

The human α_2 -plasmin inhibitor: functional characterization of the unique plasmin(ogen)-binding region

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Abstract The human α_2 -plasmin inhibitor (A2PI) possesses unique N- and C-terminal extensions that significantly influence its biological activities. The C-terminal segment, A2PIC (Asn³⁹⁸-Lys⁴⁵²), contains six lysines thought to be involved in the binding to lysine-binding sites in the kringle domains of human plasminogen, of which four (Lys⁴²², Lys⁴²⁹, Lys⁴³⁶, Lys⁴⁵²) are completely and two (Lys⁴⁰⁶, Lys⁴¹⁵) are partially conserved. Multiple Lys to Ala mutants of A2PIC were expressed in *Escherichia coli* and used in intrinsic fluorescence titrations with kringle domains K1, K4, K4 + 5, and K1 + 2 + 3 of human plasminogen. We were able to identify the C-terminal Lys⁴⁵² as the main binding partner in recombinant A2PIC (rA2PIC) constructs with isolated kringles. We could show a cooperative, zipper-like enhancement of the interaction between C-terminal Lys⁴⁵² and internal Lys⁴³⁶ of rA2PIC and isolated K1 + 2 + 3, whereas the other internal lysine residues contribute only to a minor extent to the binding process. Sulfated Tyr⁴⁴⁵ in the unique C-terminal segment revealed no influence on the binding affinity to kringle domains.

Keywords α_2 -Plasmin inhibitor · Plasminogen-binding region · Kringle · Lysine-binding site · Fluorescence titration

Abbreviations

6-AHA 6-Aminohexanoic acid
A2PI Human α_2 -plasmin inhibitor
A2PIC C-terminal moiety of A2PI (Asn³⁹⁸-Lys⁴⁵²)

rA2PIC Recombinant A2PIC (A2PIC/Pro399Ala)
FX_a Activated coagulation factor X
His-tag Peptide MNHKVH₆MELGTIEGR
K1 Kringle 1 of Pgn (Cys⁸⁴-Cys¹⁶²), generated as rK1 (Lys⁷⁸-Glu¹⁶⁴)
K2 Kringle 2 of Pgn (Cys¹⁶⁶-Cys²⁴³), generated as rK2 (Cys162Thr/Glu163Ser/Glu¹⁶⁴-Thr²⁴⁴/Cys169Gly)
K3 Kringle 3 of Pgn (Cys²⁵⁶-Cys³³³), generated as rK3 (Thr²⁵³-Ser³³⁵/Cys297Ser)
rK3mut Mutated K3 domain of Pgn (rK3/Lys311Asp)
K4 Kringle 4 of Pgn (Cys³⁵⁸-Cys⁴³⁵), generated as fragment Val³⁵⁵-Ala⁴⁴⁰ of Pgn
K5 Kringle 5 of Pgn (Cys⁴⁶²-Cys⁵⁴¹), generated as rK5 (Thr⁴⁵⁶-Ala⁵⁴³, Pro457Ala)
K1–3 Kringles 1–3 of Pgn (Cys⁸⁴-Cys³³³), generated as fragment Tyr⁸⁰-Val³³⁸ of Pgn
K4–5 Kringles 4–5 of Pgn (Cys³⁵⁸-Cys⁵⁴¹), generated as rK4–5 (Val³⁵⁵-Phe⁵⁴⁶)
LB Broth Luria–Bertani Broth
LBS Lysine-binding site
PAN Plasminogen N-terminal domain
Pgn Human plasminogen
Plm Human plasmin
RCL Reactive center loop
Serpin Serine protease inhibitor
t-AMCHA *trans*-4-(Aminomethyl) cyclohexanecarboxylic acid

Introduction

The human α_2 -plasmin inhibitor (A2PI), a member of the serine protease inhibitor (serpin) super family [1], was

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discovered in 1976 [2–4]. A2PI is a single-chain plasma glycoprotein [3, 5] synthesized in the liver [6, 7] with a carbohydrate content of 14% [5]. It has been shown that A2PI contains four fully glycosylated N-glycosylation sites (Asn⁸⁷, Asn²⁵⁰, Asn²⁷⁰, and Asn²⁷⁷) [8, 9]. Two different N-terminal forms of A2PI circulate in human plasma: Met-A2PI, a 464-residue protein with Met as N-terminus, and Asn-A2PI, a 452-residue form with Asn as N-terminus [10, 11]. The numbering in this publication is according to Asn-A2PI. Mature human A2PI, which corresponds to Asn-A2PI [10–13], contains four cysteine residues (Cys³¹, Cys⁶⁴, Cys¹⁰⁴, and Cys¹¹³), two of which form a single disulfide bridge (Cys³¹-Cys¹⁰⁴) [14] and a sulfated tyrosine residue (Tyr⁴⁴⁵) of unknown function [15]. A2PI possesses unique N- and C-terminal extensions that significantly influence its biological activities. The N-terminal portion at Gln² is cross-linked to Lys³⁰³ in the A α chain of fibrin (ogen) by activated coagulation factor XIII_a [16, 17]. The C-terminal moiety of A2PI (A2PIC) extends approximately 55 amino acids beyond the consensus serpin sequence [12] and is highly conserved among species, which indicates a distinct and specific function for the C-terminal Asn³⁹⁸-Lys⁴⁵² segment [18] (Scheme 1). It has been shown that A2PIC mediates the binding to plasmin(ogen) kringles [19–21]. The X-ray crystal structure of N-terminally truncated murine A2PI, which shares 71% sequence identity with human A2PI, shows that A2PI exhibits the typical serpin fold [22]. Unfortunately, the structure of the unique N- and C-terminal extensions still remains unsolved.

A2PI is the principal physiological inhibitor of plasmin (Plm) [2–4], the key enzyme of the fibrinolytic cascade. Plasminogen (Pgn), the zymogen of Plm, is a 90-kDa multidomain plasma glycoprotein synthesized in the liver [23]. It is composed of an N-terminal PAN module [24], five kringle domains [25], and a trypsin-like serine protease unit [26]. All kringle domains contain a lysine-binding site (LBS), which is responsible for the interaction of Pgn with, e.g., fibrin, A2PI, and small molecules with positively charged group(s) such as lysine and lysine analogous, like 6-aminoheptanoic acid (6-AHA) or *trans*-4-(aminomethyl) cyclohexanecarboxylic acid (*t*-AMCHA) (reviewed in

[27]). Plm reacts with A2PI to form an enzymatically inactive 1:1 equimolar complex [3, 5, 28, 29] upon cleavage of the scissile peptide bond (Arg³⁶⁵-Met³⁶⁶) in the reactive center loop (RCL) of A2PI [30]. Kinetic data suggest a two-step reaction for the rapid inhibition of Plm: a fast reversible second-order reaction (k_1 , $3.8 \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$) followed by a slower irreversible first-order transition (k_2 , $4.2 \times 10^7 \text{ M}^{-3} \text{ s}^{-1}$) [19, 28, 29, 31, 32]. In the second step, the scissile peptide bond in the RCL of A2PI is cleaved by the protease subunit of Plm. The first step is rate limiting and results in the formation of a non-covalent complex between A2PIC and kringle domains of Plm. The complex formation between Plm and A2PI is competitively inhibited by plasminogen kringles 1–3 (K1–3) and kringle 4 (K4) or by 6-AHA [19]. A2PI inhibits miniplasmin, a Plm fragment possessing only kringle 5 (K5) and the fully active protease domain, approximately 60-fold slower than intact Plm [31]. 30% of mature A2PI in human plasma is partially degraded lacking at least 26 residues (Gly⁴²⁷-Lys⁴⁵²) of the C-terminal segment [20]. This truncated form also inhibits Plm less rapidly than native A2PI [33, 34]. These observations emphasize the physiological relevance of this non-covalent interaction between the unique A2PIC and Plm kringle domains.

