Molecular Insight into the Steric Shielding Effect of PEG on the Conjugated Staphylokinase: Biochemical Characterization and Molecular Dynamics Simulation

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Abstract

PEGylation is a successful approach to improve potency of a therapeutic protein. The improved therapeutic potency is mainly due to the steric shielding effect of PEG. However, the underlying mechanism of this effect on the protein is not well understood, especially on the protein interaction with its high molecular weight substrate or receptor. Here, experimental study and molecular dynamics simulation were used to provide molecular insight into the interaction between the PEGylated protein and its receptor. Staphylokinase (Sak), a therapeutic protein for coronary thrombolysis, was used as a model protein. Four PEGylated Saks were prepared by site-specific conjugation of 5 kDa/20 kDa PEG to N-terminus and C-terminus of Sak, respectively. Experimental study suggests that the native conformation of Sak is essentially not altered by PEGylation. In contrast, the bioactivity, the hydrodynamic volume and the molecular symmetric shape of the PEGylated Sak are altered and dependent on the PEG chain length and the PEGylation site. Molecular modeling of the PEGylated Saks suggests that the PEG chain remains highly flexible and can form a distinctive hydrated layer, thereby resulting in the steric shielding effect of PEG. Docking analyses indicate that the binding affinity of Sak to its receptor is dependent on the PEG chain length and the PEGylation site. Computational simulation results explain experimental data well. Our present study clarifies molecular details of PEG chain on protein surface and may be essential to the rational design, fabrication and clinical application of PEGylated proteins.

Introduction

PEGylation, conjugation of polyethylene glycol (PEG) to a therapeutic protein, is a successful approach to improve therapeutic potency of proteins [1,2]. The therapeutic potency is improved by increasing serum half-life, decreasing immunogenicity, and reducing renal clearance and proteolytic degradation of the protein [3–5]. Typically, a therapeutic protein is conjugated with PEG via a covalent linkage at some reactive moieties of the protein. PEG entangles around the protein surface through hydrophobic interaction and concurrently forms hydrogen bonds with the surrounding water molecules. On one hand, the steric shielding effect of PEG can achieve the advantages mentioned above. On the other hand, PEG may sterically shield the bioactive domain (e.g., substrate or receptor binding domain) of a protein, resulting in a substantial loss of its bioactivity [4–6]. Therefore, it is of interest to understand the steric shielding effect of PEG on the interaction between a protein and its substrate/receptor.

The PEGylation site and the PEG mass are two main factors that influence the bioactivity of the protein. Site-specific PEGylation of a protein far from its bioactive domain has been demonstrated to decrease the loss of bioactivity of the PEGylated protein [7,8]. In contrast, PEGylation at the bioactive domain may result in complete loss of bioactivity of a protein. However, this approach can not completely avoid the loss of bioactivity, due to the presence of the steric shielding effect of PEG [9,10]. Some therapeutic proteins, whose pharmacological effects involve high molecular weight (Mw) substrate or receptor interactions, show in vitro bioactivity is inversely proportional to the mass of attached PEG. For example, Chiu et al. prepared PEGylated trypsin conjugated with 2, 5, 10 and 20 kDa PEG, respectively [11]. PEG with 10 kDa was demonstrated to be the optimal size to improve thermal stability and maintain bioactivity under physiological conditions. However, the molecular mechanism of the interaction between the PEGylated protein and its receptor is not adequately clear.

Molecular dynamics simulation is an effective approach to reveal atomic level details of biological interactions [12,13] and has been used for PEG-protein simulation. For example, Manjula et al. built the PEG-hemoglobin models by means of molecular dynamics simulation [14]. They found that the PEG chain was folded loosely on the surface of hemoglobin and the coverage was not proportional to the PEG chain length. Yang et al. modeled the PEG-insulin system and found the optimized conformations of the conjugates using a simulated annealing method [15]. However, these studies focus on the effects of the PEG chain length on the properties of PEGylated protein, where the effect of the
PEGylation sites is ignored. In addition, the underlying mechanism of the steric shielding effect of PEG on the protein therapeutic potency is not well understood, especially for protein interaction with its high M_w substrate or receptor.

In the present work, experimental studies and molecular dynamics simulations were used to provide a molecular insight into the interaction between the PEGylated protein and its receptor at the atomic level. Staphylokinase (Sak) is a bioactive protein that can disintegrate thrombus through activating the plasma plasminogen to plasmin and is used as the model protein here [16,17]. Compared with native Sak, Sak used here lacks 10 amino acid residues at the N-terminus and has an additional peptide of Gly-Gly-Cys at the C-terminus [18]. In this study, 5 kDa and 20 kDa PEG aldehydes were used for site-specific PEGylation of Sak at the α-amino group at the N-terminus (Fig. 1A). In addition, 5 kDa and 20 kDa PEG maleimides were used for site-specific PEGylation of Sak at the thiol group at the C-terminus (Fig. 1B). The structural and functional properties of the four PEGylated Saks were characterized and compared with the free Sak.

