ANTIPLASMIN-CLEAVING ENZYME IS A SOLUBLE FORM OF
FIBROBLAST ACTIVATION PROTEIN

Kyung N. Lee, Kenneth W. Jackson, Victoria J. Christiansen, Chung S. Lee, Jin-Geun Chun and Patrick A. McKee

William K. Warren Medical Research Center and Department of Medicine, University of Oklahoma Health Sciences Center, Oklahoma City, Oklahoma 73190, USA

Running title: soluble fibroblast activation protein

Scientific section: Hemostasis, Thrombosis, and Vascular Biology

Word count: Abstract, 199 words; total text, 4705

Correspondence:

Kyung N. Lee, PhD
W.K. Warren Medical Research Center
P.O. Box 26901, BSEB-306
Oklahoma City, Oklahoma 73190
Fax: (405) 271-3191
e-mail: kyung-lee@ouhsc.edu

Financial support: William K. Warren Medical Research Center and NIH grant HL072995

Declaration of commercial interest: None

Copyright © 2005 American Society of Hematology
ABSTRACT

Circulating antiplasmin-cleaving enzyme (APCE) has a role in fibrinolysis and appears structurally similar to fibroblast activation protein (FAP), a cell-surface proteinase that promotes invasiveness of certain epithelial cancers. To explore this potential relationship, we performed comparative structure/function analyses of the two enzymes. APCE from human plasma and recombinant FAP (rFAP) exhibited identical pH optima of 7.5, extinction coefficients ($\varepsilon_{280\text{nm}}^{1\%}$) of 20.2 and 20.5, common sequences of tryptic peptides, and cross-reactivity with FAP antibody. APCE and rFAP are homodimers with monomeric subunits of 97 and 93 kDa. Only homodimers appear to have enzymatic activity, with essentially identical kinetics toward Met-\(\alpha_2\)-antiplasmin (Met-\(\alpha_2\)AP) and peptide substrates. APCE and rFAP cleave both Pro3-Leu4 and Pro12-Asn13 bonds of Met-\(\alpha_2\)AP, but relative $k_{cat}/K_m$ values for Pro12-Asn13 are ~16-fold higher than for Pro3-Leu4. APCE and rFAP demonstrate higher $k_{cat}/K_m$ values toward a peptide modeled on P4-P4' sequence surrounding the Pro12-Asn13 primary cleavage site than for Z-Gly-Pro-AMC and Ala-Pro-AFC substrates. These data support APCE as a soluble derivative of FAP and Met-\(\alpha_2\)AP as its physiologic substrate. Conversion of Met-\(\alpha_2\)AP by membrane or soluble FAP to the more easily fibrin-incorporable form, Asn-\(\alpha_2\)AP, may increase plasmin inhibition within fibrin surrounding certain neoplasms and impact growth and therapeutic susceptibility.
INTRODUCTION

Human $\alpha_2$-antiplasmin ($\alpha_2$AP) becomes rapidly crosslinked to fibrin by activated clotting factor XIII (FXIIIa) and protects fibrin by rapidly inhibiting plasmin within the fibrin clot\textsuperscript{1,2}. Fibrin containing reduced $\alpha_2$AP activity is susceptible to fibrinolysis, while fibrin containing a normal amount of $\alpha_2$AP is strikingly resistant to lysis by plasmin\textsuperscript{2-6}. Human $\alpha_2$AP was thought to have an N-terminal Asn and a 39-residue signal peptide until later work demonstrated that $\alpha_2$AP was secreted from liver cells as a 464-residue protein with an additional 12 residues at the N-terminus (Met-$\alpha_2$AP), making the signal peptide 27 residues\textsuperscript{7-10}. The N-terminal 12-residue peptide of Met-$\alpha_2$AP was reported to be cleaved in the circulation by an unknown proteinase\textsuperscript{10}. In normal human plasma, the ratio of Met-$\alpha_2$AP to Asn-$\alpha_2$AP was estimated as $\sim$30:70\textsuperscript{9,10}. We showed that during clot formation Asn-$\alpha_2$AP becomes crosslinked to fibrin by FXIIIa $\sim$13X faster than Met-$\alpha_2$AP, and as a consequence, clot lysis rates slow in direct proportion to the ratio of Asn-$\alpha_2$AP/Met-$\alpha_2$AP in human plasma\textsuperscript{5}.

We purified the plasma proteinase that converts Met-$\alpha_2$AP to Asn-$\alpha_2$AP, and termed it antiplasmin-cleaving enzyme (APCE)\textsuperscript{5}. Its N-terminal amino acid sequence corresponded to residues 24-38 of fibroblast activation protein (FAP), also known as seprase, a type II integral membrane glycoprotein, which is predicted to have its first six N-terminal residues within the fibroblast cytoplasm, followed by a 20-residue transmembrane domain, and then a 734-residue extracellular C-terminal catalytic domain\textsuperscript{5,11,12}. FAP is a proline-specific enzyme that exhibits gelatinase activity as well as dipeptidyl peptidase activity\textsuperscript{13}. FAP and
dipeptidyl peptidase IV (DPP IV), which is another post-prolyl peptidase, are members of
the S9b peptidase family in which the order of Ser/Asp/His in the active-site is different
from that in the chymotrypsin-like serine proteinase family\(^\text{14}\). FAP is reported to be
expressed by fibroblasts of reactive stroma of epithelial-derived cancers, but not by
counterpart normal tissues\(^\text{13,15,16}\). Subsequent studies showed that FAP is also expressed by
several types of neoplastic parenchymal cancer cells and also by endothelial cells of newly
formed vessels within certain neoplasms\(^\text{17-21}\). In a murine breast cancer model, cells
transfected to express FAP form more rapidly growing tumors when compared to control-
transfectants\(^\text{22}\). While these findings suggest that FAP may be a potential target for
diagnostic and therapeutic approaches in certain cancers\(^\text{23}\), its biologic function remains
unknown. To date no certain physiologic substrate for FAP has been identified.