Human A2PIC contains six lysine residues, of which four (Lys⁴⁵², Lys⁴³⁶, Lys⁴²⁹, and Lys⁴²²) are completely and two (Lys⁴⁰⁶ and Lys⁴¹⁵) are partially conserved (Scheme 1). It is well known that internal as well as C-terminal lysine residues often exhibit a distinct affinity for LBS in kringles. Initial studies gave evidence for the participation of the C-terminal Lys⁴⁵² residue in the binding to the Plm kringle domains [18]. Furthermore, a synthetic C-terminal peptide of A2PI (Gly⁴²⁷-Lys⁴⁵²) has been shown to competitively block the reversible association of A2PI to Plm kringle domains [20, 21]. Substitution of Lys⁴⁵² with arginine or its removal with carboxypeptidase B was found to reduce the interaction between these peptides and Plm [21, 35]. These data led to the suggestion that the multiple lysine residues within A2PIC contribute, probably in a zipper-like manner, to the binding to the ‘five-kringle array’ in Plm [18, 35]. In contrast to these

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human  NPSAPRELKEQQDSPGNKDFLQSLKGFPRGDKLFPGDLKLVPPEEDYPQFGSPK
bovine NPGAQPERKEQQDSDPGKDSFQDHKGLPRGDKPFDPDLKLGPPEEDYAQPSSK
murine  NPSALPQLQEQRDSDPNRLIGQNDKADFHGGKTFGPDLKLAPRMEEDYPQFSSSK
rabbit  NPSAQPERKEQQDSDPHRDPSQPKSFPHGDKLFSPDLKLAPPSEEDYPQLSSSK
rat     DESAQPPQEQQDSDPNRRLDQNDKADIPGGKTFAPDLKLVRLEEDYPQFSSSK
      :*,*  :  :*:***.  :  *  *  *  *  *  *  *  *  *  *  *  *  *  *
      |          |          |          |          |          |
      400        410        420        430        440        450

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Scheme 1 Multiple sequence alignment of human, bovine, murine, rabbit, and rat A2PIC. The numbering is according to the Asn form of human A2PI. The Pro399Ala mutation is shaded in gray, sulfated

Tyr⁴⁴⁵ is *underlined*, Lys residues are in *bold* type and the symbols *asterisk*, *colon*, *dot*, and *blank space* stand for identity, high, average, and poor similarity, respectively

findings, kinetic data by Wang et al. [36, 37] indicate that Lys⁴³⁶ might play an important role in the interaction between Plm and A2PI.

To elucidate the exact role of the four completely conserved lysine residues (Lys⁴⁵², Lys⁴³⁶, Lys⁴²⁹, and Lys⁴²²) and the partially conserved Lys⁴¹⁵ and Lys⁴⁰⁶ in the binding to the LBS of Pgn kringle domains, we have expressed wild-type, single, double, triple, quadruple, quintuple, and sextuple Lys mutants of recombinant A2PIC(Pro399Ala) (rA2PIC) in *Escherichia coli*. Association constant (K_a) values of the different mutant rA2PIC constructs to human recombinant kringle 1 (rK1) and kringles 4–5 (rK4–5) and human K4 and K1–3, which were obtained by limited proteolysis of Pgn with porcine elastase, have been determined applying intrinsic fluorescence ligand titrations. To investigate the influence of sulfated Tyr⁴⁴⁵ in the interaction with human kringles rK3 and K4, the K_a values of synthetic peptides Glu⁴⁴²-Lys⁴⁵² of the C-terminal region of A2PI, A2PIC(Glu⁴⁴²-Lys⁴⁵²), with and without sulfated Tyr⁴⁴⁵ have also been obtained by intrinsic fluorescence ligand titrations.

Materials and methods

Materials

Restriction endonucleases, *Pfu* DNA Polymerase, T4 DNA Ligase, activated coagulation factor X (FX_a) and trypsin (porcine, seq. grade modified) came from Promega (Madison, WI, USA). Porcine elastase was purchased from Serva (Heidelberg, Germany). UltraPure™ Agarose was obtained from Invitrogen (Carlsbad, CA, USA). E.Z.N.A.® MicroSpin Gel Extraction Kit was purchased from peQLab (Erlangen, Germany). GenElute™ HP Plasmid Miniprep Kit and Luria–Bertani Broth (LB Broth) were obtained from Sigma-Aldrich (St. Louis, MO, USA). Primers were synthesized either by Microsynth (Balgach, Switzerland) or Sigma-Aldrich. Sephadex G-50m and G-50sf were purchased from GE Healthcare (Waukesha, WI, USA). Acrylamide, Bio-Gel P-6DG and Bio-Gel P-300 were from Bio-Rad (Hercules, CA, USA). Lysine Bio-Gel P-300 was prepared according to Brunisholz et al. [38] and Ni²⁺-NTA Agarose was purchased from QIAGEN (Venlo, The Netherlands).

Bacterial strains and plasmids

Escherichia coli strain XL1-Blue {*recA1 endA1 gyrA96 thi-1 hsdR17 supE44 relA1 lac* [F' *proAB lacI^qZAM15 Tn10* (Tet^r)]} was obtained from Stratagene (La Jolla, CA, USA) and used for routine transformations and plasmid

preparations. Rosetta™ [F⁻ *ompT hsdS_B(r_B⁻ m_B⁻) gal dcm* (DE3) pRARE [39] (Cam^R)] *E. coli* strain was purchased from Novagen (Madison, WI, USA) and used for the expression of recombinant A2PIC (rA2PIC) and rA2PIC mutants. The pColdII plasmid was obtained from Takara (Kyoto, Japan) and used for the expression of rA2PIC. The plasmid contains a *lac* operator, a *cspA* 5'-untranslated region, a translation enhancing element, a hexaHis Tag sequence (His-tag), a multiple cloning site, and a *cspA* 3'-untranslated region immediately upstream of the *cspA* transcription initiation site and carries an ampicillin resistance [40]. The pET100/D-TOPO® plasmid (Invitrogen) containing the rA2PIC peptide was prepared as described previously [8].

Methods

DNA manipulations

Isolation of plasmid DNA was done as described by the manufacturer. DNA fragments generated by restriction endonuclease cleavage were purified on 1.2 or 0.5% (depending on their size) agarose gels, stained with ethidium bromide and visualized by UV light. DNA fragments were recovered from the agarose gels by the E.Z.N.A.® MicroSpin Gel Extraction Kit according to the supplier's instructions.

Construction of expression vectors for wild-type and mutant rA2PIC peptides

The C-terminal region of A2PI (Asn³⁹⁸-Lys⁴⁵²) was amplified in a PCR using the pET100/D-TOPO® plasmid containing rA2PIC or rA2PIC(Lys452Ala) as a template. The 5' primer F/rA2PIC, which binds to the FX_a cleavage site of the sense strand of the pET100/D-TOPO® plasmid and the first four bases of rA2PIC, introduced a *KpnI* and a FX_a cleavage site upstream of the codon for Asn³⁹⁸. The 3' primer R/rA2PIC, complementary to the coding strand of rA2PIC, was used to introduce two stop codons and a *BamHI* cleavage site downstream of the codon for Lys⁴⁵². Primers used are listed in Table 1. The *KpnI* and *BamHI* cleaved PCR product was cloned into the *KpnI*-/*BamHI*-digested pColdII plasmid and transformed into XL1-Blue *E. coli* cells. Clones were selected and the insert was verified by DNA sequence analysis at Microsynth (Balgach, Switzerland).

For the construction of the expression vectors rA2PIC (Lys452Ala) and rA2PIC(Lys452Ser) the 3'-mismatch primers Lys452Ala and Lys452Ser (Table 1) were used, respectively. The point mutation at the position Lys⁴⁵² to Ser was introduced to improve the solubility of rA2PIC(Lys452Ala).

