Understanding Russell’s viper venom factor V activator’s substrate specificity by surface plasmon resonance and in-silico studies

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Abstract

Blood coagulation factor V (FV) is activated either by Factor X or thrombin, cleaving at three different sites viz., Site I (Arg709-Ser710), site II (Arg1018-Thr1019), and site III (Arg1545-Ser1546). Russell’s viper venom factor V activator (RVV-V) is a thrombin-like serine proteinase that activates FV with selective, single cleavage at site III. A long lasting effort is being pending in understanding the ‘selective’ binding specificity of the RVV-V towards site III. Here, we present the binding kinetic study of RVV-V with two designed peptides corresponding to the regions from site I (Gln699—Asn713) and site II (1008Lys—Pro1022), respectively, that include 15 amino acids. Our investigation for justifying the binding efficacy and kinetics of peptides includes SPR method, protein-peptide docking, molecular dynamics simulation, and principal component analysis (PCA). Surprisingly, the SPR experiment disclosed that the Peptide II showed a lower binding affinity with \( K_D \) of 2.775 mM while the Peptide I showed none. Docking and simulation of both the peptides with RVV-V engaged either rooted or shallow binding for Peptide II and Peptide I respectively. The peptide binding resulted in global conformational changes in the native fold of RVV-V, whereas the similar studies for thrombin failed to make major changes in the native fold. In support, the PCA analysis for RVV-V showed the dislocation of catalytic triad upon binding both the peptides. Hence, RVV-V, a serine protease, is incompetent in cleaving these two sites. This study suggests a transition in RVV-V from the native rigid to the distorted flexible structure and paves a way to design a new peptide substrate/inhibitor.

Introduction

According to World Health Organization, snake envenomation causes the highest deaths in India (about 11000 deaths) every year. There are 200 out of 600 venomous snakes, which are lethal, and of medical importance [1, 2]. Snake venoms (Crotalidae and Viperidae) are the rich sources of snake venom serine proteases (SVSPs) with varying molecular specificity [3, 4]. The SVSPs are both pro and anticoagulant in nature and use conserved serine protease
The SVSP is found to interfere with the human haemostasis system upon envenomation. The Russell’s viper Venom contains RVV-V a thrombin like SVSP which activates blood coagulation Factor V (FV), one of the crucial components of the blood coagulation cascade [8]. RVV-V serve as an extremely useful tool in haemostasis [9, 10].

Factor V (FV) is a well-studied large protein, consists of six domains, involved in the blood coagulation cascade. The FV is present as an inactive precursor in the blood; it requires activation by cleaving at specific sites by proteases, such as thrombin and FXa. Both the proteins cleave the FV at three different sites of FV viz., site I (Arg709–Ser710), site II (Arg1018–Thr1019), and site III (Arg1545–Ser1546). Highly glycosylated B domain of FV is removed by these proteases to achieve an active form of FV. The FV is also activated by RVV-V by cleaving the site III and does not cleave the other two thrombin-susceptible sites [11]. The thrombin is active against several molecular substrate including the FV but no protein or molecular substrate is reported so far for RVV-V other than the FV [12].

In general, serine proteases are robust and highly sensitive for the cleavage sites; they contain catalytic triad or diad in the active site. Even though they possess a distinct structure they converge at the catalytic active site e.g., chymotrypsin and subtilisin [13]. Both these proteins are active against different molecular substrates. The molecular structure and the catalytic triad arrangement of the RVV-V are analogous with that of the thrombin. Yet the RVV-V exhibits no effect on either the structure or the activity even after a prolonged incubation with fibrinogen, prothrombin, factor VIII, and FX [14, 15]. In addition, the RVV-V has been showed to possess a weak amidase activity towards the thrombin chromogenic substrate S2238 (D-Phenyl-L-pipercolyl-L-arginyl-p-nitroanilide). Similarly, no response against the serine protease inhibitor (BPTI, Antithrombin, EDTA) is observed [16, 17].

The crystal structure of the RVV-V with a substrate fragment of the site III of FV was solved by Nakayama et al. [18] wherein the peptide is cleaved and the left over remains intact on the active site. The essential molecular interaction between the protease and the substrate is observed. The narrow specificity of RVV-V has been addressed without a fulfillment. The crystal structure explains multiple interaction of the peptide fragment; and it suggested that the missing hydrophobic environment in the other two susceptible sites might cause the selective specificity of the RVV-V.

There are several similar protease/substrate/inhibitor complex structures reported [19–26]. The mode of binding of either the substrate or the inhibitor is essentially same, yet one molecule is digested whereas the other is not. Even then, the molecule, which found to be an inhibitor, does not affect the native fold of the binding protein. Here we report the molecular interaction of the RVV-V with Site I and II of factor V whose binding disrupts the native fold of the RVV-V.