Schmidt et al.\(^\text{24}\) indicated that no detectable soluble FAP, or a derivative of such,
could be found in human blood; however, our finding of APCE in human plasma\(^\text{5}\) is
inconsistent with this. Interestingly, Collins et al.\(^\text{25}\) have found FAP-like activity in
bovine serum which is in accord with our report of APCE in human plasma. APCE may
be a soluble, circulating derivative of FAP, resulting from cleavage of the Cys23-Ile24
bond in the transmembrane or extracellular domain of FAP, and if so, its presence in
plasma could impact the use of therapeutic agents designed to target FAP at the tissue
level\(^\text{23,24}\). We now report direct evidence that the two proteins are essentially identical.
We obtained sufficient quantities of APCE from human plasma and soluble human rFAP
from a yeast expression system to allow determinations and comparisons of previously
unreported molecular structure/function details for the two proteinases.
MATERIALS AND METHODS

Materials

Human plasma Met-α2AP was isolated from a mixture of Met-α2AP and Asn-α2AP using Met-α2AP-specific antibody as previously described\(^5\); a percent extinction coefficient of α2AP at 280 nm of 6.70 and a molecular weight of 70,000 were used to determine molar concentration\(^26\). Hybridoma cells secreting the F19 monoclonal antibody to FAP were purchased from American Type Culture Collection (Manassas, VA) and grown in serum-free media; the F19 antibody was purified from hybridoma cell culture media using MEP-Hypercel chromatography (Pall Corp., East Hills, NY). An EST cDNA clone (GenBank accession #: AV698972) was obtained from Dr. Ze-Guan Han, through the Chinese National Human Genome Center at Shanghai, China. Upon determining its sequence, using M13-forward and M13-reverse primers in conjunction with four specific primers, the EST clone was shown to contain full-length FAP cDNA (GenBank accession #: NM_004460). Other reagents included: Pichia pastoris expression vector pPICZαA and Pichia pastoris X-33 (Invitrogen, San Diego, CA), TPCK-treated trypsin (Sigma, St. Louis, MO), and Z-Gly-Pro-AMC and Ala-Pro-AFC (Bachem, Torrance, CA). AMC (7-amido-4-methylcoumarin) and AFC (7-amino-4-trifluoromethylcoumarin) were obtained from MP Biochemicals (Aurora, OH). A fluorescence resonance energy transfer (FRET) peptide, Arg-Lys(DABCYL)-Thr-Ser-Gly-Pro-Asn-Gln-Glu-Gln-Glu(EDANS)-Arg was synthesized in-house as previously described\(^5\). Asn-Gln-Glu-Gln-Glu(EDANS)-Arg was
prepared as follows: (a) the Pro-Asn bond of the FRET peptide was cleaved by APCE; and (b) the fluorescent fragment, Asn-Gln-Glu-Gln-Glu(EDANS)-Arg, was purified by HPLC and confirmed by mass and amino acid sequence analyses.

**Isolation of APCE**

APCE was partially purified by ammonium sulfate precipitation and hydrophobic interaction chromatography as previously described\(^5\). The activity peak was applied to an immunoaffinity column made with F19 monoclonal antibody and eluted with 5 M LiCl. Based on Z-Gly-Pro-AMC cleaving activity, the active fractions were pooled and brought to 6% ammonium sulfate, applied to a Phenyl-5PW column (Tosohaas, Montgomeryville, PA) equilibrated in 6% ammonium sulfate, and eluted with 12.5 mM sodium phosphate, pH 7.5. Before storing at \(-80^\circ\)C, glycerol was added to purified APCE to give a final concentration of 20%.

**Construction of rFAP Expression Vector and Transformation of *Pichia pastoris***

Human FAP cDNA with XhoI and NotI sites at the 5’ and 3’ ends, respectively, was amplified as described previously\(^2\), using two oligonucleotide primers (5’-ATCTCTCGAGAAAAGAATTGTCTTACGCCCTTCAAG for the XhoI site and 5’-CAGCCCGCGGCCGCTTAGTCTGACAAAGAAAC for the NotI site) and the template full length FAP cDNA. Using the introduced restriction sites, the amplified DNA was ligated into the *Pichia pastoris* vector pPICZaA for secretional expression. The
expression vector was digested with SacI and then transformed into *Pichia pastoris* X-33 by the EasyComp transformation procedure (Invitrogen instruction manual). Transformants were selected on YPDS agar plates containing 100 µg/ml Zeocin.