Table 1 Primers used for the generation of rA2PIC mutants, pColdIImut vector, rK5, and rK4–5

Primer name	DNA sequence
F/pelB	5'-GGCGACTAGTATGAAATACCTGCTGCCGACCG-3'
R/pelB	5'-CGGCACTAGTCGAATTAATTCGGATATCCATGGCC-3'
F/pColdIImut	5'-GGCGACTAGTCATCATCATCATCATCATATGGAGCTCGG-3'
R/pColdIImut	5'-CGGCACTAGTCACTTTGTGATTCATGGTGTATTACCTC-3'
F/rA2PIC	5'-CACCGGTACCATCGAGGGTAGAAACC-3'
R/rA2PIC	5'-GCGCGGATCCCTATTACTTGGG-3'
F/rK4–5	5'-CACCGGTACCATCGAGGGTAGAGTACAGGACTGCTACCAT-3'
R/rK4–5	5'-CGCGGGATCCCTATTAATAAATGAAGGGGCCGACA-3'
F/rK5	5'-CACCGGTACCATCGAGGGTAGAACTGCATCCGAAGAAGACTGTATG-3'
R/rK5	5'-CGCGGGATCCCTATTAGGCCGCACACTGAGGGACAT-3'
Lys406Ala	5'-GCGGGAGCTCGCGGAACAGCAGGATTC-3'
Lys415Ala	5'-GATTCCCCGGCAACGCGGACTTCCTCCAGAG-3'
Lys422Ala	5'-TCCTCCAGAGCCTGGCAGGCTTCCCCCG-3'
Lys429Ala	5'-GGCTTCCCCGCGGAGACGCGCTTTTCGGCCCTGACTT-3'
Lys436Ala	5'-GCTTTTCGGCCCTGACTTAGCACTTGTGCCCCCATGGAG-3'
Lys452Ala	5'-GCGCGGATCCCTATTAAGCGGG-3'
Lys452Ser	5'-CGCGGGATCCCTATTAAGAGGGGCTGCCAAACTGGGGG-3'
Pro399Ala	5'-CGAGGGTAGAAACGCCAGTGCACCGC-3'

Mismatched bases are shown in bold type

Construction of expression vectors for single, double, triple, quadruple, quintuple, and sextuple Lys to Ala mutant rA2PIC peptides by site-directed mutagenesis

The method used is based on the QuikChange[®] Site-Directed Mutagenesis Kit (Stratagene). In brief, the appropriate pColdII expression vector (0.5 µg/µL) was selected as template and used in a PCR with the suitable 5' primer in a concentration of 1.25 µg/µL. Primers used are listed in Table 1. After amplification, the template DNA was digested by adding 1 µL *DpnI* (8 h at 37°C). 4% of *DpnI*-digested PCR product was then transformed into XL1-Blue *E. coli* cells. Clones were selected and the insert was verified by DNA sequence analysis.

Substitution of Pro³⁹⁹ to Ala in all rA2PIC peptides

A point mutation at position Pro³⁹⁹ was introduced in all rA2PIC peptides using site-directed mutagenesis as described above, to improve the cleavage efficiency of FX_a [41, 42]. The Pro399Ala primer is shown in Table 1. All mentioned rA2PIC peptides in this publication are hereafter referred to as carriers of the Pro399Ala mutation.

Construction of the pColdIImut(*pelB*) vector

To overcome the poor expression yields with the pColdII expression system for certain constructs, we have modified the pColdII vector by introducing the *pelB* leader sequence, which can also be used as enhancer for the total protein expression [43], between the TEE and His-tag sequence of

the pColdII vector. pColdIImut was amplified in a PCR using the pColdII vector as template. The 5' primer F/pColdIImut and 3' primer R/pColdIImut were used to introduce a *SpeI* cleaving site (ACTAGT) between the TEE and the His-tag sequences. Cleavage with *SpeI* and ligation, followed by transformation into XL1-Blue *E. coli* cells and selection of appropriate clones, yielded the mutated pColdIImut vector. The *pelB* leader sequence was amplified in a PCR using the pET22b(+) vector as a template. The 5' primer F/pelB and 3' primer R/pelB were used to introduce a *SpeI* cleaving site up- and downstream of the *pelB* leader sequence. The *SpeI* cleaved PCR product was cloned into the *SpeI*-digested pColdIImut plasmid and transformed into XL1-Blue *E. coli* cells. Clones were selected and the insert was verified by DNA sequence analysis. The primers used are listed in Table 1.

Construction of the expression vector for rK5 and rK4–5

The rK5 construct (Thr⁴⁵⁶-Ala⁵⁴³) was amplified in a PCR using the pPLGKG plasmid (contains the cDNA of human Pgn) as a template. The 5'-mismatch primer F/rK5, which binds to the sense strand of Pgn, introduced a *KpnI* and a FX_a cleavage site upstream of the codon for Thr⁴⁵⁶. This 5'-mismatch primer was also used to introduce a point mutation at position Pro⁴⁵⁷ to Ala to enhance the cleavage efficiency of FX_a [41, 42]. The 3' primer R/rK5, complementary to the coding strand of Pgn, was used to introduce two stop codons and a *BamHI* cleavage site downstream of the codon for Ala⁵⁴³. The primers used are compiled in Table 1. The cloning, transformation, and verification of

clones was carried out according to the above-described procedure for the construction of expression vectors for wild-type and mutant rA2PIC peptides except that the pColdIImut(*pelB*) vector was used. The mentioned rK5 in this article is hereafter referred to as rK5 construct (Thr⁴⁵⁶-Ala⁵⁴³) and carrier of the Pro457Ala mutation.

The rK4–5 construct (Val³⁵⁵-Phe⁵⁴⁶) was generated as described above except that 5' primer F/rK4–5, 3' primer R/rK4–5 and pColdII were used. The mentioned rK4–5 in this publication is hereafter referred to as rK4–5 construct (Val³⁵⁵-Phe⁵⁴⁶).

Expression and isolation of wild-type and mutant rA2PIC peptides

The expression was carried out according to a slightly modified protocol of the manufacturer's instructions. The agar plates and LB Broth culture media always contained 100 μ g ampicillin/mL and 34 μ g chloramphenicol/mL. Cells were grown at 37°C in LB Broth medium in 2-L Erlenmeyer flasks to an OD₆₀₀ of about 0.4–0.6 absorbance units (AU). The cell cultures were refrigerated at 15°C for 35 min before isopropyl- β -D-thiogalactopyranoside was added to a final concentration of 2 mM. The cultures were thereafter incubated at 15°C and 220rpm for 21–24 h and finally harvested by centrifugation (Sorvall RC 3C Plus, 4,000g, 65 min, 4°C). The cell pastes were stored at –80°C until further use. The protein expression was verified by SDS-PAGE (15%) and the proteins were stained with Coomassie brilliant blue G-250.

For the isolation of wild-type and mutant rA2PIC peptides, the cell pastes were thawed and suspended in lysis buffer (6 M guanidine hydrochloride and 300 mM NaCl in 50 mM Tris, pH 8.0; 10 mL/g of wet cell paste), stirred overnight at 4°C and centrifuged (Sorvall RC 5B Plus, 12,000g, 45 min, 4°C). The supernatants were loaded onto a Ni²⁺-NTA Agarose column (2.2 cm \times 9 cm; 1 mL Ni²⁺-NTA Agarose/1,000 mL cell culture) equilibrated with the lysis buffer, pH 8.0. The column was thereafter washed with lysis buffer, pH 8.0 and 6.3. The recombinant peptides were eluted from the columns with lysis buffer, pH 4.5. The peptides were desalted by dialysis against 20 mM NH₄HCO₃, lyophilized and stored at 4°C until further use. The peptides were analyzed by RP-HPLC, amino acid analysis, ESI-MS, and Edman sequence analysis.

Expression, isolation, and characterization of recombinant kringle constructs rK1, rK3, rK5, and rK4–5

rK1 and rK3 were expressed, isolated, refolded and characterized according to Marti et al. [44, 45].

rK4–5 and rK5 were expressed and isolated according to the above-described protocol for wild-type and mutant

rA2PIC peptides except that the proteins were refolded instead of desalting by dialysis. rK4–5 was refolded and isolated according to Douglas et al. [46]. rK5 was refolded according to the same protocol as rK4–5 [46]. In addition, isolated rK5 was lyophilized and further purified by size exclusion chromatography on a Sephadex G-50sf column (1.5 cm \times 100 cm; 0.13 M formic acid). The constructs were analyzed as mentioned above.

Isolation of human plasminogen and generation of the K1–3 and K4 domains

Pgn was isolated from citrated plasma by affinity chromatography on Lysine Bio-Gel P-300 as previously described [38]. K1–3 and K4 were prepared via limited proteolysis of Pgn with porcine elastase, isolated and purified as previously described [25].

Purification and characterization of the synthetic peptides A2PIC(Glu⁴⁴²-Lys⁴⁵²) with and without sulfated Tyr⁴⁴⁵

The synthetic peptides A2PIC(Glu⁴⁴²-Lys⁴⁵²) and A2PIC(Glu⁴⁴²-Lys⁴⁵²/Tyr⁴⁴⁵[SO₄]) were purified by preparative RP-HPLC on a butyl column (Wide-PoreTM, 5 μ m, 4.6 mm \times 250 mm; J.T. Baker) with a Hewlett-Packard liquid chromatograph 1090. 50 mM ammonium acetate in water, pH 7.0, was used as solution A and 25 mM ammonium acetate, 80% (v/v) acetonitrile in water, pH 7.0, as solution B. A linear gradient (0–60% solution B in 60 min) was used at a flow rate of 1 mL/min.