In order to justify the selective binding of RVV-V we designed the peptides of 15-residue length in the regions of Site I and II of the FV. Peptide I (699QNGLAAALGIRSFGN713) and Peptide II (1008KHTHHAPLSPRTFHP1022) correspond to the region of the FV from the cleavage site I and II respectively. Both the peptides were used for SPR (Surface Plasmon Resonance) analysis with the purified RVV-V and modeled for docking and Molecular Dynamics study. Since, the thrombin is known for cleaving the FV, the designed peptides were docked, and the dynamics was performed with the existing structure of the thrombin for comparison.

**Material and methods**

**Purification of RVV-V**

The Russell viper venom was purchased from Irula Snake-Catchers Industrial Cooperative Society (ISCICS), Tamil Nadu, India. The RVV-V was purified as described by Shiffman et al.,
with some changes [14]. Lyophilized Crude venom (100 mg) was dissolved in 1 ml of 50 mM
Sodium Acetate buffer (pH 5.5). The solution was centrifuged at 12,000 rpm for 10 minutes at
4˚C and the supernatant was loaded to Sephadex S-100 column (FPLC). Peaks containing the
RVV-V were pooled and loaded for CM Sepharose FF Cation Exchange (Pharmacia) column
(equilibrated with 50 mM sodium acetate buffer pH 5.5). Proteins were eluted with 50mM Na-
acetate buffer with gradient 0 to 1 M Sodium Chloride. The fractions of RVV-V were pooled,
dialyzed in 50 mM sodium-acetate buffer, and concentrated. Further, the binding kinetics was
performed for RVV-V with both the peptides (purchased from Genei™) by SPR method.

Synthesis of Peptide I and II and model building

We designed 15-residue peptides (Fig 1) including a cleavage site, allowing few residues on
both sides of P₀ to be present to ensure proper penetration of the peptide into the cavity of the
binding pocket. Allowing the residues on both sides of the P₀ site will make more interaction
on both the ends; resulting in a proper hold and exposure of P₀ site for the cleavage by the
enzyme. It is observed that in most of the protein substrate/inhibitor complex structures more
number of interactions is found in either side of the P₀ site; accordingly, it may act on either
the substrate or the inhibitor. A model for the Peptide I and II was designed using the server
CABS-dock, which builds the model as well as performed docking with the protein.

Study of kinetics of RVV-V using SPR

The SPR is a well-known primary protein-ligand screening method for the measurement of
a real-time, label-free detection of biomolecular interactions like protein-peptide, antibody-
antigen, and protein-DNA interactions. A small amount of protein is sufficient and the affinity
data are more accurate as the recorded Sensorgram is time-dependent and a continuous flow
system [27]. In the experiment the ligand is saturated with an immobilized protein followed by
washing, and the association constant (Kₐ), and the dissociation constant (K₅) are obtained
from the binding response data [28, 29]. Biacore T200 optical biosensor was used to get
the interaction kinetics of RVV-V with the peptides. The purified RVV-V (concentration of
50 μg/ml in 50 mM sodium acetate buffer, pH 5.5) was immobilized on CM5 chip by amine-
coupling method. A constant flow rate of 10μl/min was used to activate the surface of the
flow cell for 7 min using a 1:1 mixture of 100 mM N-hydroxysuccinimide (NHS) and 100 mM
N-ethyl-N-(dimethylaminopropyl)-carbodiimide (EDC) dissolved in water. The RVV-V was
injected for 7 min, and the non-activated carboxy methyl groups on the surface were blocked
by 7 min injection of 1 M ethanolamine (pH 8.5) and totally 3000 RU of the RVV-V were
immobilized on the chip.
Biacore control software was used for obtaining all data by monitoring the change in the refractive index as a function of time at a flow rate of 30 \( \mu \)l/min. A relative quantity of peptide bound to RVV-V was quantified by calculating the amount of net increase in the refractive index over a period as compared to the initial buffer (PBS) response. The inline subtraction is performed during each run. Before changing each peptide and the peptide-concentration, the surface of the sensor chips was washed by running the buffer (PBS) and the chip was regenerated by injecting 10 mM glycine pH 2.5 for 30 sec. The association and the dissociation times are 120 and 300 sec respectively. For both the peptides, the experiment was performed by changing the concentration of the peptides ranging from 62.5 \( \mu \)M to 1 mM. The results were analyzed by T200 evaluation software version 2.0 and a two-state binding model fit the data.

**Docking and MD simulation**

The structure of RVV-V was retrieved from protein data bank (PDB ID: 3S9C) and water molecules, heteroatoms, and small Peptides were removed and then it was processed for further molecular docking and MD analysis. The interaction of RVV-V with cleavage sites I and II has yet not been reported and accordingly the blind docking was performed for Peptide I and Peptide II using CABS-Dock. The CABS-Dock server models the peptide and performs the simulation search for the binding site, permitting the flexibility of the peptide and the slight fluctuations in the receptor backbone \[30, 31\]. The best-docked poses were refined and subjected for MD simulation analysis.