**Expression and Purification of Recombinant FAP**

Zeocin resistsants (100 colonies) were screened by determining rFAP activity in expression culture medium. Five ml cultures were grown overnight in BMGY medium (buffered glycerol complex medium) at 29°C and cells were harvested by centrifugation at 2000 x g. Cell pellets were resuspended with 5 ml BMMY medium (BMGY with 5 ml of methanol per liter in place of glycerol), and then induction was continued with fresh methanol (5 ml per liter) added every 24 hours for 3 days. Z-Gly-Pro-AMC cleaving activity in the culture medium was monitored.

The yeast cells displaying highest activity were grown to \(A_{600}\) of 2-3 in 200 ml of BMGY medium in a two-liter baffled flask at 29°C. The cells were harvested and resuspended in 200 ml of BMMY medium and cultured at 29 °C for 5 days and then centrifuged at 12,800 x g for 40 min at 4°C to pellet the yeast cells. The supernatant was filtered through 0.1 µm Millipore membranes to remove solid residuals, and purified by the same procedure as described for APCE.

**Size Exclusion Chromatography**
Native molecular mass for rFAP or APCE was determined at room temperature on a Superdex-200 PC 3.2/30 column (Amersham Bioscience, Piscataway, NJ), using 50 mM sodium phosphate-150 mM NaCl, pH 7.4, at a flow rate of 45 µl/min. Commercially obtained protein standards (Bio-Rad, Hercules CA) were used for calibration.

**Extinction Coefficient**

APCE or rFAP concentration was determined by amino acid analysis and absorbance at 280 nm measured in 12.5 mM sodium phosphate buffer, pH 7.5. Percent extinction coefficient was expressed as \( (\varepsilon_{\text{molar}})^{10} = (\varepsilon_{\text{percent}}) \times \text{MW} \), where \( \varepsilon_{\text{molar}} (A_{280} \text{ nm/M}) \) and \( \varepsilon_{\text{percent}} (\varepsilon_{280nm}^{\%}) \) are molar and percent extinction coefficients and MW is the subunit molecular weight of 97,000 for APCE, or 93,000 for rFAP. Percent extinction coefficients at 280 nm were calculated as 20.2 and 20.5 for APCE and rFAP, respectively.

**Enzyme Assays**

APCE or rFAP was incubated in 25 mM sodium phosphate buffer, pH 7.5, containing 1.0 mM EDTA and 4% methanol in a total volume of 200 µl for 20 min at 22°C, using as substrates Z-Gly-Pro-AMC (25-400 µM); Ala-Pro-AFC (50-800 µM); or a FRET peptide, Arg-Lys(DABCYL)-Thr-Ser-Gly-Pro-Asn-Gln-Glu-Gln-Glu(EDANS)-Arg (5-80 µM) that was designed to contain the APCE-sensitive Pro12-Asn13 bond within the Thr9-Gln16 sequence of Met-α2AP. Fluorescence was monitored with time at excitation and emission wavelengths of 360 and 460 nm for Z-Gly-Pro-AMC and the FRET peptide,
and 360 and 530 nm for Ala-Pro-AFC, using a BIO-TEK FL600 fluorescence plate reader (Winooski, VT). To obtain standard curves, dilutions of AMC, AFC, and Asn-Gln-Glu-Gln-Glu(EDANS)-Arg were prepared in the same assay buffer and corresponding fluorescence was measured. All enzyme kinetic parameters were computed by fitting data to the Michaelis-Menten equation27, employing the commercially available program EZ-FIT (Perrella Scientific, Amherst, NH).

Zymography to measure gelatinase activity of APCE or rFAP was performed using gelatin zymograms, and renaturing and developing solutions (Invitrogen, San Diego, CA). Sample loading, electrophoresis and incubation of each gel in the renaturing and developing buffers were carried out according to the accompanying protocol. After overnight incubation in the developing buffer at 37°C, each gel was stained with Coomassie brilliant blue G-250 in order to see the area of gelatin digestion as a clear band and rFAP or APCE protein as a darker band in the background.

**Liquid chromatography-mass spectrometry (LC/MS) Analysis of Met-α2AP Digested by rFAP or APCE**

Selected concentrations of Met-α2AP were incubated with 0.85 µg rFAP or APCE in 100 µl of 25 mM sodium phosphate-1.0 mM EDTA buffer, pH 7.5. After incubating for five hours, an internal standard peptide (FRQLTSDKPNQEYQV, 0.113 µg, 144 pmol) was dissolved in the reaction mixture and cold acetone (4x sample volume) was added to precipitate proteins. The sample was incubated at -80°C for 60 min and centrifuged for
10 min at 16,000 x g at 4°C to remove precipitated proteins. The supernatant was removed and dried by vacuum centrifugation and dissolved in 5% acetic acid.

LC/MS analysis of samples was performed on a Magic 2002 HPLC system (Michrom BioResources, Auburn, CA) equipped with a Jupiter Proteo reverse-phase column (1.0 mm X 150 mm, Phenomenex, Torrance, CA). The column was equilibrated with 5% acetonitrile/water containing 0.1% TFA. Upon sample injection, the solvent composition was increased to 10% acetonitrile and a linear gradient was applied to 40% acetonitrile over 60 min. Peptides were detected at 215 nm wavelength. The HPLC was connected to an electrospray mass spectrometer (QSTAR, Applied Biosystems, Foster City, CA) equipped with an ion spray source operated in the positive ion mode. Scanning was performed over an m/z range of 350 to 1800 amu. Both the internal standard peptide and peptide products of digestion were located by reconstructed ion current analysis of data for each peptide over a 1.5 amu window for both the singly and doubly charged forms of each peptide, based upon the peptide’s predicted monoisotopic molecular mass. Quantitation was performed by summing all detected isotopes of each peptide.

