Procaspase-3 activation by a metalloprotease secreted from Vibrio vulnificus

HYO YOUNG KIM1, ALAN K. CHANG1, JUNG EUN PARK1,2, IL-SEON PARK1, SEONG MYEONG YOON3 and JUNG SUP LEE1,2,4

1Research Center for Proteineous Materials, 2BK21 Research Team for Protein Activity Control, Departments of 3Marine Life Science, 4Biotechnology, Chosun University, Gwangju 501-759, Korea

Received May 10, 2007; Accepted June 29, 2007

Abstract. Vibrio vulnificus is a marine bacterium and a human pathogen capable of causing wound infection and septicemia. We previously showed that the metalloprotease vEP secreted by V. vulnificus activates prothrombin in vitro. To further investigate the ability of vEP to activate other zymogens, we used a mutant form of procaspase-3 which lacks the native cleavage sites as a zymogen. The mutant zymogen was activated by vEP to yield a mature enzyme with a maximum increase in caspase-3 activity of approximately 14-fold in a time-dependent manner. However, the increase started to decline with prolonged incubation and with higher protease concentration as a result of further cleavage of the mature enzyme. Western blot analysis revealed a band of ~17 kDa for the cleavage product, which corresponded with the change in caspase-3 activity. The activated procaspase-3 by vEP was also able to cleave poly(ADP-ribose) polymerase in a cell-free system, and was inhibited by Ac-DEVD-CHO, a potent caspase-3 inhibitor. The results presented are the first to demonstrate the in vitro activation of one of the crucial enzymes involved in cell death by a bacterial extracellular metalloprotease.

Introduction

Vibrio vulnificus (V. vulnificus) is an opportunistic human pathogen causing wound infection and septicemia (1,2). It secretes a zinc metalloprotease that has been reported to have many biological functions. These include the degradation of various plasma proteins and vascular permeability enhancement through the generation of inflammatory mediators. The wide spectrum of its activities in an in vitro system has made it a very interesting protease. It has not yet been conclusively demonstrated whether these biological activities of the enzyme contribute to the pathogenicity exerted by the bacterium V. vulnificus, although it has been shown that injection of the purified enzyme into animals produced some of the pathogenic symptoms observed with the bacterial infection (3,4). We previously reported that the protease secreted by V. vulnificus, ATCC 29307, which we named vEP, is a broad specificity enzyme that activates prothrombin in vitro (5). The cleavage of prothrombin produced a number of fragments and one of these corresponded to that of α-thrombin as shown by Western blot analysis. The cleavage site occurred at Thr271-Ala272, which is very near the site (Arg271,Thr272) recognized by Factor Xa, the physiological prothrombin activator. Based on this property, we speculated that vEP might also activate the zymogens of other proteases if it cleaves these zymogens at sites that are very near to the native cleavage sites for their activation.

Treatment of HeLa and NIH3T3 cells with vEP resulted in cell death (unpublished data). However, the mode of cell death exerted by vEP has not yet been established, and so far there is no evidence that this enzyme causes direct apoptosis, which is a tightly regulated cellular event that plays a critical role in the development and tissue homeostasis of metazoans (6,7). At the molecular level, apoptosis is tightly controlled and is mainly orchestrated by the activation of the aspartate-specific cysteine proteases known as caspases (8,9). The degradation of certain proteins by members of the caspase family is a general event taking place in cells undergoing apoptosis. The most prevalent caspase in the cell is caspase-3. It is a mediator caspase that is ultimately responsible for the majority of apoptotic effects. Inhibition of caspase-3 or caspase-3-like proteases in various cells has been shown to block apoptosis (10,11). In addition to its role in cell death, caspase-3 is important for survival, as caspase-3-knockout mice are born at a low frequency and die after only a few weeks (12).

