Data Article

Structural basis of specific inhibition of tissue-type plasminogen activator by plasminogen activators inhibitor-1

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Abstract

Thrombosis is a leading cause of death worldwide [1]. Recombinant tissue-type plasminogen activator (tPA) is the FDA-approved thrombolytic drug for ischemic strokes, myocardial infarction and pulmonary embolism. tPA is a multi-domain serine protease of the trypsin-family [2] and catalyses the critical step in fibrinolysis [3], converting the zymogen plasminogen to the active serine protease plasmin, which degrades the fibrin network of thrombi and blood clots. tPA is rapidly inactivated by endogenous plasminogen activators inhibitor-1 (PAI-1) [4] (Fig. 1). Engineering on tPA to reduce its inhibition by PAI-1 without compromising its thrombolytic effect is a continuous effort [5]. Tenecteplase (TNK-tPA) is a newer generation of tPA variant showing slower inhibition by PAI-1 [6]. Extensive studies to understand the molecular interactions between tPA and PAI-1 have been carried out [7–18], however, the precise details at atomic resolution remain unknown. We report the crystal structure of tPA·PAI-1 complex here. The methods required to achieve these data include: (1) recombinant expression and purification of a PAI-1 variant (14-1B) containing four mutations (N150H, K154T, Q319L, and M354I), and a tPA serine protease domain (tPA-SPD) variant with three mutations (C122A, N173Q, and S195A, in the chymotrypsin numbering) [19]; (2) formation of a tPA-SPD·PAI-1 Michaëlis

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complex in vitro [19]; and (3) solving the three-dimensional structure for this complex by X-ray crystallography [deposited in the PDB database as 5BRR]. The data explain the specificity of PAI-1 for tPA and uPA [19,20], and provide structural basis to design newer generation of PAI-1-resistant tPA variants as thrombolytic agents [19]. © 2016 Elsevier Inc.. Published by Elsevier Inc. This is an open access article under the CC BY license (http://creativecommons.org/licenses/by/4.0/).

Specifications table

| Subject area                      | Biology                                                                 |
|----------------------------------|-------------------------------------------------------------------------|
| More specific subject area       | Protein structure and biochemistry                                      |
| Type of data                     | X-ray crystal structure, Mass spectrometry                             |
| How data was acquired            | X-ray diffraction data were collected at Shanghai Synchrotron Radiation Facility. Mass spectra of MALDI-TOF-MS were obtained on a Bruker REFLEX III MALDI-TOF-MS (Bruker-Franzen, Bremen, Germany). |
| Data format                      | Processed                                                              |
| Experimental factors             | Recombinant proteins were purified to high homogeneity before use.    |
| Experimental features            | The structure of the tPA·PAI-1 complex was determined by X-ray crystallography. |
| Data source location             | City, Country and/or Latitude & Longitude (& GPS coordinates) for collected samples/data if applicable |
| Data accessibility               | The data is available from the related publication by Gong et al. (http://www.ncbi.nlm.nih.gov/pubmed/26324706), and the structure deposited in the Protein Data Bank (entry 5BRR). |

Value of the data

- Determines the crystal structure of the Michaëlis complex between tPA and PAI-1.
- Provides insight on the specificity of PAI-1 for tPA and uPA.
- Identifies key residues of tPA for binding to PAI-1.
- Explains the PAI-1-resisting property of Tenecteplase.
- Offers important clues to design newer generation of PAI-1-resistant tPA variants.

1. Data, experimental design, materials and methods

1.1. Data and experimental design

We have determined the structure of tPA·PAI-1 Michaëlis complex and identified key residues of tPA for binding to PAI-1 by X-ray crystallography, and the data are summarized in the original publication [19].

We expressed the recombinant PAI-1 variant 14-1B (N150H, K154T, Q319L, and M354I) [21], using the expression vector pT7-PL and BL21 cells as soluble protein [22]. The choice of this particular variant is to obtain PAI-1 in active form, advantageous for crystallization, because the wild type PAI-1
has a half life of only 2 h and has propensity to spontaneously convert into an inactive, so-called latent form, and to aggregate at high concentration [23,24].