The stability of the docked complexes was analyzed by molecular dynamic simulation using Gromacs 5.0 package on work station with pentium i7 octacore. The topology of protein was generated using “gromos96 54a7” force field. A cubic box was created (edges 1.2 nm away from protein surface) and was solvated with the water model (SPC216). The total charge was neutralized for the system by adding counterions. The energy minimization for the protein complex was achieved using the steep-descent followed by the conjugate method for 50,000 steps with a tolerance of 1000 kJ mol\(^{-1}\) nm\(^{-1}\). The treatment of long range electrostatic forces was utilized by PME (particle mesh ewald) with a cut-off 1 nm \[32\]. LINCS algorithm was performed for constrains of all the bonds. To equilibrate temperature, pressure, density, and total energy of the system, the position restrain MD for 100 ps was carried out for NVT and NPT ensemble. To regulate temperature (310K) and pressure (1 atm), V-rescale and Parrinello-Rahman algorithms were employed respectively. After the equilibration, the system was subjected to MD simulation production and the coordinates of trajectory were saved at every 2ps. The Gromacs analysis package was utilized to analyze the trajectory and the xmgrace for plotting the results. A similar docking and MD simulation was performed for the Peptide III (1535DPDNIAAWYLRSNNG1551) with RVV-V in order to conform the authentic binding of this region with RVV-V.

Both the peptides viz., site I and site II, were the susceptible cleavage sites for the thrombin as well. To compare the docked and the simulation results of RVV-V, the peptide models obtained from CABS-dock were docked with the thrombin using HADDOCK server, which returned eight clusters for Peptide I and four clusters for Peptide II with four pdb files for each. The best-docked pose, for each peptide was chosen for MD analysis with the same MD parameters for 100ns.

**Dihedral PCA (dPCA) analysis**

Dihedral PCA (principal component analysis) was used to describe the high-amplitude concerted motion from the MD trajectories of protein based on eigenvectors calculated using covariance matrix \[33, 34\]. The dihedral angles of protein were used to define the atomic
fluctuation throughout the MD simulation, which is described by the cosine values of the PC of covariance matrix. The cosine values are used to check whether the trajectory has ensembled enough to show the free energy landscape obtained from the PCA analysis [35, 36]. The range of cosine value from 0 to 1 in the total time of MD simulation (T) is given by,

\[ C_i = \frac{2}{T} \left( \int_0^T \cos \left( \frac{\pi t}{T} \right) p_i(t) \, dt \right)^2 \left( \int_0^T p_i^2(t) \, dt \right)^{-1} \]

Where \( p_i(t) \) is the \( i \)th PC’s value.

Therefore, we can measure absolute and sensitive measurements of trajectory by getting numerous free-energy minima, which relates to conformations mapping with their respective energy basins as available in the free energy landscape of the selected PCs. Generally, the first few PCs contributions define the protein nature. However, in the most of the eigenvector cosine, the values are near one, which is the result of a large-scale motion in the protein dynamics and hence are not used [37, 38].

**Result and discussion**

**SPR result**

In order to know the binding affinity of RVV-V with both the peptides, the binding kinetics was performed using the SPR method. Different concentrations of peptides were prepared for the SPR analysis and the response plot was obtained. The binding responses for the Peptide II were obtained, fitted with a two-state interaction curve with \( K_D \) value of 2.775 mM (Table 1) (Fig 2) whereas the Peptide I showed no response. The result shows that the Peptide II has affinity with RVV-V. The two-state fit for the Peptide II shows conformational changes, which is evident from the given Table 1. The kinetic rate constants show that the first reaction is faster (\( k_{a1} = 760.4/\text{Ms} \)) than the second (\( k_{a2} = 2.809 \times 10^{-4}/\text{s} \)). Hence, at the beginning an
unstable (kd1 = 2.414/s) RVV-V/Peptide II complex is quickly formed. This complex undergoes a slow (ka2 = 2.809x10^{-4}/s) conformational change into a more stable complex (kd2 = 1.942x10^{-3}/s).

It is evident from the SPR results that the Peptide I shows no response and the Peptide II shows a weak binding affinity towards the RVV-V. The affinity for Peptide II has to be established and the behaviour of the Peptide I to be judged. We performed model building and docking of both the peptides with the RVV-V using CABS-DOCK server to establish the binding behaviour and the affinity.