Like other caspases, caspase-3 is initially synthesized as an inactive 32-kDa precursor (procaspase-3). During activation, procaspase-3 is converted to 20-kDa and 12-kDa

Correspondence to: Professor Jung Sup Lee, Department of Biotechnology, College of Natural Sciences, Chosun University, Gwangju 501-759, Korea
E-mail: jslee@mail.chosun.ac.kr

Abbreviations: CAPS, 3-(cyclohexylamino)-1-propanesulfonic acid; DTT, dithiothreitol; EDTA, ethylenediaminetetraacetic acid; EGTA, ethylene glycol-bis(2-aminoethyl ether)-N,N,N’,N’-tetraacetic acid; PARP, poly(ADP-ribose) polymerase; PBS, phosphate-buffered saline; PMSF, phenylmethanesulfonyl fluoride; pNA, p-nitroaniline; PVDF, polyvinylidene fluoride

Key words: Vibrio vulnificus, metalloprotease, procaspase-3
peptides via proteolysis at the cleavage site, Ile^{175}-Glu-Thr-Asp\_Ser^{190} (arrow indicates the cleavage site). The 20-kDa peptide undergoes further cleavage at the site, Glu^{208}-Ser-Met-Asp\_Ser^{209} (arrow indicates the cleavage site), generating the 17-kDa peptide (13). The active caspase-3 is a tetrameric protein consisting of two p17 and two p12 subunits. Recent reports have shown that procaspase-3 is catalytically active with lower \( k_\text{cat} \) and similar \( K_m \) compared to that of caspase-3 (14,15). In these reports, an uncleavable procaspase-3 mutant (D9A, D28A, or D175A) was used instead of natural pro-caspase-3. Using the same procaspase-3 [procaspase-3(D3A)], which is incapable of auto-activation when expressed in \( E. coli \), we were able to show that vEP could activate this caspase-3zymogen to generate mature caspase-3 which exhibited the same substrate and inhibitor specificity as wild-type caspase-3 in a cell-free system.

Materials and methods

**Materials.** Ac-DEVD-pNA and Ac-DEVD-CHO were purchased from Bachem. Mouse monoclonal anti-caspase-3 antibody was obtained from Sigma. Rabbit polyclonal anti-PARP antibody was obtained from Santa Cruz Biotechnology. Anti-rabbit and anti-mouse IgGs were obtained from Jackson Immunoresearch. Ni^{2+}-NTA was purchased from Amersham Biosciences. All other chemicals used were obtained from Sigma.

**Enzyme expression and purification.** The construction of procaspase-3(D3A) has been previously described (15). The aspartate at residues 9, 28, and 175 of the procaspase coding region was substituted with alanine to prevent auto-processing of thezymogen. The enzyme contained a His tag at the N- and C-termini. Procaspase-3(D3A) will henceforth be referred to as D3A for simplicity. D3A was expressed in the BL21(DE3) \( E. coli \) strain (Novagen), and protein induction was carried out with 0.2 mM IPTG at 25°C overnight. The enzyme was purified from a Ni-NTA column and finally exchanged into a buffer containing 25 mM Tris-HCl, pH 7.5, and 40% glycerol and stored in small aliquots at -70°C. vEP was purified from the culture supernatant of \( V. vulnificus \) ATCC 29307 as described previously (5).

**Activation of D3A by vEP.** Activation of D3A was performed by incubating thezymogen (0.16-0.58 mg/ml) with 0-8 \( \mu \text{g/ml} \) vEP in 25 mM Tris-HCl, pH 7.5 at room temperature. The reaction was terminated by addition of 1 mM 1,10-phenanthroline, a potent inhibitor of vEP (5) and then subjected to SDS-PAGE and Western blot analysis with anti-caspase-3 antibody or activity assay with the caspase-3-specific chromogenic substrate, Ac-DEVD-pNA in a 100 \( \mu \text{l} \) reaction volume consisting of 20 mM HEPES, pH 7.5, 1 mM EDTA, 1 mM EGTA, 20 mM NaCl, 10 mM DTT and 1 mM 1,10-phenanthroline. The release of pNA was continuously monitored by absorbance at 405 nm at 37°C using a 96-well plate reader (Molecular Devices). For assay with the inhibitor Ac-DEVD-CHO, 0.5 mg/ml zymogen was treated with 2 \( \mu \text{g/ml} \) vEP for 10 min at room temperature and then terminated by the addition of 1 mM 1,10-phenanthroline. Aliquots were extracted and assayed for caspase-3 activity as described above in the absence or presence of various concentrations of the inhibitor. 