RESULTS

Expression and Purification of rFAP

We have suggested that APCE may be the N-terminally-cleaved form of FAP, partially because the N-terminal 15 amino acids of APCE are identical to residues 24-38 of FAP5.
Therefore the 23 residues were removed by amplifying the coding region of full-length FAP cDNA between amino acid residues 24 and 760. Using the amplified DNA and *Pichia pastoris* vector pPICZαA, the expression vector was constructed for secretion of rFAP into media. Figure 1A shows that purified rFAP or APCE gives a single band of 93 kDa or 97 kDa, respectively, on reduced SDS-PAGE. Identical patterns for both enzymes were also observed in the absence of reducing agent. Figure 1B shows that the F19 monoclonal antibody to FAP13 cross-reacts with both APCE and rFAP.

Our purification process for rFAP produced ~42 µg of pure protein from one liter of culture media to give a 17% yield. Protein concentration of rFAP or APCE was calculated using percent extinction coefficients of 20.5 or 20.2, respectively, at 280 nm as determined by us. Figure 2 shows the effects of pH on the specific activity of rFAP and APCE. At this time we have no explanation for the wide range of enzyme activity over the pH spectrum examined these studies. For each enzyme, the pH optimum was 7.5 for the cleavage of the Pro-Asn bond in the FRET peptide substrate, Arg-Lys(DABCYL)-Thr-Ser-Gly-Pro-Asn-Gln-Glu-Gln-Glu(EDANS)-Arg. Both enzymes also showed the same pH optima with the substrate Z-Gly-Pro-AMC (data not shown).

**Sequence Analysis of rFAP and APCE**

The N-terminal amino acid sequences of rFAP and APCE were determined by using an automated protein sequencer. As shown in Figure 3, the N-terminal sequence of purified rFAP (EENTMRALTL, FAP residues 35-44) is different from the N-terminal sequence of
APCE (IVLRPSPVHNSEENT, FAP residues 24-38). Although the N-terminal amino acid of rFAP was expected to be Ile24, it in fact was Glu35. Various constructs were designed with a His(6)- or Glu-Ala-His(6)-tag attached N-terminally, or not at all, and every instance resulted in the expression of a stable, active enzyme, but with the same dominant N-terminal sequence. A dominant amino acid sequence of 10 amino acid residues (EENTMRALTL, residues 35-44) and a minor sequence containing residues 29-37 (SRVHNXEEN) were obtained in ~80:20 ratio, indicating the homogeneity of purified rFAP and suggesting that 11 N-terminal residues had been deleted or cleaved during culture or enzyme preparation. As can be seen in Figure 1A, molecular size determination of rFAP (93 kDa) and APCE (97 kDa) by SDS-PAGE support rFAP as being smaller, probably due to glycosylation differences between human and yeast-expressed proteins and the deletion of 11 N-terminal residues. To compare the internal sequences of rFAP and APCE, each protein was subjected to SDS-PAGE, reduced, alkylated and digested with trypsin. Each trypsin digest was analyzed by liquid chromatography-mass spectrometry-mass spectrometry (LC/MS/MS) to obtain molecular weights and MS/MS fragment ion spectra to utilize with the MASCOT MS/MS ion search engine to query the NCBI comprehensive non-identical protein database (http://www.matrixscience.com/cgi/search_form.pl?FORMVER=2&SEARCH=MIS).

As shown in Figure 3, amino acid sequences of peptides obtained from trypsin digests of rFAP and APCE are identical to the sequence derived from FAP cDNA sequence.