RP-HPLC

RP-HPLC separations were carried out on an Aquapore butyl column (BU-300, 7 μ m, 300 Å, 2.1 mm \times 100 mm; Perkin Elmer) or on a Nucleosil 300 phenyl column (7 μ m, 2 mm \times 100 mm; Dr. Maisch HPLC, Ammerbuch, Germany) with a Hewlett-Packard liquid chromatograph 1090. 0.1% (v/v) trifluoroacetic acid (TFA) in water was used as solution A and 0.1% (v/v) TFA, 80% (v/v) acetonitrile in water as solution B. A linear gradient (0–100% solution B in 60 min) was used at a flow rate of 0.3 mL/min.

Amino acid analysis

Samples were hydrolyzed in the gas phase with 6 M HCl containing 0.1% (v/v) phenol for 22 h at 115°C under N₂ vacuum according to Chang and Knecht [47]. The liberated amino acids were reacted with phenylisothiocyanate, and the resulting phenylthiocarbamoyl amino acids were analyzed by RP-HPLC on a Nova Pack C18 column (4 μ m, 3.9 mm \times 150 mm; Waters) with a Dionex Summit[®] HPLC system (Dionex, Olten, Switzerland) equipped with

an automatic injection system according to Bidlingmeyer et al. [48]. The corresponding ammonium acetate buffer replaced the 0.14 M sodium acetate buffer, pH 6.3.

Mass spectrometry

The mass of the peptides was determined by ESI-MS (VG Platform mass spectrometer; Micromass Instruments, UK). The samples were dissolved in 0.5% (v/v) formic acid, 50% (v/v) acetonitrile in water.

Removal of the His-tag sequence

To remove the His-Tag, constructs were incubated with FX_a at 37°C for 16 h in 50 mM Tris, pH 8.0, containing 100 mM NaCl and 5 mM CaCl₂ at an enzyme/substrate ratio 1:800 (w/w). Cleaved peptides were separated from His-tag and undigested constructs on a Ni²⁺-NTA Agarose column (2.2 cm × 9 cm). rA2PIC and rK5, both without His-tag, were finally desalted either on a Sephadex G-50m or on a Bio-Gel P-6DG column (1.5 cm × 100 cm) (50 mM NH₄HCO₃, pH 8.0).

Tryptic peptide mass mapping

For an explicit identification of the diverse rA2PIC mutants, the peptides were digested with trypsin at 37°C overnight in 0.1 M NH₄HCO₃ at an enzyme/substrate ratio 1:100 (w/w). The peptide mass mapping was carried out by ESI-MS.

Intrinsic fluorescence titration

The influence of wild-type and mutant rA2PIC peptides on the intrinsic fluorescence of different Pgn kringles was measured in 50 mM sodium phosphate, pH 8.0, at 25°C with a Cary Eclipse fluorescence spectrophotometer (Varian) as reported by Menhart et al. [49]. The ligand (wild-type and mutant rA2PIC) was added to a 5-μM kringle solution at intervals of 1.66, 3.3, 5, 10, 20, 33, or 40 μM depending on the binding affinity of the investigated kringle until the intrinsic fluorescence was constant. Fluorescence was measured at 340 or 344 nm, depending on the investigated kringle, with excitation at 296 nm. *K_a* values were determined according to Scatchard [50].

A threefold determination of the peptide content in the ligand solutions by amino acid analysis allowed an accurate evaluation of the *K_a* values.

The *K_a* values for the interaction of A2PIC (Glu⁴⁴²-Lys⁴⁵²) and A2PIC(Glu⁴⁴²-Lys⁴⁵²/Tyr⁴⁴⁵[SO₄]) with Pgn rK3 and K4 were obtained as described above.

Results

Recombinant expression of rA2PIC and rA2PIC mutants

We have successfully applied the pColdII expression system to improve the expression of rA2PIC and rA2PIC mutants. The identification of rA2PIC peptides by SDS-PAGE showed a main band indicating a relatively high level of expression (data not shown). The expression yield of the rA2PIC mutants was in the range of 5–35 mg of pure peptide/2 L of cell culture due to variable expression levels of the rA2PIC constructs. The introduction of the point mutation at position Pro³⁹⁹ to Ala dramatically enhanced the cleavage efficiency of FX_a, since it has been shown that the P2' position at the FX_a cleavage site (IEGR) is important for the enzymatic activity of FX_a [41, 42]. We could overcome the poor solubility of rA2PIC(Lys452Ala) by introducing a Lys⁴⁵² to Ser mutation. All rA2PIC peptides eluted as sharp, symmetrical peaks on RP-HPLC columns. After removal of the His-tag, the measured masses agree well with the calculated values (Table 2). Amino acid analyses and Edman sequence analyses yielded the expected compositions and N-terminal sequences (data not shown), respectively, and both methods were applied for the characterization of all following constructs.

Recombinant expression, purification, and refolding of Pgn rK1, rK3, rK5, and rK4–5

rK1 and rK3 were expressed in *E. coli*, purified, refolded, and characterized as previously described [44, 45]. The isolation over Lysine Bio-Gel P-300 ensured that the

Table 2 Characterization of rA2PIC peptides by ESI-MS (experimental and theoretical values)

Construct	Mass (exp.) (Da)	Mass (theor.) (Da)
rA2PIC(wt)	6,145.0 ± 0.4	6,145.9
rA2PIC/K406A	6,087.4 ± 0.5	6,088.8
rA2PIC/K415A	6,087.5 ± 0.8	6,088.8
rA2PIC/K422A	6,087.9 ± 0.6	6,088.8
rA2PIC/K429A	6,088.0 ± 0.6	6,088.8
rA2PIC/K436A	6,088.8 ± 1.2	6,088.8
rA2PIC/K452A	6,088.0 ± 0.1	6,088.8
rA2PIC/K452S	6,104.3 ± 0.2	6,104.8
rA2PIC/K436/452A	6,031.1 ± 0.3	6,031.7
rA2PIC/K429/452A	6,031.4 ± 0.2	6,031.7
rA2PIC/K429/436A	6,031.8 ± 0.2	6,031.7
rA2PIC/K422/429/436A	5,974.3 ± 0.4	5,974.6
rA2PIC/K415/422/429/436A	5,917.3 ± 0.7	5,917.5
rA2PIC/K406/415/422/429/436A	5,860.4 ± 0.4	5,860.4

disulfide bridges were folded correctly. The measured masses of $10,167.3 \pm 0.6$ Da for rK1 and $9,503.8 \pm 1.0$ Da for rK3 correspond well with the calculated masses of 10,167.2 and 9,505.4 Da, respectively.

rK5 was successfully expressed in *E. coli* using the pColdIImut(*pelB*) vector with which we achieved a significantly higher expression yield compared to the unmodified pColdII plasmid. A point mutation at position Pro⁴⁵⁷ to Ala was introduced to enhance the cleavage efficiency of FX_a as mentioned above. The identification of rK5 by SDS-PAGE showed a main band indicating a relatively high level of expression (data not shown). The expression yield was about 19 mg of pure protein/2 L of cell culture. rK5 was isolated as described above and refolded according to Marti et al. [44, 45]. The measured mass of $9,769.0 \pm 0.6$ Da corresponds well with the calculated value of 9,768.7 Da.

rK4-5 was successfully cloned into the pColdII vector and expressed in *E. coli* [51]. Approximately 5–10 mg of pure protein/2 L of cell culture was obtained. rK4-5 was isolated and refolded as described for rK5 and the measured mass of $23,734.1 \pm 4.9$ Da corresponds well with the calculated mass of 23,735.3 Da.

Characterization of the purified synthetic peptides
A2PIC(Glu⁴⁴²-Lys⁴⁵²/Tyr⁴⁴⁵[SO₄])
and A2PIC(Glu⁴⁴²-Lys⁴⁵²)

The measured masses of $1,295.8 \pm 0.3$ Da for A2PIC(Glu⁴⁴²-Lys⁴⁵²) and $1,376.1 \pm 0.3$ Da for A2PIC(Glu⁴⁴²-Lys⁴⁵²/Tyr⁴⁴⁵[SO₄]) correspond well with the calculated masses of 1,296.4 and 1,376.4 Da, respectively.

