3 shows an experiment in which initial rates of 0.1 μM prothrombin/prethrombin 1 activation by 4.5 ng/ml ecarin or 10 ng/ml multisquamase were determined at varying NaCl concentrations. Due to the fact that multisquamase-dependent prethrombin 1 activation was extremely slow the latter reaction was followed at a 50-fold higher concentration (5 μM). At 30 mM NaCl, ecarin was a somewhat better prothrombin activator than multisquamase but, as can be seen, both prothrombin and prethrombin 1 activation by multisquamase were strongly inhibited at higher ionic strength. Furthermore, at approximate physiological ionic strength, prothrombin activation by multisquamase was some 50-100 fold faster than prethrombin 1 activation. In contrast, ecarin-catalyzed activation of prothrombin or prethrombin 1 was much less affected by variation of the ionic strength and even at 400-500 mM NaCl, prothrombin and prethrombin 1 activation occurred at ~40% of the rate observed at 30 mM NaCl (Figure 3).

The kinetic parameters of prothrombin and prethrombin 1 activation by ecarin and multisquamase,

![SDS-PAGE experiment](image-url)
Table 2. Rate of prothrombin and prethrombin 1 activation by ecarin and multisquamase

<table>
<thead>
<tr>
<th>Activator</th>
<th>Prothrombin (nM activated/min/ng activator)</th>
<th>Prethrombin 1 (nM activated/min/ng activator)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ecarin</td>
<td>1.52</td>
<td>1.01</td>
</tr>
<tr>
<td>Multisquamase</td>
<td>0.51</td>
<td>0.001</td>
</tr>
</tbody>
</table>

The initial rate of activation of 1μM prothrombin or prethrombin 1 by 1 ng/ml ecarin or multisquamase was determined at 37°C in 25 mM Hepes (pH 7.7), 175 mM NaCl, 5 mg/ml BSA, and 1 mM CaCl₂, as described under "Materials and Methods."

determined at 30 mM NaCl (I=0.04) and at 175 mM NaCl (I=0.2), are summarized in Table 3. The Vmax of prothrombin and prethrombin 1 activation were minimally affected by an increase in the ionic strength. The low rate of prethrombin 1 activation by multisquamase is explained by its unfavourable Km which is approximately 200-fold higher than the Km for prothrombin. It appeared that the inhibition observed for multisquamase at high ionic strength was mainly due to an increase of the Km. The Km for prethrombin 1 activation by multisquamase at high ionic strength was increased so much that it was impossible to obtain an accurate Km value (Table 3). It appears, therefore, that the interaction of multisquamase with prothrombin or prethrombin 1 is driven (at least for a major part) by electrostatic interactions and for efficient interaction to occur, the presence of fragment 1 is required.

However, prethrombin activation by multisquamase appears to be also strongly dependent upon the presence of Ca²⁺ as shown in Figure 4. Other metal ions known to induce the conformational change in prothrombin (Mg²⁺ or Mn²⁺ but not Ba²⁺) [16] also stimulated multisquamase-catalyzed prothrombin activation (data not shown). In contrast, ecarin-dependent prethrombin or prethrombin 1 activation was hardly influenced by addition of divalent metal ions.

2.3. Activation of Prothrombin and Descarboxyprothrombin by Ecarin and Multisquamase

The above data strongly suggest that multisquamase not only requires an intact but also a carboxylated prothrombin molecule for efficient activation. To test this, prothrombin present in normal plasma and in plasma from a patient treated with vitamin K antagonists was activated with multisquamase. For comparison the same plasma samples were also activated with ecarin (ecarin activates prothrombin, prethrombin 1, and descarboxyprothrombin equally [6-9]. The immunoblot presented in Figure 5 shows that both activators converted all prothrombin present in normal plasma and bands indicative for meizothrombin-des-fragment 1 and (meizo) thrombin-antithrombin III complexes became visible. In patient plasma, ecarin also activated all prothrombin and descarboxyprothrombin, but with multisquamase only part of the prothrombin present was converted. This indicates that multisquamase indeed only activated carboxylated prothrombin.

Thus, in principle the combination of prothrombin activation with ecarin and multisquamase should yield relevant information concerning the coagulant status of prothrombin present in a given plasma sample. Figure 6 shows the result of an experiment in which the prothrombin and descarboxyprothrombin present in a number of different plasma samples...
Table 3. Kinetic parameters of prothrombin and prethrombin 1 activation

<table>
<thead>
<tr>
<th></th>
<th>30 mM NaCl</th>
<th></th>
<th>175 mM NaCl</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>$K_m$ (μM)</td>
<td>$V_{max}$ (nmol activated min/μg act)</td>
<td>$K_m$ (μM)</td>
</tr>
<tr>
<td>Ecarin</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prothrombin</td>
<td>0.05</td>
<td>2.55</td>
<td>0.15</td>
</tr>
<tr>
<td>Prethrombin 1</td>
<td>0.03</td>
<td>1.55</td>
<td>0.12</td>
</tr>
<tr>
<td>Multisquamase</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prothrombin</td>
<td>0.09</td>
<td>0.61</td>
<td>0.8</td>
</tr>
<tr>
<td>Prethrombin 1</td>
<td>22.92</td>
<td>1.11</td>
<td>&gt;100</td>
</tr>
</tbody>
</table>