Molecular models of the PEGylated Saks were constructed and molecular dynamics simulations of the PEGylated Saks performed. Docking analyses were carried out to investigate interaction between the PEGylated Sak and its receptor, plasminogen. Plasminogen was represented by micro-plasminogen (micro-plg) for docking purpose. Molecular dynamics simulations and docking analyses explain well the experimental results. Our study is expected to facilitate understanding of the steric shielding effect of PEG on a protein, especially the ones involving interactions with their high M_w substrate or receptor.

Materials and Methods

Materials

The recombinant staphylokinase (Sak) was produced in a transformed E. coli and purified by ion exchange chromatography and size exclusion chromatography as described previously [18]. Methoxyl PEG propionaldehyde of 5 kDa (ald5k) and 20 kDa (ald20k), methoxyl PEG maleimide with 5 kDa (mal5k) and methoxyl PEG maleimide with 20 kDa (mal20k) were purchased from Jenkem Biotech (Beijing, China). Tris(2-carboxyethyl)phosphine (TCEP) was obtained from Pierce (USA). NaCNBH_3 was obtained from Sigma (USA).

Preparation of the PEGylated Sak

C-terminally PEGylated Sak. Sak was incubated with TCEP at the Sak/TCEP molar ratio of 1:10 in 20 mM sodium phosphate buffer, pH 7.2 (PB buffer). The incubation was conducted at room temperature for 3 h to prevent the S-S dimerization of Sak. Then, TCEP was removed by centrifugation at 5000 rpm with 3 kDa cutoff membrane against PB buffer for three times. Sak (1.5 mg/ml, 0.1 mM) in PB buffer was incubated with 0.4 mM mal5k and 0.4 mM mal20k at 4°C overnight, respectively.

N-terminally PEGylated Sak. TCEP treated Sak was dialyzed against 20 mM sodium acetate buffer (pH 5.0). Afterwards, Sak (0.1 mM) in PB buffer was incubated with 0.4 mM mal5k and 0.4 mM mal20k in the presence of 10 mM NaCNBH_3, respectively. The incubation was in 20 mM sodium acetate buffer (pH 5.0) at 4°C overnight.

Purification of the PEGylated Saks. The reaction mixtures were subjected to an SP Sepharose HP column (1.6 cm x 25 cm, GE Healthcare, USA) that was equilibrated with 20 mM NaAc-HAc buffer (pH 5.0, Buffer A). As for mal5k and mal20k treated
Sak, the column was eluted with Buffer A for removal of the free PEG reagent, followed by elution with a linear salt gradient of 0–0.2 M NaCl in Buffer A for 30 min at a flow rate of 1.0 ml/min. The effluent was detected at 254 nm. The fractions corresponding to Sak-mal5k and Sak-mal-20k were pooled and concentrated for further experiment, respectively.

As for ald5k and ald20k treated Sak, the column was eluted with 12 ml of Buffer A for removal of the free PEG reagent, followed by elution with 18 ml of 0.5 M NaCl in Buffer A at a flow rate of 1.0 ml/min. The effluent was detected at 254 nm. The fractions corresponding to the mixture of Sak and PEGylated Sak were pooled, respectively. The fractions were loaded on a Superdex 200 column (2.6 cm ×60 cm, GE Healthcare, USA) based on size exclusion chromatography (SEC). The column was equilibrated and eluted by 20 mM phosphate buffer (pH 7.2) at a flow rate of 3.0 ml/min. The fractions corresponding to Sak-ald5k and Sak-ald-20k were pooled and concentrated for further experiment.

Analysis of PEGylated Sak

SEC analysis of the Sak samples was carried out at room temperature on a Superdex 200 column (1 cm×30 cm, GE Healthcare, USA). The column was equilibrated and eluted by PB buffer at a flow rate of 0.5 ml/min. The effluent was detected at 280 nm. SDS-PAGE analysis was performed on a 14% polyacrylamide gel. The gel was stained with Coomassie blue R-250.

Circular dichroism spectroscopy

Circular dichroism (CD) spectra of the products were recorded on a Jasco-810 spectropolarimeter (Jasco, Japan) from 260 to 195 nm at the room temperature. A cuvette with 0.2 cm pathlength was used. All samples were at a Sak concentration of 0.2 mg/ml in PB buffer.