Binding of wild-type and mutant rA2PIC to human Pgn K4

Wild-type rA2PIC exhibits a K_a value of 34 ± 2 mM⁻¹ for Pgn K4, which is in the range of previously published data [18]. Replacing the C-terminal Lys⁴⁵² residue by either Ser or Ala substantially decreases the affinity (rA2PIC(K452S), $K_a = 11 \pm 2$ mM⁻¹) by a factor of ~ 3 . Our data confirm that Lys⁴⁵² is the main binding partner in the interaction with the isolated K4 domain in solution, which is in good agreement with published data [18]. The single Lys to Ala mutants of the totally conserved internal Lys residues Lys⁴³⁶ ($K_a = 32 \pm 2$ mM⁻¹), Lys⁴²⁹ ($K_a = 33 \pm 2$ mM⁻¹), and Lys⁴²² ($K_a = 33 \pm 3$ mM⁻¹), and the partially conserved Lys⁴¹⁵ ($K_a = 44 \pm 6$ mM⁻¹) and Lys⁴⁰⁶ ($K_a = 42 \pm 6$ mM⁻¹) contribute only to a minor extent to the total binding affinity to the LBS of K4. The double rA2PIC(Lys 429Ala/Lys436Ala) ($K_a = 29 \pm 0.5$ mM⁻¹), triple rA2PIC(Lys422Ala/Lys429Ala/Lys436Ala) ($K_a = 28 \pm 1$ mM⁻¹), quadruple rA2PIC(Lys415Ala/Lys422Ala/Lys429Ala/

Lys436Ala) ($K_a = 24 \pm 1$ mM⁻¹), and quintuple mutants rA2PIC(Lys406Ala/Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala) ($K_a = 23 \pm 2$ mM⁻¹) exhibit a steady decrease of the K_a values with a decreasing number of Lys residues. The data confirm the weak affinity for internal Lys residues to the LBS of K4 [18, 52, 53]. The sextuple rA2PIC(Lys406Ala/Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala/Lys452Ala) mutant exhibits a very weak affinity ($K_a = \sim 3$ mM⁻¹). A comparison of the fluorescence titration and Scatchard plot of wild-type rA2PIC with mutants rA2PIC(Lys436Ala), rA2PIC(Lys452Ser), and rA2PIC(Lys406Ala/Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala) is given in Fig. 1 and all K_a values are summarized in Table 3. The determined K_a value for 6-AHA of 46 ± 4 mM⁻¹ was in good agreement with published data [18].

Binding of wild-type and mutant rA2PIC to Pgn rK1

The wild-type rA2PIC exhibits a K_a value of 77 ± 4 mM⁻¹ for Pgn rK1 and replacement of the C-terminal Lys⁴⁵²

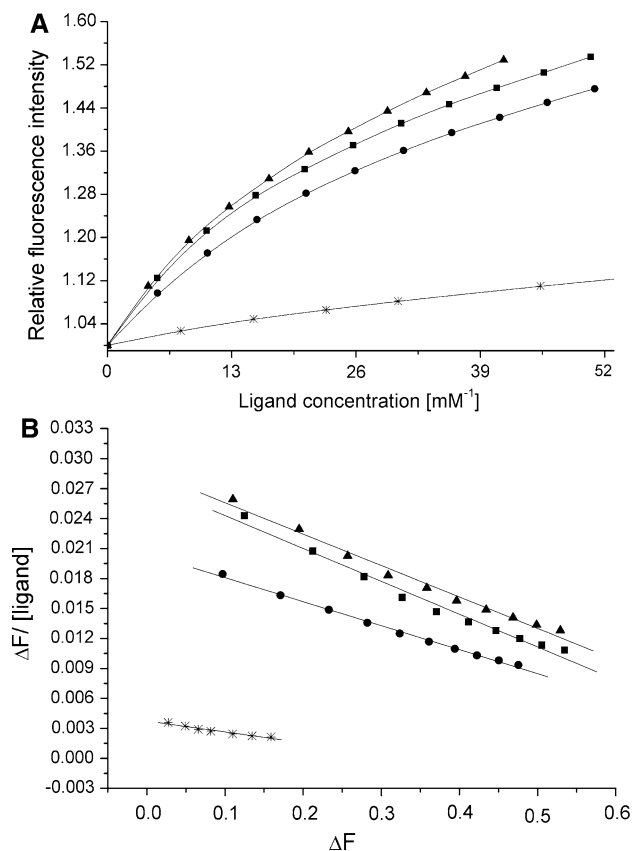


Fig. 1 **a** Averaged fluorescence titration of K4 with wild-type rA2PIC (filled squares), single Lys to Ala mutants rA2PIC(Lys436Ala) (filled triangles), and rA2PIC(Lys452Ser) (star), and quintuple mutant rA2PIC(Lys406Ala/Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala) (filled circles). The fluorescence intensity is normalized to 1 and the x-axis is enlarged. **b** Corresponding Scatchard plots and calculated regression curves

residue by Ser or Ala decreases the affinity (rA2PIC(Lys452Ser), $K_a = 8 \pm 0.5 \text{ mM}^{-1}$) by a factor of ~ 10 , which is in good agreement with published data [18]. The double mutant rA2PIC(Lys429Ala/Lys452Ala) ($K_a = 8 \pm 0.5 \text{ mM}^{-1}$) exhibits the same K_a value as the single mutant rA2PIC(Lys452Ser), while the single mutant of internal Lys⁴³⁶ rA2PIC(Lys436Ala) ($K_a = 70 \pm 1 \text{ mM}^{-1}$) shows no significant difference compared to the wild-type rA2PIC. The quintuple mutant rA2PIC(Lys406Ala/Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala) ($K_a = 53 \pm 7 \text{ mM}^{-1}$) exhibits a partially decreased affinity compared to the wild-type rA2PIC. The sextuple mutant rA2PIC(Lys406Ala/Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala/Lys452Ala) shows a weak K_a value of $\sim 3 \text{ mM}^{-1}$. Our data confirm that Lys⁴⁵² is the main binding partner in the interaction with the isolated rK1 domain in solution, which is in good agreement with published data [18], and that rK1 exhibits only a weak but distinct affinity for internal Lys residues [18, 52, 53]. A selection of the fluorescence titrations and Scatchard plots of wild-type rA2PIC, single rA2PIC(Lys436Ala) and rA2PIC(Lys452Ser), double rA2PIC(Lys429Ala/Lys452Ala), and quintuple rA2PIC(Lys406Ala/Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala) mutants is shown in Fig. 2 and all K_a values are summarized in Table 3. The K_a value of $75 \pm 11 \text{ mM}^{-1}$ for 6-AHA was determined by Frank et al. [18] and is in the same range as wild-type rA2PIC.

Binding of wild-type and mutant rA2PIC to human Pgn K1–3

The K_a values of wild-type rA2PIC ($K_a = 176 \pm 30 \text{ mM}^{-1}$) and internal single mutants of the totally conserved Lys⁴²⁹ ($K_a = 170 \pm 9 \text{ mM}^{-1}$) and Lys⁴²² ($K_a = 177 \pm 22 \text{ mM}^{-1}$) for Pgn K1–3 are in a similar range. The K_a values drop dramatically after replacement of the C-terminal Lys⁴⁵² (rA2PIC(Lys452Ser), $K_a = 17 \pm 4 \text{ mM}^{-1}$) or the internal Lys⁴³⁶ (rA2PIC(Lys436Ala), $K_a = 26 \pm 10 \text{ mM}^{-1}$) residue by Ser or Ala, respectively. The double rA2PIC(Lys429Ala/Lys436Ala) ($K_a = 17 \pm 6 \text{ mM}^{-1}$) and rA2PIC(Lys429Ala/Lys452Ala) ($K_a = 30 \pm 5 \text{ mM}^{-1}$), triple rA2PIC(Lys422Ala/Lys429Ala/Lys436Ala) ($K_a = 22 \pm 7 \text{ mM}^{-1}$), quadruple rA2PIC(Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala) ($K_a = 12 \pm 2 \text{ mM}^{-1}$), and quintuple rA2PIC(Lys406Ala/Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala) ($K_a = 46 \pm 12 \text{ mM}^{-1}$) mutants exhibit a highly reduced affinity, thus confirming the importance of the Lys⁴⁵² and Lys⁴³⁶ residues in the interaction with the isolated triple-kringle domain K1–3 in solution. The fluorescence titrations and Scatchard plots of wild-type rA2PIC, single rA2PIC(Lys436Ala) and rA2PIC(Lys452Ser), double rA2PIC(Lys429Ala/Lys452Ala), and quintuple rA2PIC(Lys406Ala/Lys415Ala/Lys422Ala/

Lys429Ala/Lys436Ala) mutants are shown in Fig. 3. The sextuple mutant rA2PIC(Lys406Ala/Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala/Lys452Ala) exhibits a very weak affinity ($K_a = \sim 4 \text{ mM}^{-1}$) and the Lys analog 6-AHA exhibits a K_a value of $107 \pm 11 \text{ mM}^{-1}$, which is considerably smaller compared to that of wild-type rA2PIC. All K_a values are summarized in Table 3.