**Cleavage of PARP.** To prepare the cell lysate, NIH3T3 cells from two 10 cm dishes were harvested, washed twice with PBS, re-suspended in 100 \( \mu \text{l} \) of lysis buffer (10 mM Tris-HCl, pH 7.5, 0.5% Triton X-100, 5 mM EDTA, 1 mM PMSF) and then incubated on ice on a shaker for 15 min. Ten \( \mu \text{l} \) of the lysate (150 \( \mu \text{g protein} \) ) was mixed with 10 \( \mu \text{l} \) of vEP-activated D3A (as described above) for different time periods at room temperature, and the reaction was terminated by the addition of 20 \( \mu \text{l} \) of 2X SDS-PAGE sample buffer. The samples were heated at 100°C for 1 min and then subjected to SDS-PAGE with subsequent Western blot analysis using an antibody against PARP.

**SDS-PAGE and Western blot analysis.** SDS-PAGE was performed according to the method of Laemmli (16). Samples to be analyzed were mixed with an equal volume of 2X SDS-PAGE sample buffer and heated at 100°C for 1-2 min, and then loaded onto 15% gel. After electrophoresis, protein bands were visualized by staining the gel with Coomassie blue. For Western blot analysis, protein samples were subjected to SDS-PAGE and then transferred to a PVDF membrane. The membrane was blocked at room temperature for 2 h with blocking solution (PBS containing 5% skim milk and 0.2% Tween-20) and then incubated for 1 h with either mouse monoclonal anti-caspase-3 antibody diluted 1:5000 or with rabbit polyclonal anti-PARP antibody diluted 1:500. The blot was washed and incubated with a 1:5000 dilution of peroxidase-conjugated anti-rabbit or anti-mouse IgG antibody for 1 h. The blot was washed, treated with ECL Western Blotting Detection reagents (Amersham Biosciences), and then exposed to Hyperfilm ECL (Amersham Biosciences).

**N-terminal sequencing.** Protein samples were subjected to electrophoresis on 15% SDS-PAGE gel. After electrophoresis, proteins were transferred to a PVDF membrane in 10 mM 3-(cyclohexylamino)-1-propanesulfonic acid (CAPS) buffer, pH 11.0 containing 10% methanol. The blot was stained with Coomassie blue, followed by destaining. Target bands were excised from the blot and subjected to N-terminal sequencing using an Applied Biosystem Precise sequencer (Applied Biosystem).

**Results**

**Activation of D3A.** The substitution of Ala for Asp in the cleavage sites of D3A prevented thezymogen activation since simple expression of wild-type caspase zymogens in \( E. coli \) usually results in their activation (17,18). The zymogen itself showed some activity, but the activity clearly varied with the concentrations of vEP used during the activation process, with a maximum increase of ~14-fold over thezymogen at 2 \( \mu \text{g/ml} \) vEP (Fig. 1A). vEP exhibited no detectable activity toward Ac-DEVD-pNA (data not shown). At higher concentrations of vEP, the activity decreased probably due to additional cleavage of the mature enzyme by vEP. At a fixed concentration of vEP (2 \( \mu \text{g/ml} \) ), the caspase-3 activity of activated D3A also exhibited an increase that paralleled the
concentrations of zymogen used in the activation reaction, but the activity appeared to reach a maximum at 0.5 mg/ml zymogen (Fig. 1B). At 0.5 mg/ml zymogen and 2 μg/ml vEP, the activity of caspase-3 increased with increasing activation time reaching a peak at 10 min and then started to decline with increasing activation time (Fig. 2A). The loss of activity correlated with the loss of a band that was approximately the same size as the p17 subunit of mature caspase-3 as revealed by Western blot analysis (Fig. 2B), and at the same time resulted in the accumulation of a band of ~14 kDa, which could be derived from the cleavage of the larger band via further proteolysis by vEP. This band did not belong to the p12 subunit since the anti-caspase-3 antibody reacted only with the p17 subunit. The band corresponding to 24-kDa appeared to be quite stable over a 40 min incubation period, whereas much of the zymogen had disappeared by this time point. The disappearance of the zymogen with time did not result in a corresponding increase for the mature enzyme, suggesting that cleavage of the zymogen to yield active caspase-3 was transient and the active caspase-3, once formed, was highly susceptible to further cleavage by vEP.