PAI-1 inhibits tPA by a suicide-substrate mechanism common to all SERPIN members [23,25] – see Fig. 1A in the original publication [19]. In this SERPIN mechanism, a long flexible loop of PAI-1 (reaction center loop, or RCL) inserts into the active site of tPA to form a transient Michaëlis complex. The RCL is cleaved by tPA through the classical serine proteolytic mechanism. tPA forms a covalent acyl-enzyme intermediate with PAI-1 by cleaving the scissible bond of PAI-1 RCL, following the Michaëlis complex. Before the hydrolysis of this acyl-enzyme intermediate, the PAI-1 RCL undergoes major conformational changes and inserts itself into the PAI-1 β-sheet A. At the same time, the tPA in the intermediate is pulled to the other side of PAI-1, distorted, and deactivated before the hydrolysis of the acyl-enzyme intermediate can take place.

Human tPA contains a fibronectin type II domain (amino acids 1–50), a growth factor domain (amino acids 51–91), two kringle domains (amino acids 92–261), an interdomain linker (amino acids 262–275) and a serine protease domain (SPD, amino acids 276–527) [2] – see Fig. 1B in the original publication [19]. The tPA-SPD is the catalytic domain responsible for the plasminogen activation and is inhibited by PAI-1. Thus, we used only the recombinant tPA-SPD domain to form the Michaëlis complex with PAI-1. We generated three mutations in tPA-SPD: S478A (or S195A in the chymotrypsin numbering) to render the tPA-SPD catalytically inactive, so the Michaëlis complex does not proceed to a stable, covalent complex; N448Q (or N173Q in the chymotrypsin numbering) to remove the glycosylation on tPA-SPD, increasing the homogeneity of the recombinant protein and facilitating
protein crystallization; and C395A (or C122A in the chymotrypsin numbering that will be used throughout the rest of text) mutation to remove the disul
dide bond linked to K2 domain – see Fig. 1B in the original publication [19]. The recombinant tPA-SPD mutant was expressed in P. pastoris and confirmed by SDS-PAGE and mass spectrometry after trypsin digestion (Table 1).

The recombinant PAI-1 14-1B and tPA-SPD were respectively dialysed into a high-concentration salt (1 M NaCl) and low pH (20 mM Mes pH 6.1) buffer before the Michaëlis complex formation. This condition is required to stabilize PAI-1 at its active form. Subsequently, these two proteins in high salt concentrations and low pH buffer were mixed in a 1:1 M ratio, followed by a dialysis into a low-concentration salt (150 mM NaCl) and neutral pH (20 mM Tris–HCl pH 7.4) buffer. This dialysing step ensures the complex formation similar to that in physiologic condition. A further gel filtration chromatography purification yielded a complex of greater than 99% purity.

2. Materials and methods

2.1. Recombinant protein production

The recombinant PAI-1 mutant 14-1B [21] containing four point mutations (N150H, K154T, Q319L, and M354I), and a hexa-His-tag was expressed in E. coli, using the expression vector pT7-PL and BL21 cells as previously described [22]. The recombinant tPA-SPD was expressed in P. pastoris X-33. This strain facilitates the formation of five disulfide bonds (C42-C58, C50-C111, C100-C182, C136-C201, C168-C182 in chymotrypsin numbering) in tPA-SPD with a high yield about 50 mg recombinant protein per liter medium – see in the original publication [19].

2.2. The peptide mass fingerprinting of tPA-SPD by MALDI-TOF mass spectrometry

The SDS-PAGE was performed using 15% polyacrylamide gels. Following SDS-PAGE, the gels were stained with 0.1% (w/v) Coomassie brilliant blue R-250 in 25% (v/v) ethanol and 10% (v/v) acetic acid. The gel digestion was performed using a modified version of previously published protocol [26]. Briefly, the gel band containing 100 ng tPA-SPD was excised from the 15% two-dimensional SDS-PAGE gel, cut in pieces, and destained by washing with 50% (v/v) acetonitrile in 100 μl of 25 mM NH₄HCO₃ for 30 min at room temperature. The gel pieces were then dried in a SpeedVac Vacuum (Savant Instruments, Holbrook, NY, USA) and rehydrated at 4 °C for 15 min in 3–5 μl digestion solution (25 mM NH₄HCO₃ and 12.5 ng/μl modified sequence-grade trypsin). Then 3–5 μl of digestion solution without trypsin was added to keep the gel pieces wet during the digestion. After overnight incubation at 37 °C, the digestion was stopped with 5% trifluoroacetic acid (TFA) for 20 min. The peptides were extracted by 20 μl of 5% TFA for 1 h at 37 °C and then by 20 μl of 2.5% TFA/50% acetonitrile for 1 h at 37 °C. The combined supernatants were evaporated in the SpeedVac Vacuum and dissolved in 4 μl 0.5% aqueous TFA for MS analysis.