Binding of wild-type and mutant rA2PIC to Pgn rK4–5 and rK5

The K_a value of wild-type rA2PIC ($K_a = 18 \pm 1 \text{ mM}^{-1}$) for the rK4–5 double-kringle domain is smaller than for K4 alone. As in K4 and rK1, the C-terminal Lys⁴⁵² (rA2PIC(Lys452Ser), $K_a = 1.4 \pm 0.5 \text{ mM}^{-1}$) is the main binding partner for the interaction with the LBS of isolated rK4–5 in solution. The single rA2PIC(Lys436Ala) ($K_a = 16 \pm 2 \text{ mM}^{-1}$), double rA2PIC(Lys429Ala/Lys436Ala) ($K_a = 16 \pm 5 \text{ mM}^{-1}$), triple rA2PIC(Lys422Ala/Lys429Ala/Lys436Ala) ($K_a = 18 \pm 5 \text{ mM}^{-1}$), quadruple rA2PIC(Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala) ($K_a = \sim 17 \text{ mM}^{-1}$), and quintuple rA2PIC(Lys406Ala/Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala) ($K_a = 17 \pm 4 \text{ mM}^{-1}$) mutants exhibit comparable affinities to the LBS of rK4–5 as wild-type rA2PIC. The sextuple mutant rA2PIC(Lys406Ala/Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala/Lys452Ala) shows no detectable change in intrinsic fluorescence and an accurate K_a value could not be determined. 6-AHA exhibits a K_a value of $\sim 52 \text{ mM}^{-1}$, which is significantly higher than the K_a value of wild-type rA2PIC. All K_a values are summarized in Table 3.

Due to the low affinity of K5 for Lys residues, no significant fluorescence change could be detected. This is consistent with earlier studies, which showed the very weak affinity of K5 for C-terminal and internal Lys residues [18, 54].

Binding of A2PIC(Glu⁴⁴²-Lys⁴⁵²/Tyr⁴⁴⁵[SO₄]) and A2PIC(Glu⁴⁴²-Lys⁴⁵²) to Pgn rK3 and K4

rK3 was used for intrinsic fluorescence titrations with the short synthetic C-terminal segments A2PIC(Glu⁴⁴²-Lys⁴⁵²) and A2PIC(Glu⁴⁴²-Lys⁴⁵²/Tyr⁴⁴⁵[SO₄]) to investigate the influence of the sulfated Tyr⁴⁴⁵ residue. The A2PIC(Glu⁴⁴²-Lys⁴⁵²/Tyr⁴⁴⁵[SO₄]) exhibits a very weak affinity with a K_a value of $\sim 3 \pm 1 \text{ mM}^{-1}$ and is in a comparable range to the non-sulfated peptide A2PIC(Glu⁴⁴²-Lys⁴⁵²) ($K_a = \sim 5 \pm 1 \text{ mM}^{-1}$).

K4 was used to confirm the ability of A2PIC(Glu⁴⁴²-Lys⁴⁵²) and A2PIC(Glu⁴⁴²-Lys⁴⁵²/Tyr⁴⁴⁵[SO₄]) to bind to the LBS of Pgn kringles. The K_a values of A2PIC(Glu⁴⁴²-Lys⁴⁵²/Tyr⁴⁴⁵[SO₄]) ($K_a = 29 \pm 4 \text{ mM}^{-1}$) and A2PIC(Glu⁴⁴²-Lys⁴⁵²) ($K_a = 24 \pm 4 \text{ mM}^{-1}$) are in a

Table 3 Association constants K_a (mM^{-1}) of wild-type and mutant rA2PIC peptides to various kringle constructs of Pgn

Ligand	rK1	K1–3	K4	rK4–5
6-AHA	75 ± 11^a	107 ± 11	46 ± 4	~ 52
Wild type	77 ± 4	176 ± 30	34 ± 2	18 ± 1
K406A			42 ± 6	
K415A			44 ± 6	
K422A		177 ± 22	33 ± 3	
K429A		170 ± 9	33 ± 3	
K436A	70 ± 1	26 ± 10	32 ± 2	16 ± 2
K452S	8 ± 0.5	17 ± 4	11 ± 2	1.4 ± 0.5
K452A	11.6 ± 0.5^a		9.1 ± 0.2^a	
K429/36A		17 ± 6	29 ± 0.5	16 ± 5
K429/52A	8 ± 0.5	30 ± 5		
K422/29/36A		22 ± 7	28 ± 1	18 ± 5
K415/22/29/36A		12 ± 2	24 ± 1	~ 17
K406/15/22/29/36A	53 ± 7	46 ± 12	23 ± 2	17 ± 4
K406/15/22/29/36/52A	~ 3	~ 4	~ 3	
A2PIC(E442-K452/Y445[SO ₄])			29 ± 4	
A2PIC(E442-K452)			24 ± 4	

^a From [18]

similar range (see Table 3). The quintuple mutant rA2PIC (Lys406Ala/Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala) ($K_a = 23 \pm 2 \text{ mM}^{-1}$) and the synthetic C-terminal peptide A2PIC(Glu⁴⁴²-Lys⁴⁵²) ($K_a = 24 \pm 4 \text{ mM}^{-1}$), both containing only the C-terminal Lys⁴⁵², show very similar binding affinities.

Discussion

Our investigations of wild-type and single, double, triple, quadruple, quintuple, and sextuple Lys to Ala mutants of the Pgn-binding region of A2PI (Asn³⁹⁸-Lys⁴⁵²), rA2PIC, via intrinsic fluorescence titrations, identify the C-terminal Lys⁴⁵² as the main binding partner in the interaction with the LBS of isolated K4 in solution, which is consistent with published data [8, 18]. We could show that all internal Lys residues (Lys⁴³⁶, Lys⁴²⁹, Lys⁴²², Lys⁴¹⁵ and Lys⁴⁰⁶) contribute only to a minor but distinct extent to the binding to the LBS of K4, which is demonstrated by a steady decrease of the K_a values with a decreasing number of Lys residues. We find similar results for isolated rK1, which is also consistent with published data [8, 18], and the double-kringle domain rK4–5 and the triple-kringle domain K1–3 in solution. The K_a value for wild-type rA2PIC and rK4–5 is smaller than for wild-type rA2PIC and K4, which could be due to a shielding effect of the rK5 domain in the flexible rK4–5 double-kringle domain or due to the N-terminal His-tag, which was still present. Due to the weak binding affinity of K5 to internal and C-terminal Lys residues [18, 54], we could not detect a significant change in the intrinsic fluorescence of isolated rK5 in solution. Frank

et al. [18] found that the K_a values for wild-type rA2PIC with isolated rK2, rK3mut (contains an active LBS in contrast to wild-type rK3, which is devoid of a functionally active LBS [45]) and K5 in solution are in the range of 3–4 mM^{-1} , which is indicative for a minor importance of these Pgn kringles in the interaction with wild-type rA2PIC. In contrast to these findings with isolated kringle domains, we identified Lys⁴⁵² and Lys⁴³⁶ as important binding partners for the interaction with the isolated triple-kringle domain K1–3 in solution. The association constant of K1–3 for wild-type rA2PIC is two- to threefold increased compared to rK1, and replacing of the conserved internal Lys⁴²² and Lys⁴²⁹ with Ala does not affect this enhanced binding affinity for K1–3. In addition, we could also demonstrate that the other internal Lys residues (Lys⁴¹⁵ and Lys⁴⁰⁶) do not contribute significantly in the binding to the LBS of K1–3 and that the quintuple mutant rA2PIC(Lys406Ala/Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala) exhibits a K_a value which is in the same range ($K_a = \sim 50 \text{ mM}^{-1}$) as for the interaction with isolated rK1. This enhanced binding affinity of K1–3 for wild-type rA2PIC points towards a cooperative effect of Lys⁴⁵² and Lys⁴³⁶, since it has been shown that the binding affinity of isolated rK2 in solution for Lys residues is very low [18, 44]. Our data underscore the fact that the Lys residues in the unique A2PIC mediate the binding to Pgn kringle domains [19–21, 31–34]. Wang et al. [36, 37] found in their kinetic studies with A2PI mutants that Lys⁴³⁶ is important for the interaction with plasmin in solution, immobilized plasminogen, or immobilized K1–3. They showed, in contrast to our findings, that Lys⁴⁵² is not involved in this non-covalent interaction. The question