Since vEP-activated D3A probably consisted of more than one species with activity toward DEVD, no kinetic properties were determined for the mature enzyme. However, the $K_m$ for Ac-DEVD-pNA determined from the mixed species was 30.5 μM compared with the values of 2.0-16.8 μM reported for the recombinant wild-type mature caspase-3 (14,15,19). The $K_m$ for D3A (12.8 μM) was similar to the reported value of 13.2 μM (15). The inhibitory effect of Ac-DEVD-CHO on vEP-activated D3A was also confirmed, with an apparent $K_i$ of 18.3 nM at 0.2 mM substrate.

Cleavage fragments of D3A. From the cleavage of D3A, it seemed clear that vEP cleaved at multiple sites within the zymogen and the rate of cleavage for each site might differ. The purified zymogen consisted mainly of a 32-kDa protein on SDS-PAGE with minor contaminating bands of ~20 kDa and an even fainter band near the 17-kDa regions (Fig. 3A). Both bands could be a product of proteolysis by E. coli proteases during expression and/or purification. Cleavage of the zymogen by vEP yielded various fragments, with a 24-kDa fragment being the most abundant. Various fragments of 18-12 kDa that were absent in the case of non-cleaved zymogen were present at a low amount. These fragments covered the size of both the large and small subunits of the mature caspase-3. With the non-activated D3A, Western blotting detected a band of ~20 kDa in addition to the zymogen (Fig. 3B); therefore, this fragment appeared to be caspase-3 protein derived probably from the cleavage of the zymogen during expression or purification. Notably, at longer activation times, the cleavage of the zymogen may proceed with higher non-specificity without yielding active mature caspase-3. Two fragments (F1 and F2, Fig. 3A) derived from the cleavage of D3A by vEP with sizes similar to those of the p17 had the same N-terminal sequence of Ile20-Ile-His-Gly-Ser24. Therefore Lys19-Ile20 was one of the sites within D3A cleaved by vEP, and since both F1 and F2 had the same N-terminal

---

**Figure 1.** Effect of vEP and zymogen concentration on the activation of D3A. (A) D3A (0.5 mg/ml) was activated with different concentrations of vEP at room temperature for 10 min. (B) Different concentrations of D3A were activated with 2 μg/ml vEP at room temperature for 10 min. Caspase-3 activity was assayed with Ac-DEVD-pNA as described in Materials and methods. Data are the means of two experiments performed in duplicate.

**Figure 2.** Time-dependent activation of D3A by vEP. D3A (0.5 mg/ml) was activated by vEP (2 μg/ml) at room temperature for different time intervals and then subjected to activity assay toward Ac-DEVD-pNA (A) and Western blot analysis (B). The arrow in B indicates the position of the p17 subunit of mature caspase-3.
Discussion

Activation of procaspase-3 by other proteases has previously been shown to be catalytically competent but the catalytic efficiency is ~130- to 200-fold lower than its mature form despite having similar $K_m$ values (14,15). Although the crystal structure of caspase-3 has been determined (26,27), no crystal structure for procaspase-3 is yet available. However, the crystal structure obtained for procaspase-7 (28) shows that the presence of cleavage at the junction between the large and small subunits is necessary to form the correct active site for binding to substrate. It appears that the retention of a correct active conformation by the processed enzyme is crucial for maintaining substrate specificity even though the zymogen might not be cleaved at the precise cleavage site. This is a significant event that emphasizes the versatility of a protease which has broad specificity, whereby random cleavage of a zymogen leads to an active enzyme that further amplifies a particular reaction cascade.