All mass spectra of MALDI-TOF-MS were obtained on a Bruker REFLEX III MALDI-TOF-MS (Bruker-Franzen, Bremen, Germany) in positive ion mode at an accelerating voltage of 20 kV with the matrix of α-cyano-4-hydroxy cinnamic acid. The spectra were internally calibrated using trypsin autolysis products. The peptide mass fingerprinting obtained was used to search through the SWISS-PROT and NCBI database by the Mascot search engine (http://mascot.proteomics.com.cn/) with a tolerance of ~±0.3 D and one missed cleavage site.

| Mr observed (Da) | Mr calculated (Da) | Peptide sequence |
|-----------------|-------------------|-----------------|
| 1387.1          | 1386.8004         | $^{53}$FPHHLTLVGLGR$^{64}$ |
| 1335.8          | 1335.6328         | $^{142}$HEALSPFYSTER$^{152}$ |
| 1179.4          | 1179.6157         | $^{238}$TNLYLDWIR$^{247}$ |
| 878.8           | 878.4618          | $^{231}$DPGVYTK$^{238}$ |
| 722.4           | 722.3831          | $^{160}$LYPSSR$^{165}$ |
2.3. X-ray crystallography

The tPA-SPD·PAI-1 Michaëlis complex was formed by mixing tPA-SPD and PAI-1 in a 1:1 M ratio at low concentration (∼ 0.5 mg/ml), followed by dialysis into 20 mM Tris–HCl pH 7.4, and 150 mM NaCl, concentration to 0.5 ml volume for a further gel filtration chromatography purification, which yielded to a complex of greater than 99% purity. The purified complex was then concentrated to 10 mg/mL before setting up crystallization trials. Crystals of the tPA-SPD·PAI-1 Michaëlis complex were grown at 20 °C with the sitting drop method by mixing equal volumes of protein solution and precipitant solution (8% PEG-6000 and 0.1 M Tris pH 7.4), and appeared quickly within one day. However, the crystals always appeared as very thin plates, and decayed rapidly in the X-ray beam, posing great difficulty for X-ray data collection. Most crystals diffracted to only 4–5 Å at Shanghai Synchrotron Radiation Facility (SSRF) BL-17U beam line, and the diffracting spots often appeared as elongated or splitted shapes. After many crystallization and data collection trials for one and half years, one 3.16 Å data set was finally obtained at SSRF beam line BL17U using 25% glycerol as cryoprotectant at a wavelength of 0.979 Å. The data were processed and scaled using the HKL2000 program package [27]. The crystal belongs to P2₁2₁2₁ space group with one complex in the crystallographic asymmetric unit. The structure of the tPA-SPD·PAI-1 Michaëlis complex was solved by molecular replacement method using MolRep program [28], which gave very strong and unambiguous solutions. A tPA-SPD molecule was first positioned inside the crystal lattice using the structure of the tPA-SPD catalytic domain (PDB code 1A5H) [29] as a searching model and all the X-ray data up to 3.3 Å. The molecular replacement using MolRep gave a contrast of 12.33, a signal to sigma ratio for translational function of 16.02, and a correlation coefficient of 0.365. Next, the position of PAI-1 was searched using the model of active stable variant of PAI-1 (Protein Data Bank code 1DVM) [30] while fixing the already positioned tPA-SPD model, giving only one translational function with a signal to sigma ratio of 19.4, and a correlation coefficient of 0.538. The molecular replacement model was subjected to iterative refinement and manual model rebuilding using Refmac [31] and Coot [32], respectively, giving a final R factor and Rfree factor of 0.20 and 0.27, respectively. The structure was validated with PROCHECK [33] and analyzed by PyMOL [34] and PISA [35]. The final refined crystal structure of tPA-SPD·PAI-1 Michaëlis complex was deposited in PDB with the code 5BRR.

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Appendix A. Supplementary material

Supplementary data associated with this article can be found in the online version at http://dx.doi.org/10.1016/j.dib.2015.12.050.

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