$K_m$ and $V_{max}$ were obtained by fitting the Michaelis-Menten ($v$ vs. $S$) curves using non-linear least squares regression analysis. $k_{cat}$ was calculated assuming a Mr of 55000 for ecarin [6] and 67000 for multisquamase (see Results). Final activation mixtures contained 25 mM Hepes (pH 7.7), 1 mM CaCl$_2$, 5 mg/ml BSA, 30 or 175 mMNaCl, appropriate dilutions of venom activators, and varying concentrations of prothrombin or prethrombin 1. n.d. = not determined.

(randomly chosen from orally anticoagulated individuals, based on the INR values and independent of medical background) was activated with crude E. carinatus or with E. multisquamatus venom. Since multisquamase is the only prothrombin activator present in E. multisquamatus venom it is not necessary to isolate multisquamase from the crude venom for specific activation of prothrombin in a plasma sample. To yield accurate measurement of prothrombin levels (without interference of endogenous plasma inhibitors) the plasma samples were diluted 2000-fold before activation. A good correlation was observed when the ratio of the amidolytic activity obtained with E. carinatus (prothrombin plus descarboxyprothrombin) and with E. multisquamatus venom (prothrombin) was plotted against the INR value.

3. Discussion

The venom from the viper Echis multisquamatus contains a strong procoagulant which is capable of activating prothrombin. This procoagulant, which we call multisquamase, is a metalloproteinase that consists of two subunits of 58000 and 23000 Mr. The smaller subunit is comprised of two polypeptide chains linked together via a disulfide bridge.

The structural and functional properties of the E. multisquamatus activator are similar to those of carinactivase, a second prothrombin activator that was recently discovered to be present in the venom from E. carinatus [7]. However, the catalytic efficiency of multisquamase is not only dependent on the presence of Ca$^{2+}$ but is also strongly affected by salt concentration. In addition, multisquamase is the only enzyme with prothrombin-converting activity present in the crude venom of E. multisquamatus. The fact that the crude venom did not activate prethrombin 1 at physiological ionic strength is indicative for the absence of an ecarin-like prothrombin activator in E. multisquamatus venom.

The catalytic efficiency of prothrombin activa-
Fig. 5. Immunoblot analysis of plasma prothrombin activation by ecarin and by the purified activator from E. multisquamatus. 40-fold dilutions of normal human plasma (lanes 1-3) and plasma from a patient on oral anticoagulant therapy (lanes 4-6) in a buffer containing 25 mM Hepes (pH 7.7), 175 mM NaCl, 1 mM CaCl2 were activated with ecarin or multisquamase to result in complete activation of prothrombin within 5 min. After the amidolytic activity reached a constant level, 16 μl aliquots were subjected to SDS-PAGE (13% gel) according to Laemmli [11]. Prothrombin and related products were visualized after transblotting to immobilon membranes and immunologic staining as described under “Materials and Methods.” The relative positions of reference molecular weight markers are indicated. Lane 1, normal plasma; lane 2, normal plasma+ecarin; lane 3, normal plasma+multisquamase; lane 4, patient plasma; lane 5, patient plasma+ecarin; and lane 6, patient plasma+multisquamase.

The kinetic parameters of prothrombin or prethrombin 1 activation by ecarin are hardly affected by the reaction conditions and the enzyme does not show a similar requirement for the presence of Ca2+ and Gla residues.

The small pilot study, in which patient plasma prothrombin levels were determined both with the crude venoms from Echis carinatus and Echis multisquamatus, shows that the combination of these measurements correlated reasonably well with the INR values obtained for the individual plasma samples. These data indicate that Echis multisquamatus venom may be of future use as a tool to quantitate functional prothrombin in those cases where vitamin K-dependent carboxylation of prothrombin is hampered.

This work was supported by Program Grant 900–526–192 from the Dutch Organization for Scientific Research (NWO). R.J. Petrovan was supported by the Deutscher Akademischer Austauschdienst and Deutsche Akademie der Naturforscher “Leopoldina.” We thank Dr. Hamulyak (Dept. of Hematology, University Hospital Maastricht) for donating the plasma samples of orally anticoagulated patients.
Fig. 6. Estimation of anticoagulation in plasma from orally anticoagulated patients. Plasma samples (diluted 1/2000) were activated with crude E. carinatus and E. multisquamatus venom. Amidolytic activity present after 30 minutes was determined using S2366 as a substrate, and the ratio of amidolytic activities formed by E. carinatus and E. multisquamatus (EC/EM) is plotted vs. the INR values. The solid line was obtained by linear regression analysis (R=0.83). Further details are given under “Materials and Methods.”

References