**Homodimer Formation Required for APCE or rFAP Activity**
FAP appears to be composed of two 97 kDa identical subunits in a homodimeric structure, suggested as necessary for both dipeptidyl peptidase and gelatinase activities\(^{11,13}\). Definitive data, however, showing that dimerization is required for activity are lacking. As shown in Figure 4, we examined this directly using zymography and size exclusion chromatography. When boiled in SDS, APCE and rFAP showed 97 and 93 kDa bands, respectively, and given complete denaturation, no gelatinase activity was observed (Figure 4A). However, non-boiled samples, from which SDS was removed by incubating the zymogram-gelatin gel in a renaturing buffer after electrophoresis (Figure 4A), clearly showed both 170 kDa and 180 kDa gelatinase activities for APCE and rFAP, respectively. Also noted were remaining inactive 97 kDa and 93 kDa monomer bands for both enzymes, which despite renaturation efforts, lacked gelatinase activity. The molecular size of APCE was consistent with prior reports of 175 kDa for enzymatically active FAP homodimer and 95 kDa for the inactive monomer\(^{11,13,29}\). Without SDS in the samples, the gelatinase activity band of rFAP migrated slightly slower than that of APCE, and both band intensities increased ~20-fold compared with non-boiled SDS-containing sample bands; monomeric bands were not apparent (Figure 4B). Although by SDS-gel electrophoreses the monomer band of APCE migrates slower than that of rFAP (Figure 4A), the dimeric form of APCE migrates faster than that of rFAP in the non-SDS gel (Figure 4B). We speculate that this may be due to the negative surface charge of APCE being higher than for rFAP due to differences in glycosylation. Similar size discrepancy has been reported in native and recombinant DPP IV proteins, which are in the same S9b peptidase family as FAP\(^{30}\). Using size exclusion chromatography, we also assessed whether APCE or rFAP remains as an enzymatically active dimer in phosphate-buffered
saline, pH 7.4. As shown in Figure 4C, APCE is present as a 175 kDa protein peak, while rFAP gives a 153 kDa peak. No other peaks are present, particularly in the monomeric 95 kDa range. The rFAP peak showed a broader mass distribution than APCE that is probably due to heterogeneous glycosylation by the yeast-expression system. The broader peak observed for rFAP on size exclusion chromatography is also consistent with the broader band for rFAP by SDS-PAGE when compared to APCE (Figure 1). Both the 175 kDa and 153 kDa peak fractions have gelatinase and Ala-Pro-AFC cleaving activities. These data from zymography and size exclusion chromatography suggest that homodimers are required for enzymatic activity and monomers lack such function.

**Substrate Specificity of rFAP and APCE**

To determine if rFAP has the same specificity as APCE, enzymatic activities of rFAP and APCE were compared using selected synthetic peptides as substrates (Table 1). These analyses clearly demonstrated that rFAP possesses the same, or closely similar, kinetic parameters as APCE. For Z-Gly-Pro-AMC substrate, human rFAP and APCE have $K_m$ values of 0.101 mM and 0.124 mM. These values are ~two-fold smaller when compared to a $K_m$ of 0.270 mM for bovine FAP toward Z-Gly-Pro-AMC. Using Z-Gly-Pro-AMC, we determined $k_{cat}$ values for human rFAP and APCE to be 30.5 min$^{-1}$ and 32.2 min$^{-1}$; $k_{cat}$ for bovine FAP has not been reported. For Ala-Pro-AFC substrate, human rFAP and APCE have $K_m$ and $k_{cat}$ values of 0.323 mM and 64.7 min$^{-1}$, and 0.272 mM and 59.7 min$^{-1}$, respectively, in contrast to 0.20 mM and 120 min$^{-1}$, which Sun et al. reported for full-
length human rFAP. The modest discrepancies between our data and theirs are likely due to different buffer pHs used in the two sets of experiments. Also, compared to their full-length rFAP, our rFAP was N-terminally shorter by 34 amino acids, thus lacking the cytosolic and transmembrane domains, which could have diminished solubility and affected kinetic parameters. The kinetic efficiencies \(k_{cat}/K_m\) values of rFAP and APCE for cleavage of Z-Gly-Pro-AMC, Ala-Pro-AFC, and FRET peptide are essentially the same towards each substrate. However, the \(k_{cat}/K_m\) value for cleavage of FRET peptide, remembering that it is modeled on the sequence surrounding the APCE-cleavage site (Pro12-Asn13) of Met-\(\alpha_2\)AP, is ~8-fold higher than for Z-Gly-Pro-AMC and ~12-fold higher than for Ala-Pro-AFC.

**Met-\(\alpha_2\)AP Cleavage by rFAP**

Because our kinetic studies and FRET peptide data suggested that rFAP, just as APCE\(^5\), may specifically cleave Met-\(\alpha_2\)AP at Pro12-Asn13, rFAP was incubated with Met-\(\alpha_2\)AP and the mixture examined at selected times by Western blot analysis. As demonstrated in Figure 5A, the intensity of the Met-\(\alpha_2\)AP band decreased with time when immunoblotted with a specific antibody to the 12-residue N-terminal peptide of Met-\(\alpha_2\)AP\(^5\). Figure 5B, however, shows that a blot immunostained with an antibody that detects both N-terminal forms of \(\alpha_2\)AP did not change through the entire time course, and bands suggesting fragmentation were never generated. N-terminal analyses of the 0 h sample showed only the Met-\(\alpha_2\)AP sequence, MEPLGRQLTSGP, while the 7 h sample had only the Asn-
α₂AP N-terminal sequence, NQEQVSPLTLLK. These data indicate that rFAP, like APCE, specifically cleaves the Pro12-Asn13 bond of Met-α₂AP.