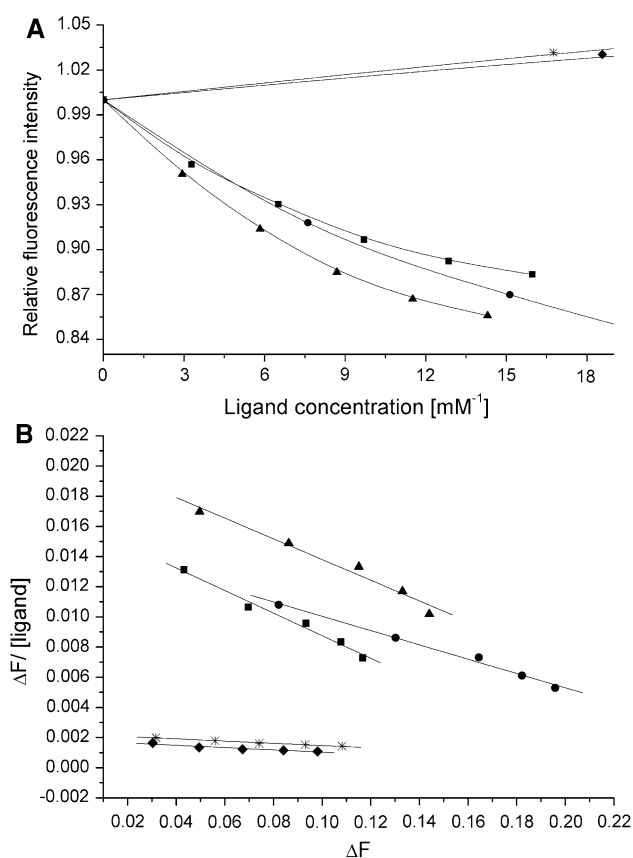


Fig. 2 **a** Averaged fluorescence titration of rK1 with wild-type rA2PIC (filled squares), single Lys to Ala mutants rA2PIC (Lys436Ala) (filled triangles), and rA2PIC(Lys452Ser) (star), double rA2PIC(Lys429Ala/Lys452Ala) (filled diamonds), and quintuple rA2PIC(Lys406Ala/Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala) (filled circles) mutants. The fluorescence intensity is normalized to 1 and the x-axis is enlarged. **b** Corresponding Scatchard plots and calculated regression curves

arises, why the C-terminal Lys⁴⁵² should be of no importance since it has been shown that the LBS of rK1 and K4 exhibit a much higher affinity for *N*^ε-acetyllysine, a ligand that mimics C-terminal lysines, than for *N*^ε-acetyllysine methyl ester, a mimic of internal Lys residues [18, 44, 52, 53]. Furthermore, another interesting fact supports the importance of Lys⁴⁵². As Frank et al. [18] have already discussed, the distance between the last two lysines (Lys⁴³⁶ and Lys⁴⁵²) in fully extended linear A2PIC is ~30 Å, which would match the distance required to bridge the LBSs of K1 and K2 as estimated from the K1–3 crystal structure [55], where the distance from Trp¹⁴⁴ (K1 LBS) to Trp²²⁵ (K2 LBS) is ~30 Å. This bridging seems to point towards an induced-fit model, where the binding of Lys⁴⁵² to the LBS of K1 is reinforced by the binding of Lys⁴³⁶ to the LBS of K2, which results in a conformational change of the K1–3 triple-kringle construct as a whole, rather than structural changes within domains, resulting in an enhanced binding compared to isolated rK1 or rK2 in

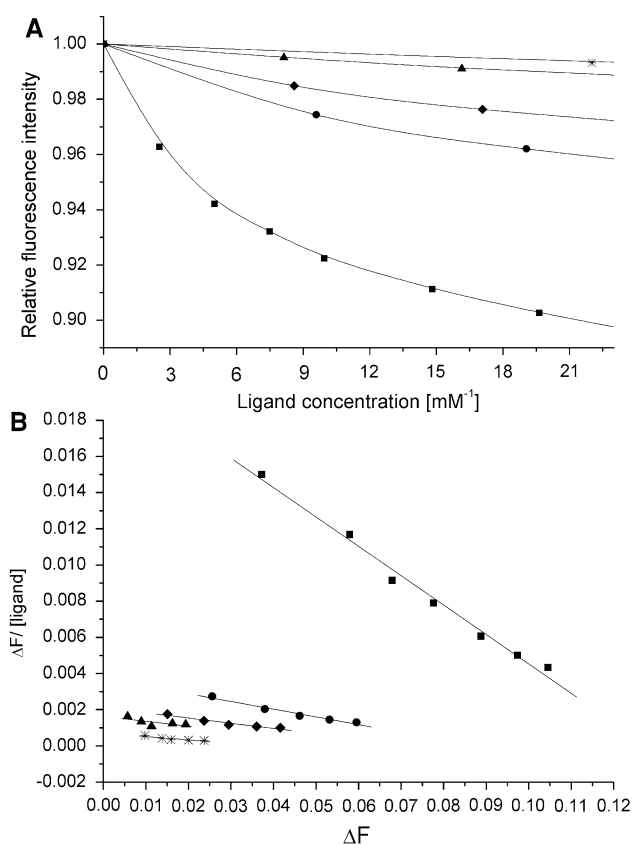


Fig. 3 **a** Averaged fluorescence titration of K1–3 with wild-type rA2PIC (filled squares), single Lys to Ala mutants rA2PIC (Lys436Ala) (filled triangles) and rA2PIC(Lys452Ser) (star), double rA2PIC(Lys429Ala/Lys452Ala) (filled diamonds), and quintuple rA2PIC(Lys406Ala/Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala) (filled circles) mutants. The fluorescence intensity is normalized to 1 and the x-axis is enlarged. **b** Corresponding Scatchard plots and calculated regression curves

solution. This hypothesis is supported by the fact that the multi-kringle domains are very flexible structures, since the conformational change upon activation of Pgn to Plm is quite dramatic (reviewed in [27]). Another fact derived from the X-ray crystal structure, supporting the importance of the Lys⁴⁵² and Lys⁴³⁶ residues, is that the C-terminal extension lies close to the RCL in such a way that the more distal conserved Lys residues (Lys⁴⁵² and Lys⁴³⁶) could readily engage the kringles in binding [22]. In addition, Sofian et al. [56] prepared three peptides, peptide 1 (Asn³⁹⁸-Leu⁴²¹), peptide 2 (Asp⁴¹⁶-Leu⁴³⁵), and peptide 3 (Leu⁴³⁰-Lys⁴⁵²), and the corresponding antibodies to these different regions of A2PIC, and investigated their effects on plasmin–antiplasmin kinetics and fibrinolysis. They could show that peptide 3 and antibody 3 exhibit a ~10-fold decrease of the rate of association between A2PI and Plm, which underscores the importance of Lys⁴⁵² and Lys⁴³⁶, whereas peptides 1 and 2 and antibodies 1 and 2, corresponding to the remaining internal Lys residues