**Peptide docking with RVV-V using CABS dock**

For each peptide, CABS-dock result was collected (10 docked poses for each peptide, which were obtained from 10-cluster ranked based on the cluster density). The best-docked pose for Peptide I and Peptide II is shown in Figs 3 and 4 respectively and their interaction details are

![Docked pose of Peptide I with RVV-V](https://doi.org/10.1371/journal.pone.0181216.g003)
given in Tables 2 and 3. Both the peptides bind the RVV-V such that the Arginine at position $P_0$ falls in subsite $S_0$, which is formed by two loops (220-Loop and 192-Loop) and the residues in $P_{-2}$ position occupies the subsite $S_{-1}$ formed in the middle of the three loops (220-Loop, 174-Loop, and 97-Loop). Whereas the $P_2$ and the $P_3$ residues of Peptide I lie in subsite $S_1$ (surrounded by 192-Loop, 149-Loop, and 36-Loop). The Docked pose of Peptide I with RVV-V (Complex R1) and the RVV-V with Peptide II (Complex R2) show comparable interactions. However, the Complex R2 seems to have more number of weaker interactions, which may add strength to the overall binding of Peptide II to RVV-V. The MD simulation was performed for both the peptide complexes in order to assess the stability and dynamic behaviour of the RVV-V with these peptides.

Usually the substrates bind in the serine proteases in such a way that the susceptible cleavage site $P_0$ comes in a close proximity of the catalytic triad. The NH1 and the NH2 of $P_0$ may
interact with Asp 189, which would anchor the residue, and the transfer of electron in oxy-
anion hole will proceed to further the hydrolysis of the peptide bond. The Peptides I and II with RVV-V after docking do not seem to have any such instances; in the case of Peptide I, the Arg709 is oriented in vertical axis to the catalytic site and is located at a distance, more than the standard non-bonded interactions from the active site residues. This pose would not bring the carboxyl site closer to the position of His57 and Ser195. Similarly, the Arg1018 of Peptide II is penetrated into the binding cavity, nevertheless the carboxyl site is not occupying the close proximity of active site residues. Hence, both the peptides docked at a distal position, which is not fair enough for the hydrolysis.

**MD simulation analysis for RVV-V with peptides**

Both the Complexes were subjected for MD simulation for 60ns. The backbone RMSD of both the peptide Complexes (Fig 5[E]) reveals that the Complex R2 deviates more than R1. From the profile, the Complex R2 deviates up to a level of 0.6 nm and stays in the range 0.55 nm. Similarly, the RMSD for R1 went up to 0.5 nm and stays in the same range. For both the complexes, the RMSD fluctuation after 20 ns remains within 0.1 nm.

Table 2. The non-bonded interaction for the best-docked pose of Peptide I with RVV-V.

<table>
<thead>
<tr>
<th>SL N</th>
<th>ATOM 1 RVV-V</th>
<th>ATOM 2 Peptide</th>
<th>Distance</th>
<th>Category</th>
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<td>THR39:N</td>
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<tr>
<td>3</td>
<td>SER38:CB</td>
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</tr>
<tr>
<td>4</td>
<td>HIS192:CE1</td>
<td>ARG709:O</td>
<td>3.42154</td>
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</tr>
<tr>
<td>5</td>
<td>ASN97:O</td>
<td>GLY701:CA</td>
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<td>H-Bond</td>
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<tr>
<td>6</td>
<td>ALA56</td>
<td>LEU702</td>
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<tr>
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<td>LEU702</td>
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<tr>
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https://doi.org/10.1371/journal.pone.0181216.t002

Table 3. The non-bonded interaction for the best-docked pose of Peptide II with RVV-V.

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</table>

https://doi.org/10.1371/journal.pone.0181216.t003
Fig 5: MD simulation analysis for RVV-V. A) An average inter-molecular hydrogen bonds (1000 Frames) throughout simulation between RVV-V and peptides. B) An average intramolecular hydrogen bond for peptides. C) A solvent accessible surface area (SASA) for the complexes (RVV-V with peptides). D) Shows SASA for only peptides. All the parameters show more fluctuation for Complex R1 as compared to R2. (E and F) represent RMSD and radius of gyration (Rg) of the Complexes and apo-form (Black), respectively. RMSD and Rg are stabilized for both the Complexes after 20ns. A rise in Rg value at 10ns for R2 is the result of the structural modification in RVV-V due to Peptide II.

https://doi.org/10.1371/journal.pone.0181216.g005
The complex R1 throughout simulation maintains the same range of radius of gyration, signifying no major structural changes due to interaction with the Peptide I. The complex R2 shows an increment in the Rg values (1.75 nm to 1.83 nm) and later decreases back to 1.75 nm after 20 ns which signifies the binding of the Peptide II is stabilized (Fig 5[F]).