Fluorescence measurement

Intrinsic fluorescence measurements were performed using Hitachi F-4500 fluorescence spectrometer with a cuvette of 1 cm pathlength. The emission spectra were excited at 280 nm and recorded from 300 to 400 nm. Excitation and emission slit widths were both 5 nm. All samples were at a protein concentration of 0.1 mg/ml in PB buffer.

Table 1. Parameters used for HEX docking.

| Parameter    | Value | Meaning of the Parameter                                                                 |
|--------------|-------|-----------------------------------------------------------------------------------------|
| Correlation type | Shape+Electrostatics | Functions to calculate the docking correlation between two molecules                     |
| FFT Mode     | 3D    | Three-dimensional (3D) rotational correlation, for accelerating the rotation part of the search |
| Grid Dimension | 0.6   | The sampling grid size                                                                   |
| Receptor range | 30    | The range angle of scanning at the receptor’s surface                                     |
| Ligand Range | 30    | The range angle of scanning at the ligand’s surface                                       |
| Twist range  | 40    | The intermolecular twist angle                                                            |
| Distance Range | 0.8   | The limit of intermolecular separation from the initial distance                           |
| Scan step    | 20    | The scan unit of the distance                                                             |
| Steric Scan  | 25    | The spherical polar docking expansion order (N) used for calculate electrostatic potential at the initial scan |
| Final Scan   |       | The higher N used for final scan to refine the search                                     |

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Molecular dynamics simulations were performed in GROMACS 4.5.4 [21] with GROMOS96 53a6 force field [22]. PEG parameterization was performed with quantum computation within Gaussian 09W [23], resultant parameters were integrated to the 53a6 force field to form a new force field, named 53a6_PEG. Its suitability for PEG simulation was validated by comparing its performance in PEG simulation to that of force field 53a6_OE developed in [24]. Figures S1 and S2 show that 53a6_PEG results in more extended PEG chains than 53a6_OE. Figures S3 and S4 show that radii of gyration (Rg) of PEG chains based on 53a6_PEG are larger than those based on 53a6_OE. Figures S1-S4 thus show that 53a6_PEG is better able to represent PEG’s good solubility in water (see File S1 for details). Simulation visualization was performed in Visual Molecular Dynamics.

Figure 3. Characterization of the four PEGylated Saks. (A) SEC analysis was carried out on a Superdex 200 column (1 cm × 30 cm). The column was equilibrated and eluted with 20 mM sodium phosphate buffer (pH 7.2) at a ow rate of 0.5 mL/min. (B) SDS-PAGE analysis of the samples. Lanes 1–6 were standard protein marker, Sak, Sak-mal5k, Sak-ald5k, Sak-mal20k and Sak-ald20k, respectively.

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Figure 4. Structural characterization of the PEGylated Saks. Circular dichroism analysis was carried out for the two C-terminally PEGylated Saks (A) and the two N-terminally PEGylated Saks (B). Intrinsic fluorescence analysis was performed for the two C-terminally PEGylated Saks (C) and the two N-terminally PEGylated Saks (D).

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further linearly increased to 450 K within 3 ns and kept constant
3 ns and then kept constant at 400 K for 2 ns. Temperature was
linearly increased from 300 to 400 K within
Simulated annealing was adopted to speed up simulation.
Steepest descent method was used for energy minimization.
NVT and 100 ps NPT equilibrations; after which a simulated
reduce inappropriate solute-solvent contacts, followed by 100 ps
Time step was set to 2 fs. Energy minimization was run first to
Ewald (PME) method was employed for electrostatic interactions.
The van der Waals and electrostatic interactions were both
Docking analyses
Configuration of PEGylated Sak at the end of simulation
process was docked to micro-plg in HEX 6.12. Crystal structure of
micro-plg was obtained from the Protein Data Bank (1QRZ) [26].
The centroids of PEGylated Sak and micro-plg were set within
30 Å of each other, which is the best working range for HEX
docking [27]. The known crystal structure of micro-plg-Sak-micro-
plg complex (1BU1) [28] provides Sak and micro-plg binding
domains and their orientation toward each other, that information
is used to set up initial positions of PEGylated Sak and micro-plg

![Figure 5. In vitro bioactivity of the PEGylated Saks.](Image)
The in vitro bioactivity of the PEGylated Saks was tested by fibrin plate assay. The bioactivities of the PEGylated Saks were compared with that of the unmodified Sak, which was set to 100%.
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(VMD) from the theoretical and computational biophysics group at UIUC [25]. To evaluate initial PEG conformation’s effect on simulation results, we had performed test simulation runs using PEGs with different initial conformations. Fig. S5A shows that the system attains a similar size starting from different initial PEG conformations. Fig. S5B shows that differences between final structures resulted from different initial conformations are small, all less than 1 nm. Figure S5 therefore shows that initial PEG conformation does not have much influence on its final wrapping state over the protein surface. Repeated MD trajectories were thus not generated in this work.