Given cleavage of the Pro12-Asn13 bond in Met-α₂AP, we questioned whether the Pro3-Leu4 bond might also be cleaved during incubation with rFAP. This prompted the use of LC/MS to analyze peptides generated during incubation of rFAP with Met-α₂AP. Figure 6A (internal standard peptide(IS) plus Met-α₂AP incubated without rFAP) and Figure 6B (IS plus Met-α₂AP incubated with rFAP) each depicts LC/MS total ion current analysis of the digest between retention times of 9 and 23 min. In Figure 6A, IS shows the expected peak. In Figure 6B, when Met-α₂AP was incubated with rFAP, the control IS and the N-terminal 12 amino acid peptide (M1-P12), MEPLGRQLTSGP, and its oxidized form, M1(o)-P12, were observed. As shown in Figure 6C, a nine-residue L4-P12 peptide (LGRQLTSGP) that is clearly a cleavage product of the N-terminal 12 amino acids was located by reconstructed ion current analysis over a 1.5-atomic mass unit window for the doubly charged form of the peptide based on its predicted monoisotopic molecular mass. The amount of M1-P12, M1(o)-P12 and L4-P12 peptides were determined by comparison of each peak area with the IS peak area, 144 pmol of IS having been added to each sample. As shown in Figure 7, cleavage rates of the Pro3-Leu4 bond and the Pro12-Asn13 bond in Met-α₂AP by rFAP or APCE were determined by the amount of generated L4-P12 peptide, and the sum of L4-P12, M1-P12, and M1(o)-P12 peptides, assuming that the Pro12-Asn13 bond is cleaved before Pro3-Leu4. Cleavage rates of Pro3-Leu4 and Pro12-Asn13 increased linearly with substrate concentrations up to 8 µM. Under competitive conditions, the ratio of hydrolytic rates of
each cleavage site can be described as a relative $k_{cat}/K_m$ value.$^{33}$ Based on cleavage rates of Pro3-Leu4 and Pro12-Asn13 in the presence of 2, 4 or 8 µM Met-$\alpha_2$AP, the relative $k_{cat}/K_m$ values for Pro3-Leu4 and for Pro12-Asn13 were determined for each substrate concentration and averaged for each of the bond cleavages to yield: for rFAP, 1 and 16.6 ± 1.2 (mean ± SD), while for APCE, the relative $k_{cat}/K_m$ for Pro3-Leu4 and for Pro12-Asn13 were 1 and 16.3 ± 3.2 (mean ± SD).

**DISCUSSION**

The major goals of this study were (i) to determine if APCE is a soluble form of FAP and (ii) to systematically characterize the molecular and kinetic properties of both enzymes. Despite the identity of limited amino acid sequences we previously reported for APCE and FAP$^5$, concern remained about differences between the two; namely, FAP has a 23-residue N-terminal extension with Met at the N-terminus, while APCE has Ile as its N-terminus (corresponding to Ile24 in FAP), cleaves Met-$\alpha_2$AP and circulates in blood, unlike what has been reported for FAP$^{24}$. Our initial approach was to develop methods for obtaining sufficient amounts of apparently homogenous APCE from human plasma for characterization. Purification yields of APCE from human plasma were improved four-fold, using a modification of our previous method$^5$. To date efforts at structure/function analyses have been limited by low yields of FAP from baculovirus or mammalian cell expression systems and, in addition, by uncertain purity. We chose the *Pichia pastoris* expression system and found that rFAP was secreted into culture media from which we could obtain ~42 µg of pure rFAP per liter. Pure rFAP, possessing only amino acid
sequence deduced from FAP cDNA, was found to be a soluble, stable and active enzyme. Our yield of purified protein was ~5-fold greater than that reported for insect cells with recombinant baculovirus expressing the 760-residue full-length protein\textsuperscript{32}. Chinese hamster ovary cells expressing full-length FAP provided only trace amounts of recombinant protein\textsuperscript{34}.

Based on comparison of APCE and rFAP amino acid sequences to the published sequence for FAP cDNA, we found that APCE and rFAP each contained only the extracellular C-terminal catalytic domain and no N-terminal cytoplasmic or transmembrane domains; although rFAP is N-terminally shorter by 11 amino acids, it has the same specific activity as APCE. In-gel tryptic digestion and then LC/MS/MS analyses were used to characterize peptides from APCE and rFAP. The analyses identified 30 peptides from APCE and 21 from rFAP, equivalent to 48% and 30%, respectively, of the amino acid sequence predicted from FAP cDNA (Figure 3). APCE and rFAP were both purified using the F19 monoclonal antibody specific for FAP\textsuperscript{35}, and both enzymes were recognized by the same antibody in Western blot analyses (Figure 1B). Hence, data comparing amino acid sequences and antibody cross-reactivities support APCE being a soluble derivative of membrane-bound FAP.

We examined whether a homodimeric structure for APCE or rFAP is required for gelatinase and dipeptidyl peptidase activities, using gelatin zymography and size exclusion chromatography. The homodimeric form of rFAP or APCE cleaves gelatin, Ala-Pro-AFC and Z-Gly-Pro-AMC, whereas monomers of either had no such activities. While
previous data suggested that homodimer formation was needed for enzymatic function\textsuperscript{11,13,29}, this could only be inferred from zymography that could not provide molecular weight data and no size-exclusion chromatography was done; hence, homodimer structure remained uncertain. The recently published crystal structure of FAP showed that the enzymatically active, soluble form of recombinant FAP expressed by a baculovirus system is a homodimer, but whether monomers have enzymatic activity was not reported\textsuperscript{36}. Each subunit comprises two distinct domains, an $\alpha/\beta$-hydrolase domain (residues 27-53 and 493-760) and a beta-propeller domain (54-492)\textsuperscript{36}. Based on its crystal structure, a small region of the propeller (residues 232-258) and three regions of the hydrolase domain (residues 651-657, 706-731 and 737-757) can be predicted to participate in the dimerization of FAP. One might speculate that based on this information, homodimers should be separable into monomers by site-directed mutagenesis or peptides that disrupt dimerization so that it could be determined if monomers possess enzymatic activity, but as yet, this has not been done with FAP. Data in the present study, however, suggest that monomers lack enzymatic activity, reinforcing the speculation that if a peptide inhibitor were designed to compete with dimer interface sequences, and thereby prevent dimerization, then it might be possible to abate FAP activity in pathologic conditions such as certain neoplasms\textsuperscript{23,37,38}.