Solvent Accessible surface area (SASA) for the Complex R2 (Fig 5[C]) initially rises and then falls but after 25 ns remains within 135 to 142 nm$^2$ whereas for the Complex R1 it fluctuates within 125 nm$^2$ to 135 nm$^2$ after 10 ns. Here, the higher value of SASA in the Complex R2 is most likely due to the distortion introduced in the RVV-V by the Peptide II and the decreasing SASA can be the outcome of the increased interaction between the Peptide II and the RVV-V. In other words, the peptide and the solvent molecules stabilized their interaction during the course of simulation. The SASA for the peptide alone (Fig 5[D]) in Complex R1 shows a fluctuation for Peptide I whereas for Peptide II it remains stable throughout the simulation trajectory. The fluctuation in SASA can be the outcome of the unstable docked pose whereas the stable SASA shows a stable docked pose.

The number of inter and intra molecular hydrogen bonds between the RVV-V and the peptides throughout the simulation has been plotted (Fig 5[A] and 5[B]). The number of intermolecular H-bonds of RVV-V with Peptide I was found to be fluctuating throughout the simulation whereas for Peptide II it maintains the number of H-bonds consistently except for the three regions. On an average, the overall number of intermolecular H-bonds with Peptide II was higher in comparison to Peptide I. The higher and stable H-Bond pattern between the RVV-V and the Peptide II support for a stable and better binding Pose. In the case of Peptide I the fewer intermolecular H-bond pattern does not support for the stable and better binding pose. Instead, it leads to a loosely bound nature with RVV-V. A consistent intra-molecular H-Bonding of Peptide II may support in tight binding with RVV-V when compared with Peptide I. Thus, we can conclude that the Peptide II binds better at the active site than the Peptide I.

The root mean-square fluctuation (RMSF) profile (Fig 6) for both the Complexes indicates a significant elevation in comparison with the native protein. The loop and the coil regions show a higher deviation indicating the binding of peptide with RVV-V. It can be observed that the region 21–26, 71–76, 91–95, 125–132, in Complex R2 shows more deviation than the Complex R1. However, in the region of N- and C terminal R1 showed a higher fluctuation.

The Peptide II shows a better binding with RVV-V. The hydrogen bond pattern observed for Complex II supported the minimal solvent exposure to the protein, which enhances the tight binding. Similarly, we can conclude that the Peptide I showed a weak binding with RVV-V.

**PCA analysis of RVV-V with peptides**

To understand the structural and the interaction behaviour of RVV-V with the peptides the dihedral principle component analysis ($\phi$PCA) was performed. The free energy landscape (FEL) was drawn using the first two principle components with cosine value less than 0.5 derived from calculated PCs. The FEL of complex R1 (Fig 7[A]) showed one large cluster and various small clusters, a representative structure from the largest cluster minima was extracted and analyzed for non-bonded interaction. Here, the Peptide I makes fifteen hydrogen bonds, one electrostatic bond, and five hydrophobic interaction. The active site residue His57 shows pi-cation interaction (4.2A) with NH1 of Arg709 of Peptide I and the other active site residues Ser195 and Asp189 show no interaction as the Arg709 is moving away from subsite $S_0$. The residues at position $P_{-1}$, $P_{-2}$, and $P_{-3}$ also moved away from $S_1$ thus shows no interaction with $S_1$ subsite; but the residues at $P_2$, $P_3$, and $P_4$ bind in $S_1$ subsite. The complete and detailed
interactions between the RVV-V and the Peptide I are given in S1 Table. If we compare the docked pose with the FEL, we can say that the course of simulation the peptide orients itself and induces slight conformational changes in RVV-V. The docked pose showed only five residues of Peptide I (Asn713, Arg709, Ala705, Leu702, and Gly701) interacting with the RVV-V but in the FEL minima conformation, most of the residues of the peptide interact except Gly701, Ala704, Ala705, Ile708. A change in Psi angle (dock to FEL) of Gly701 from 6.56˚ to 116.48˚ flips Gln699 and Asn700 towards the 97-Loop which induces the loop distortion and moves slightly away from S_{-1}, thus shifts Asp102 away from Ser195. In-silico alanine scanning mutation for the interacting residues (S1 Fig) shows that the residues Ser38, Ile40, Gln221A (with ΔΔG greater than one) play an important role in stabilizing the Peptide I binding to RVV-V. It is evidenced that the residues at P_{0} (cleavage site) are away from the catalytic triad and the other residues play a major role in holding the peptide with RVV-V.

A minimum energy conformation of Complex R2 was extracted from the largest FEL cluster and the interaction pose is depicted in Fig 7[B]. The Complex shows a better and stable interaction with twenty six H-bond, and eight hydrophobic interactions. Here, Arg1018 (P_0) interacts in the vicinity of the subsite S_{0} with 30.5% of all non-bonded interaction and 42.8% of the conventional H-bonds. The interaction of P_{-2}, P_{-1}, P_{0}, P_{1}, P_{2}, and P_{3} residues with 220-Loop stretches and moves it toward 174-Loop. The 97-Loop moves away from 220-Loop to accommodate the residues at P_{4} (Pro1014), P_{5} (Ala1013), and P_{6} (His1012) position of the Peptide II. The change in 97-Loop conformation displaces Asp102 away from Ser195 and His57 and modifies subsite S_{-1} into a deep groove (the details of interactions are given in

![Image](https://doi.org/10.1371/journal.pone.0181216.g006)
Fig 7. The FEL for complex R1 (A) and R2 (B) based on the $\omega$PCA analysis. One representative structure is displayed from the most populated free energy minimum clusters. In Complex R1, the Arg709 has moved away from S0 and show a pi-cation interaction with His57. The distance between O$_{\gamma}$ atom of Ser195 and C' atom of carboxyl oxygen of Arg (P0) has been shown for both the peptides.