Here we have shown the activation of a procaspase-3 mutant that is incapable of self-activating due to the absence of the critical aspartate residues in the cleavage sites. The repertoire of proteolysis and showed that the cleavage of the zymogen occurred within the linker region between the large and small subunits converting a single-chain zymogen to two-chain forms. vEP is a broad specificity protease which can cleave at the carboxyl side of Asp, Thr and Tyr, with the surrounding amino acid sequence playing an important part in the cleavage sites (5). A particular feature among the various known cleavage sites of vEP is the presence of a hydrophobic amino acid which is immediately adjacent to the point of cleavage. Although we have no data to show the structural change (if any) that might possibly be incurred by the Ala substitutions in D3A, we do not think that the ability of vEP to cleave D3A and activate it was afforded by the structural change produced by these substitutions. Instead, the formation of active caspase-3 from D3A was due more to the cleavage at sites that are near Asp28 and Asp175. Unfortunately, sequencing data were not successfully obtained to reveal the cleavage at the p17 and p12 junction of D3A.

Cleavage of PARP. PARP is an important 116-kDa zinc-finger protein for DNA repair, transcription and chromosomal stability (20). Caspase-3 is known to cleave PARP during apoptosis to generate 85- and 24-kDa fragments. Addition of D3A to a cell-free extract prepared from NIH3T3 cells resulted in the cleavage of PARP (Fig. 4). Since D3A has some activity toward the peptide substrate, the cleavage of PARP in cell-free extract by D3A was not unexpected. However, if D3A was first treated with vEP and then added to the cell-free extract, more cleavage of PARP was observed for the same incubation time (Fig. 4). This increase in the amount of PARP cleavage confirms that D3A activated by vEP also retained the same substrate specificity as mature caspase-3. Cleavage of PARP has become a standard way to identify the presence of caspase-3 and its activity. The mature caspase-3 resulted from the cleavage of D3A by vEP, yet still preserved the same specificity despite being processed at a different cleavage site.

Figure 4. Cleavage of PARP by vEP-activated D3A. D3A (0.5 mg/ml) was incubated without or with vEP (2 μg/ml) at room temperature for 10 min, and the reaction was terminated with 1,10-phenanthroline (1 mM). Ten μl of cell lysate (150 μg protein) was incubated with 10 μl of non-activated D3A for 30 min or with 10 μl of vEP-activated D3A for 5, 20, 30 and 60 min at room temperature. The control sample contained only cell lysate incubated for 30 min at room temperature.

Sequence but had different molecular masses, they could have different C-terminal sequences, suggesting a cleavage occurring probably near the residue corresponding to Asp175. Although sequencing data were not successfully obtained to reveal the cleavage at the p17 and p12 junction of D3A.
activation of procaspase-3 exhibited a similar feature with that of prothrombin activation (5) in that the mature enzyme was further subjected to rapid degradation by vEP. These activation events suggest that zymogen activation by vEP in both procaspase-3 and prothrombin may not be a specific reaction, but could be a by-product arising from the result of cleavage of the zymogen at sites close to those recognized by their natural activators. This property of vEP is largely attributed to its broad specificity. Although the activation may not be a significant event in the case of wild-type procaspase-3, which is capable of auto-activation, this finding shows that the cleavage site recognized by vEP does not converge on the specificity associated with the critical aspartate residues essential for auto-activation.

There is currently no evidence to suggest that activation of procaspase-3 by vEP occurs inside the cell. In order for vEP to have any activity toward procaspase-3 it needs to enter the cell. There is yet no direct evidence to show that entrance of vEP into the cell actually takes place. However, confocal microscopy images of NIH3T3 cells treated with 5 μg/ml fluorescein-labeled recombinant vEP indicates some penetration of vEP into the cells after a treatment time of 1 h (data not shown). The activation of procaspase-3 by vEP as described here emphasizes the ability of vEP (as a protease) to activate different types of zymogens rather than being a direct link to apoptosis, and certainly by no means suggests that vEP activates procaspase-3 and causes cell death. Furthermore, the activation of procaspase-3 is a response to the cell death pathway and not a cause of cell death. However, the ability of vEP to activate certain proteases through cleavage of the respective zymogens makes it a formidable enzyme to be present in the host body during V. vulnificus infection. The in vitro activation described here extends the capability of the V. vulnificus extracellular metalloprotease with respect to zymogen activation, and also opens up new ground for future investigation into the function of this broad specificity protease.

Acknowledgements

This study was supported by the ERC program of MOST/ KOSEP through Research Center for Proteoeneous Materials (RCPM) of Chosun University, and by research funds from Chosun University, 2003.

References