While our data support that APCE is soluble FAP and that its homodimeric structure appears to be required for activity expression, we also explored whether the enzymes shared substrate specificities. The synthetic substrates, Ala-Pro-AFC and Z-Gly-Pro-AMC, and the FRET peptide, synthesized to contain P4-P4’ residues of Met-
α2AP (Table 1), were used for kinetic assessments. We determined that each enzyme had a pH 7.5 optimum at which all kinetic analyses were then performed. Enzyme concentrations and $k_{cat}$ values were calculated using the subunit molecular weights and extinction coefficients as we determined for APCE and rFAP. The kinetic parameters for each enzyme were identical within acceptable error, which again supports rFAP and APCE being essentially the same enzyme. FAP has been reported to prefer Ala-Pro-AFC over Lys-Pro-AFC, and Gly-Pro-AFC$^{13}$. Z-Gly-Pro-AMC, essentially representing a tripeptide, has a 2-3x higher affinity for both rFAP and APCE than does Ala-Pro-AFC (Table 1), which suggests that both enzymes are actually endoproteinases with the ability to cleave bonds very near the N-terminus of the substrate. The FRET peptide has the highest affinity among the three substrates for rFAP and APCE, and the kinetic efficiency ($k_{cat}/K_m$) of rFAP and APCE towards this substrate is the highest, indicating that other residues within the P4-P4’ region must contribute to the substrate specificity of both enzymes$^{39}$.

rFAP, just as APCE, also cleaves the Pro12-Asn13 bond of Met-α2AP. Initially for calculating kinetic parameters, we tried to quantitate a large fragment (residues 13-464) from the cleavage of Met-α2AP by densitometric scanning of time-dependent disappearance of Met-α2AP bands on immunoblots. But due to the large inherent error we encountered with this method, we elected to quantitate the N-terminal cleavage product (residues 1-12; MEPLGRQLTSGP) of the Pro12-Asn13 bond of Met-α2AP by LC/MS. We found small amounts of cleavage products corresponding to L4-P12 peptide and an oxidized version of the M1-P12 peptide along with the expected peptide M1-P12,
all of which complicated data analyses for determining reliable individual $K_m$ and $k_{cat}$ values for rFAP or APCE toward the Pro12-Asn13 bond of Met-$\alpha_2$AP. Under competitive situations, however, the substrate preference of an enzyme can be expressed as relative $k_{cat}/K_m$ values\(^{33}\), which yields a value ratio of the Pro3-Leu4 bond and the Pro12-Asn13 bond as $\sim 1:16$. The big difference in the relative $k_{cat}/K_m$ values can be explained by Gly and Asn (P2 and P1’) being more preferred by APCE (soluble FAP) than are Glu and Leu (P2 and P1’), the latter characterizing the Pro3-Leu4 bond region\(^{39}\).

Soluble FAP (APCE) circulating in human plasma appears to have a role in making $\alpha_2$AP more efficient for protection of extravascular fibrin that forms as a host defense to staunch hemorrhage\(^5\). Since the source of the plasma form of this enzyme was uncertain, it seemed important to know if cell-membrane-bound FAP, wherever expressed in vivo under normal conditions, might be shed to somehow enter the circulation. We reasoned that if APCE is indeed a soluble form of FAP, the two should be virtually identical. Endothelial cells might produce this enzyme, but while cultured endothelial cells have been reported to express FAP mRNA, translation of FAP protein was not documented\(^{17}\). FAP is not expressed by normal human liver, but in the cirrhotic liver a strong correlation exists between FAP expression and severity of liver fibrosis\(^{34,40}\). Our results support soluble FAP as a derivative of membrane-bound FAP, with a role in human fibrinolysis, but otherwise, the origin and additional functions of membrane-bound or soluble FAP under normal conditions remain enigmatic.
REFERENCES


14. David F, Bernard AM, Pierres M, Marguet D. Identification of serine 624, aspartic acid 702, and histidine 734 as the catalytic triad residues of mouse dipeptidyl-peptidase


Table 1. Kinetic Parameters for rFAP and APCE

<table>
<thead>
<tr>
<th>Substrate</th>
<th>rFAP</th>
<th></th>
<th></th>
<th>APCE</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$K_m$ (mM)</td>
<td>$k_{cat}$ (min$^{-1}$)</td>
<td>$k_{cat}/K_m$ (min$^{-1}$/mM)</td>
<td>$K_m$ (mM)</td>
<td>$k_{cat}$ (min$^{-1}$)</td>
<td>$k_{cat}/K_m$ (min$^{-1}$/mM)</td>
</tr>
<tr>
<td>Z-Gly-Pro-AMC</td>
<td>0.101±0.018*</td>
<td>30.5±3.2</td>
<td>306±38</td>
<td>0.124±0.024</td>
<td>32.2±7.1</td>
<td>260±13</td>
</tr>
<tr>
<td>Ala-Pro-AFC</td>
<td>0.323±0.052</td>
<td>64.7±4.6</td>
<td>202±18</td>
<td>0.272±0.014</td>
<td>59.7±8.1</td>
<td>218±19</td>
</tr>
<tr>
<td>FRET peptide#</td>
<td>0.029±0.004</td>
<td>72.2±12.5</td>
<td>2491±159</td>
<td>0.026±0.005</td>
<td>63.4±10.1</td>
<td>2445±109</td>
</tr>
</tbody>
</table>

* Each value is the mean ± SD of three experiments.