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Alanine scanning showed that the residues His41, Glu218, Gln221A, and Thr229 (with ΔΔG greater than one) play an important role for binding the Peptide II with RVV-V (S1 Fig).

In-conclusion, the Peptide I binds superficially with the lesser number of non-bonded interactions and unstable conformations (Energy Landscape, H-Bond) resulting in the movement of Arg709 away from subsite S0 whereas the binding of Peptide II deeper in the cavity induces the structural modification mainly in 174-Loop, 97-Loop, and catalytic triad position (S2 Fig). Despite Arg1018 plays a major role in anchoring the Peptide II in subsite S0, the RVV-V does not cleave the peptide, as the modification in the position of catalytic triad does not favour it.

Peptide docking with thrombin using HADDOCK

The peptides were docked with thrombin viz., complex T1 (with Peptide I) and T2 (with Peptide II). The peptides in both the complexes had residues of P1, P2, and P3 positions in subsite S1, P1, P2, and P3 in S-1, and P0 (Arg) in subsite S0 (S3 Fig) and exhibited numerous non-bonded interaction (S3 and S4 Tables). To check the dynamic stability of all the complexes, the MD simulation was performed for 100ns.

MD simulation analysis for thrombin with peptides

The backbone RMSD profile of the thrombin apo-form, Complexes T1, and T2 gets stabilized around 40 ns and maintains the deviation within ±0.05nm whereas the radius of gyration (Rg) showed a consistency throughout MD (Fig 8[E] and 8[F]). The peptide in Complex T1 and T2 maintained the intra-molecular hydrogen bond around four and three respectively whereas the inter-molecular H-bond between the thrombin and the peptide was maintained around six for both the Complexes throughout the MD trajectory (Fig 8[A] and 8[B]). The solvent accessible surface area for the Complexes T1 and T2 was maintained between 135 to 140 nm² and for the peptide alone it was around 20 nm² (Fig 8[C] and 8[D]).

The RMSF for thrombin (Fig 9) showed more fluctuation in the coil region, the highest in the region 60-Loop (for apo-form and T1) and 192-Loop (apo-form and T2). The key residues His57 and Asp189 showed the least fluctuation but Asp102 and Ser195 (for apo-form and T2) showed a significant fluctuation.

PCA analysis of thrombin with peptides

The FEL of thrombin Apo-form (S5 Fig) shows three major clusters out of which the largest one has two minima. The coordinates were extracted from each minima and analyzed; two distinct conformations were recognized as open and closed. The FEL for the Complex T1 (Fig 10) has one major energy basin and a representative conformation was extracted for the analysis whereas the FEL for the Complex T2 (Fig 11) had two major energy basins and the representative frames were extracted. The conformation shows the peptides in both the Complexes, Arg (at P0) binds deeper in subsite S0 and residues P1, P2, and P3 at S1. It is notable that the 97-Loop shows no interaction with Peptide II while the interaction is negligible with Peptide I. In both complexes, the residue Trp215 of thrombin plays an important role in holding the peptide in S1 subsite (S5 and S6 Tables).

The NH2 of Arg in Peptide II is making a salt bridge with Asp 189 of the thrombin (Fig 11). The NH1 of Arg makes H-bond with the hydroxyl group of Tyr 228. Both these interactions may anchor the Arg for cleavage in the active site. The cleavage site is at a close proximity of the active site residues His 57 and Ser 195, which is important for the cleavage. Hence, we can
Fig 8. MD simulation analysis in terms of H-Bond, SASA, RMSD, and Rg for thrombin. A) An average (1000 Frames) of H-bond throughout simulation between thrombin and peptide. B) The intramolecular H-bond for peptides. C) The solvent accessible surface area (SASA) for the Complexes T1 and T2. D) SASA for only the peptides. The parameters show a stable curve throughout simulation. (E and F) shows the RMSD and the Rg values respectively for both the complexes and the apo-form. The RMSD for all is stabilized after 40ns. The Rg values remain same for both complex and apo-form.