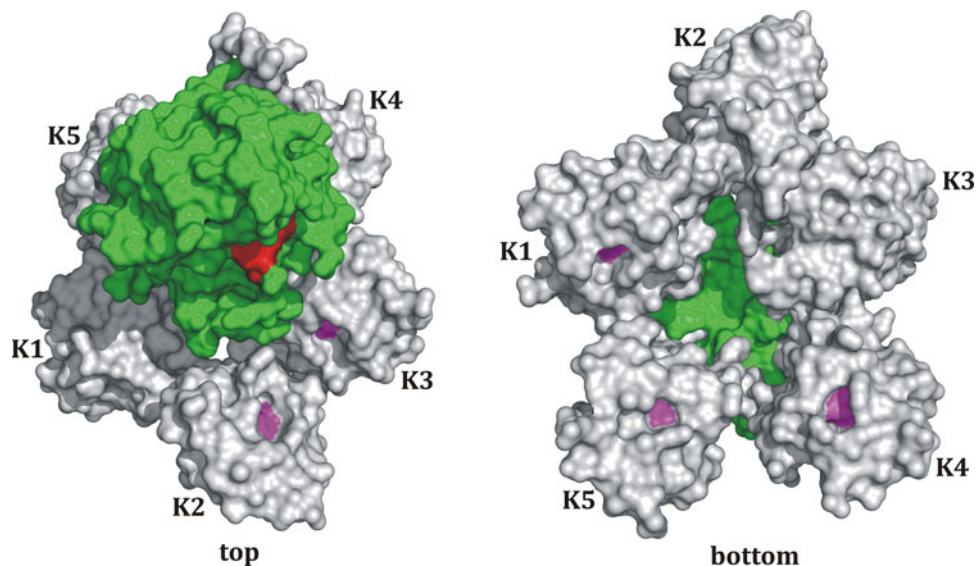


Fig. 4 Model structure of the whole Pgn molecule. The model is based on the amino acid sequence of Pgn, which was used as a template for the automated structure modeling using SWISS-MODEL [57–59]. Connected and overlapping 3D structures of Pgn fragments (K1–K2; Leu⁸¹-Thr²⁴⁵, parts of K3–K4; Thr²⁹¹-Ser⁴⁴¹, parts of K4–K5; Leu⁴⁰²-Ala⁵⁴³, K1–K2–K3; Leu⁸¹-Cys³³³, and K5; Asp⁴⁶¹-Pro⁵⁴⁴)

(Lys⁴²⁹/Lys⁴²² and Lys⁴¹⁵/Lys⁴⁰⁶, respectively), seemed to have no significant effect on the serpin inhibition reaction. Nevertheless, it might be possible that the C-terminal segment adopts different conformations in the whole A2PI or as peptide in solution. These conformational differences could result in different binding affinities for kringle domains. However, the structure of the unique A2PIC remains unsolved [22], leaving the question of different conformations open. In addition, we have expressed recombinant A2PI and Lys to Ala mutants in the C-terminal segment of A2PI at high levels in *E. coli* [51]. No binding affinities could be determined due to extremely low solubility of the recombinant A2PI constructs at physiological conditions (pH and salts). Furthermore, Frank et al. [18] demonstrated by NMR and CD measurements that rA2PIC is a rather flexible polypeptide of low β -structure supporting the discussed induced-fit model.

We also investigated the effect of sulfated Tyr⁴⁴⁵, which is present in the unique C-terminal extension of native human A2PI [15]. A possible function of this posttranslational modification could be an interaction with Pgn K3, which contains no active LBS [45] but a heparin-binding site (Prof. Dr. M. Llinás, personal communication). It is therefore possible that sulfated Thy⁴⁴⁵ in A2PIC might interact with the heparin-binding site in K3. We applied intrinsic fluorescence titrations to investigate the effect of two synthetic peptides A2PIC(Glu⁴⁴²-Lys⁴⁵²) and A2PIC(Glu⁴⁴²-Lys⁴⁵²/Tyr⁴⁴⁵[SO₄]) on the interaction with isolated Pgn rK3 in solution. The K_a values of $\sim 3 \pm 1 \text{ mM}^{-1}$

were selected and thereafter assembled and attached to the 3D structure of the catalytic domain of Plm (1BML) using the visualization and rendering software PyMOL. Each kringle domain (*K*) is shown in *gray*, the central Trp of each LBS is presented in *pink*, the protease domain is colored in *green*, and the active site residues are marked in *red*

for the sulfated and $\sim 5 \pm 1 \text{ mM}^{-1}$ for the non-sulfated A2PIC(Glu⁴⁴²-Lys⁴⁵²) revealed that sulfated Tyr⁴⁴⁵ has no detectable effect on the interaction of rK3 and A2PIC(Glu⁴⁴²-Lys⁴⁵²/Tyr⁴⁴⁵[SO₄]) in solution. To ensure that the two synthetic peptides are capable of binding to the LBS in kringle domains, we investigated the binding of A2PIC(Glu⁴⁴²-Lys⁴⁵²) and A2PIC(Glu⁴⁴²-Lys⁴⁵²/Tyr⁴⁴⁵[SO₄]) to isolated Pgn K4 in solution. The achieved K_a values of $29 \pm 4 \text{ mM}^{-1}$ for the sulfated and $24 \pm 4 \text{ mM}^{-1}$ for the non-sulfated A2PIC(Glu⁴⁴²-Lys⁴⁵²) are both in a similar range. An interesting finding is that the K_a value of A2PIC(Glu⁴⁴²-Lys⁴⁵²) is in the same range as the quintuple mutant rA2PIC(Lys406Ala/Lys415Ala/Lys422Ala/Lys429Ala/Lys436Ala), which also contains only the C-terminal Lys⁴⁵².

We further attempted to model the 3D structure of the whole Pgn. The model is based on the amino acid sequence of Pgn, which was used as a template for the automated structure modeling using SWISS-MODEL [57–59]. Connected and overlapping 3D structures of Pgn fragments (K1–K2; Leu⁸¹-Thr²⁴⁵, parts of K3–K4; Thr²⁹¹-Ser⁴⁴¹, parts of K4–K5; Leu⁴⁰²-Ala⁵⁴³, K1–K2–K3; Leu⁸¹-Cys³³³, and K5; Asp⁴⁶¹-Pro⁵⁴⁴) were selected and thereafter assembled and attached to the 3D structure of the catalytic domain of Plm (1BML) using the visualization and rendering software PyMOL. The obtained model of the whole plasmin(ogen) molecule is shown in Fig. 4. The 3D structure model exhibits a spiral-type shape, which resembles the known shape of Glu¹-Pgn visualized by

electron microscopy [60, 61]. An interesting finding is that the LBS of K1, K4, and K5 are orientated opposite the active site of the protease domain whereas the LBS of K2 and K3 are orientated towards it. The opposite orientation of the LBS of K1 and K4 would allow Plm to bind to the target surface and leave the active site in the protease domain freely accessible. The model supports the findings that the K1- and K4-LBS act as independent and complementary domains both able to support Pgn binding and activation, and both seem to be required to ensure Pgn anchorage and subsequent proteolytic activity [62]. The remaining kringles seem to be involved in the modulation of Plm activity rather than anchorage of Pgn to the target surface since it has been shown that K5 interacts with long-chain fatty acids [63] as well as various cell surface proteins [64–66] and that K3 exhibits a heparin-binding site (Prof. Dr. M. Llinás, personal communication). The model is also consistent with our findings, that isolated rK1 and K4 exhibit a strong affinity for C-terminal Lys⁴⁵², which supports the independence of both domains [62], and that the reinforced interaction between C-terminal Lys⁴⁵² and internal Lys⁴³⁶ in the interaction with the isolated K1–3 domain (two- to threefold increase compared to rK1) underscores the high flexibility of multi-kringle domains (reviewed in [27]) allowing Lys⁴³⁶ to interact with the LBS of K2.

Conclusions

We could identify the C-terminal Lys⁴⁵² as the main binding partner in the interaction of rA2PIC with isolated rK1, K4, and rK4–5 in solution. We could show a cooperative, zipper-like enhancement of the interaction between Lys⁴⁵² and internal Lys⁴³⁶ of the rA2PIC and the isolated K1–3 triple-kringle domain in solution. The data allow the assumption that the C-terminal Lys⁴⁵² is the primary binding partner whereas Lys⁴³⁶ functions as enhancer rather than as the primary binding partner. A co-crystallization of rA2PIC and Pgn or the K1–5 multi-kringle domain would definitively solve the interaction mechanism between the kringle domains and A2PIC. Our findings point towards a major interaction of the C-terminal Lys⁴⁵² of rA2PIC with both LBS in K1 and K4, which act as independent and complementary domains both required to ensure Pgn anchorage and subsequent proteolytic activity, and that the remaining kringles (K2, K3, and K5) seem to be involved in the modulation of Plm activity rather than anchorage of Pgn to the target surface. The function of sulfated Tyr⁴⁴⁵ still remains to be solved.

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