# FRET peptide is Arg-Lys(DABCYL)-Thr-Ser-Gly-Pro-Asn-Gln-Glu-Gln-Glu(EDANS)-Arg.
FIGURE LEGENDS

**Figure 1.** SDS-PAGE and Western blot analysis of purified APCE and rFAP. (A) Coomassie-stained reduced SDS-PAGE analysis on a 4-12% gradient gel. (B) Western blot of a duplicate gel to that in (A), using a monoclonal antibody (F19) to native FAP. Although the loading in each lane (1.4 µg protein) is equivalent, the F19 antibody appears to bind rFAP to a greater extent, possibly relating to differences in glycosylation.

**Figure 2.** pH curve. FRET peptide, Arg-Lys(DABCYL)-Thr-Ser-Gly-Pro-Asn-Gln-Glu-Gln-Glu(EDANS)-Arg was incubated with purified APCE or rFAP at the indicated pH. Two buffer systems were used: 0.1 M sodium phosphate for the pH range 5.8-7.5 and 0.1 M Tris-HCl for the pH range 7.9-9.0; each value represents the average of duplicate incubations. One unit of enzyme activity is defined as pmoles of Asn-Gln-Glu-Gln-Glu(EDANS)-Arg released per minute at 22°C.

**Figure 3.** Sequence Alignment. FAP sequence from GenBank (accession #NM_004460) aligned with APCE and rFAP N-terminal sequences and tryptic peptides characterized by LC/MS/MS.

**Figure 4.** Comparison of APCE with rFAP for molecular size and gelatinase activity. A protein load of 1.4 µg was applied to each electrophoretic lane. Each was stained with Coomassie brilliant blue G-250. Clear zones are due to gelatinase activity. 

*Panel A.* Electrophoresis in the presence of SDS on polyacrylamide gel containing
copolymerized gelatin. **Panel B.** Electrophoresis in the absence of SDS, but on the identical type of gel. Comparable gelatinolysis is observed for the two enzymes; however, the mobilities of the single broad zones of lysis differ slightly and cannot be related to MW standards since electrophoresis was done without SDS. **Panel C.** Analytical size exclusion chromatography of APCE (blue) and rFAP (red). Standard (black): a, thyroglobulin (670 kDa); b, γ-globulin (158 kDa); c, ovalbumin (44 kDa); d, myoglobin (17 kDa). The inset shows a standard plot of the logarithmic molecular weight against the elution volume (Ve)/void volume (Vo).

**Figure 5. Immunoblots of Met-α₂AP cleaved by rFAP to Asn-α₂AP.** Met-α₂AP was incubated with purified rFAP (0.2 µg) for selected times as shown at the top of each lane. At each time the incubation mixture was analyzed using two different antibodies. (A) rFAP diminished Met-α₂AP-band intensity with time as determined by immunoblotting with an antibody specific for the N-terminal peptide of Met-α₂AP. (B) α₂AP-band intensity was not diminished when blotted by an antibody to the C-terminal region common to both Met-α₂AP and Asn-α₂AP.

**Figure 6. LC/MS analysis of peptides released from the rFAP digest of Met-α₂AP.** (A) Total ion current (m/z = 350-1800 amu) of a control sample containing Met-α₂AP but lacking rFAP, showing only an internal standard peak (IS). (B) Total ion current of Met-α₂AP incubated with rFAP: IS, oxidized N-terminal 12 amino acid M1(o)-P12 peptide and the 12 amino acid M1-P12 peptide are observed. (C) Reconstructed ion currents (464.4-465.9 amu) of the rFAP-treated sample for the doubly charged ions represented
the L4-P12 peptide, indicating the presence of an additional cleavage site within the N-terminal 12 residue peptide.

**Figure 7.** rFAP or APCE-catalyzed cleavage of Met-α2AP as a function of Met-α2AP concentration. Cleavage rates are given in pmoles of peptides released per hour for one pmol of enzyme.
Lee_fig6

A

B

C

Intensity (cps x 10^{-4})

Intensity (cps x 10^{-3})

Time (min)
Lee_fig7

![Graph showing cleavage rate vs Met-α2AP (μM)]

- **Cleavage rate (pmol peptide/h)**
- **Met-α2AP (μM)**
- **APCE** (blue triangles)
- **rFAP** (red circles)
- Key points:
  - Pro12-Asn13 Bond
  - Pro3-Leu4 Bond
Antiplasmin-cleaving enzyme is a soluble form of fibroblast activation protein

Kyung N Lee, Kenneth W Jackson, Victoria J Christiansen, Chung S Lee, Jin-Geun Chun and Patrick A McKee