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Fig 9. The RMSF for each residue of thrombin. The Apo-form shows a significant fluctuation at region 60B-62 (60-Loop) and 190–196 (192-Loop). T1 fluctuates in 192-Loop region. The key residues His57 and Asp189 fluctuate less whereas Asp102 and Ser195 (for apo-form and T2) show a significant fluctuation.

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Fig 10. FEL for T1. One representative structure is displayed from the most populated free energy minimum clusters. The peptide is shown in stick and the thrombin in cartoon. The loops are labeled in brown the subsites are marked by yellow circles. The structure is zoomed to show distances (with red dotted line) between key residues/atoms.

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conclude that both the peptides make a favourable interaction with the active site residues for the hydrolysis.

**MD simulation & PCA analysis for Peptide III with RVV-V**

The discussions above clearly indicated that the Peptides I and II induced local and global structural change which brought instability to the protein. Ultimately RVV-V turned incapable of cleaving both sites. Whereas the same peptides induced no structural change in thrombin and better suitable for cleavage. The crystal structure of RVV-V with third site is available and the measurement of structural flex by MD will help to determine the expected structural stability of RVV-V. Hence the docking and MD was performed with third site i.e Peptide III. The results from MD simulation and PCA analysis showed that RVV-V is intact and Peptide III was identified in a location suitable for cleavage. Corresponding figures are given as supplementary (S8 and S9 Figs). The result further confirmed that both Peptides I and II induce structural instability in RVV-V.

**Binding specificity of RVV-V with site I and site II**

To understand the specificity and the structural difference the RVV-V was compared with the thrombin. The structural alignment was made (S5[C] Fig) and the sequence reflecting the alignment is shown in S5[A] Fig. In alignment, we found five inserts in thrombin and were named as I1 (60D-62), I2 (129C-134), I3 (148-149A), I4 (186D-188), I5 (204–210), and one tail

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extension in RVV-V. The inserts marked in hot pink modify the corresponding loops/regions 60-Loop, 129-Loop, 149-Loop, 186-Loop, and 206-Loop, which make the thrombin more flexible. The comparison of the closed and the open forms of the thrombin shows a significant deviation at 60-Loop (Residue Asp60E (Cα separated by 7.4 Å), 149-Loop (1.8 Å for Trp148), 186-Loop (6.3 Å for Glu186B), and 220-Loop (3.7 Å for Arg221A). The 60-Loop and 149-Loop give the mechanical support for the peptide binding in thrombin, which is not observed in the RVV-V. Thus, the peptide interaction is more dependent on the hydrophobic surface distribution of the active site cavity (S4 Fig). The hydrophobicity based surface showed S1 slightly hydrophobic and S1 Hydrophilic. The 60-Loop and the residue Trp215 also shield the 97-Loop from interacting with the peptides while binding in active site resulting in maintaining the catalytic triad in thrombin.

The intermolecular interaction of both the peptides with the thrombin is more stable than with the RVV-V. It is evident that the susceptible cleavage sites Arg 709 and Arg 1018 of Peptides I and II in the Complexes R1 and R2 are not in the vicinity of His 57 and Ser 195 (Fig 7). Binding of both the peptides dislocates the catalytic triad in RVV-V; whereas the same peptide found near the catalytic triad in thrombin (Figs 10 and 11) does not disrupt the catalytic triad. Moreover, it is an ideal binding pose for the Peptide II with the thrombin. However, in RVV-V, the binding pose disrupts both the local and the global structure. After simulation, the numerous interactions found in the RVV-V peptide complexes favour only the global/local structural change of RVV-V, which does not guarantee the peptide as a substrate. For the cleavage at carboxyl group of Arg the C’ atom should lie closer towards His 57. In R1 and R2 complex structures, they are beyond the normal H-bonding distances (Fig 7). However, in the case of T1 and T2 (Figs 10 and 11) this distance is apt and ostensible for the cleavage. In R1 and R2, even after 60 ns simulation both the peptides do not select a favourable conformation, which is apt for the cleavage. At the same time, both the peptides are engaged with a numerous interactions away from cleavage site (S1 and S2 Tables). Thus, the peptide binding influences the global/local structural changes in RVV-V leaving the peptides with numerous perpetual interactions that are not supportive for the hydrolysis. This explains that the Peptide II shows a weak affinity in SPR experiment while the Peptide I showed none.

Conclusion

The RVV-V, a serine protease, specifically cleaves the site III of the FV and not the sites I and II. The reason for such a selective specificity is not explained so far. The present study captures the scenario of the RVV-V with the sites I and II at a molecular detail. The SPR binding kinetic study showed that the Peptide II exhibits a weak binding affinity and induces the conformational changes towards the RVV-V with K_D of 2.775 mM and the Peptide I showed none. This weak affinity corroborates the binding of the peptide with the RVV-V. The docking and the MD simulation studies put an insight into molecular interaction and binding mechanism of these peptides with the RVV-V. The docking result showed the peptide binding near the active site of the RVV-V with fewer interactions. The MD simulation revealed an unstable binding of Peptide I, which also induces local structural modification in RVV-V whereas the Peptide II binding was found to be stable, and showed the local/global changes in the RVV-V. The binding pose of Peptide I and II with the RVV-V is inapt for the cleavage and leave the peptide with a numerous perpetual interactions, which was exemplified by the apt binding pose of the same with the thrombin. Thus, the weaker K_D of Peptide II could be due to the perpetual interaction and not by proper docking of peptide for the cleavage.

Other features that could define specificity of RVV-V over thrombin can be summarized as:
1. Thrombin has larger loops due to inserts and hence it is more flexible compared to RVV-V and shuttles between the open and the closed conformation.

2. The 60-Loop and the large/flexible 149-Loop in thrombin give mechanical support to the binding peptides. Lack of which in RVV-V makes it more dependent on non-bonded interaction with the peptide.

3. The 60-Loop and the residue Trp215 shield 97-Loop in thrombin thus the catalytic triad is undistorted in the presence of binding peptide. In contrary, Catalytic triad is modified due to the exposed 97-Loop in RVV-V.

4. The hydrophobic surface of the RVV-V supports only the peptide with the cleavage site III to interact without interfering 97-Loop.

   The study suggests a transition in RVV-V from the native rigid to the distorted flexible structure in the presence of peptides and paves a way to design a new peptide substrate/inhibitor.

Supporting information

S1 Table. The non-bonded interaction for the Peptide I with RVV (Complex R1) extracted minima conformation from FEL. (PDF)

S2 Table. Non-bonded interaction for Peptide II with RVV (Complex R2) extracted minima conformation from FEL. (PDF)

S3 Table. The Non-bonded interaction for the best docked pose of Peptide I with thrombin (Complex T1). (PDF)

S4 Table. The Non-bonded interaction for the best docked pose of Peptide II with thrombin (Complex T2). (PDF)

S5 Table. The Non-bonded interaction for the Peptide I with the thrombin (Complex T1) extracted minima conformation from FEL. (PDF)

S6 Table. The Non-bonded interaction for the Peptide II with the thrombin (Complex T2) extracted minima conformation from FEL. (PDF)

S1 Fig. A summary for computational alanine scanning for RVV-V with Peptide I (A) and Peptide II (B). Positive binding free energy differences indicate a potential important residue for binding with the peptides. (TIF)

S2 Fig. Show distances (in Angstrom) between the loops and catalytic triad residues (His57, Asp102, and Ser195) in R1 (A) and R2 (B) after MD simulation. In R2, the 174-Loop gets modified and 97-Loop move away from 220-Loop and 174-Loop. (TIF)

S3 Fig. The docked pose of the Peptide I (stick) with the thrombin (cartoon). The peptide residue are labeled in black, loops are labeled brown, the subsites are labeled in red and shown
in yellow shade. In both the complexes Arg at P₀ (709 in Complex T1 and 1018 in Complex T2) binds in S₀.

S4 Fig. The hydrophobic surface for RVV-V (A) and open form of thrombin (B) showing the position of subsite S₁₋₁, S₀, S₁. RVV-V shows hydrophobic S₁ and hydrophilic S1 whereas thrombin shows hydrophilic S₁ and hydrophobic S₁. Trp215 almost covers S₁ in thrombin.

S5 Fig. A comparison of RVV-V and thrombin open form, by structural alignment (C) and reflecting the sequence alignment (A). The FEL for thrombin apo-form (B) and the structural alignment of extracted open and closed forms (D) are shown in the cartoon. RVV-V is in copper colour, the thrombin closed form is in green and the open form in magenta colour. The inserts are shown in hot-pink colour and labeled in it, and the distances are measured in Angstrom.

S6 Fig. Chromatogram for size exclusion and cation exchange. The crude venom was first fractionated by FPLC size exclusion column which yielded seven peaks (P₁ – P₇) where P₃ (marked with circle) was pooled and used for cation exchanged purification (down). Here we obtained mainly three peaks (P₁, P₂ and P₃) where P₁ and P₂ (marked by red square) both showed positive activity for RVV-V.

S7 Fig. SDS-PAGE after cation exchange FPLC purification. Lane-1 marker, Lane-2 P₁, Lane-3 P₂, and Lane-4 P₃.

S8 Fig. MD simulation analysis in terms of RMSD, and Rg and RMSF for RVV-V apo form and with peptides. Here the RMSD, Rg and RMSF is lowest for the complex RVV-V with Peptide III.

S9 Fig. FEL for RVV-V with Peptide III. One representative structure is displayed from the most populated free energy minimum clusters. The peptide is shown in stick and the thrombin in cartoon. The loops are labeled in brown the subsites are marked by yellow circles. The structure is zoomed to show distances (with red dotted line) between key residues/atoms.

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References