The four PEGylated Saks were solvated by the simple point charge (SPC) water model and neutralized by adding sodium ions (Na\(^+\)). Triclinic cell was used for periodic boundary conditions. The van der Waals and electrostatic interactions were both switched at 0.9 nm and then cut off at 1.4 nm. Particle mesh Ewald (PME) method was employed for electrostatic interactions. Time step was set to 2 fs. Energy minimization was run first to reduce inappropriate solute-solvent contacts, followed by 100 ps NVT and 100 ps NPT equilibrations; after which a simulated annealing procedure with a NVT ensemble was performed. Temperature was controlled by velocity rescaling method. Steepest descent method was used for energy minimization. Simulated annealing was adopted to speed up simulation. Temperature was linearly increased from 300 to 400 K within 3 ns and then kept constant at 400 K for 2 ns. Temperature was further linearly increased to 450 K within 3 ns and kept constant at 450 K for another 2 ns, followed by a symmetric decrease to 300 K. The system was then kept at 300 K for additional 7 ns.

### Docking analyses

Configuration of PEGylated Sak at the end of simulation process was docked to micro-plg in HEX 6.12. Crystal structure of micro-plg was obtained from the Protein Data Bank [1QRZ] [26].

| Sample   | \( S_{20,w} \) \(^a\) | \( f/f_0 \) \(^b\) |
|----------|------------------|-----------|
| Sak      | 1.75±0.15        | 1.20      |
| Sak-mal5k| 1.36±0.05        | 1.48      |
| Sak-ald5k| 1.34±0.09        | 1.65      |
| Sak-mal20k| 1.21±0.04       | 2.35      |
| Sak-ald20k| 1.21±0.04       | 2.39      |

\(^a\) The sedimentation coefficient in a standard state of water at 20°C.  
\(^b\) The ratio of frictional coefficient.

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**Table 2. Sedimentation velocity coefficients of the PEGylated Saks.**

for HEX docking. Actual HEX docking parameters are listed in Table 1.

**Results**

**Characterization of the products**

Under the present experimental conditions, the PEGylated products by mal5k and mal20k were a mixture containing approximately 90% mono-PEGylated Sak and 10% unPEGylated Sak (data not shown). As shown in Fig. 2A, the free PEG was removed from the PEGylated products by ald5k and ald20k by the SP Spherarose HP column. PEGylated products by ald5k and ald20k were a mixture containing 50–55% mono-PEGylated Sak, 10–15% highly PEGylated forms and 30–40% unPEGylated Sak (Fig. 2B). The mono-PEGylated Saks (Sak-ald5k and Sak-ald20k) were separated from the reaction mixtures by Superdex 200 column (2.6 cm×60 cm). The PEGylation site of Sak-ald5k and Sak-ald20k was characterized by tryptic peptide mapping according to Liu et al. [18] and demonstrated to be the N-terminus of Sak (data not shown).

The purified PEGylated proteins were further analyzed by an analytical Superdex 200 column (1.0 cm×30 cm). As indicated by the SEC analysis (Fig. 3A), Sak-ald5k and Sak-mal5k are both eluted as single and symmetric peaks, which are left-shifted as compared with Sak. The elution peaks corresponding to Sak-ald20k and Sak-mal20k are further left-shifted as compared with Sak-ald5k and Sak-mal5k. This indicates that the hydrodynamic volume of Sak is enhanced by PEGylation and is dependent on the conjugated PEG mass. Moreover, the peak corresponding to Sak-mal5k is slightly left-shifted as compared to Sak-ald5k. Similarly, Sak-mal20k is eluted earlier than Sak-ald20k. This suggests that the PEGylation sites may alter the hydrodynamic volume of the PEGylated Sak.

As indicated by SDS-PAGE analysis (Fig. 3B), Sak shows a single electrophoresis band corresponding to an \( M_w \) of ~15 kDa (Lane 2). As compared to Sak, Sak-ald5k and Sak-mal5k show a single band with slower migration than Sak and slightly faster than Sak-mal5k (Lane 3). This indicates that the band migration of Sak is retarded by the conjugated PEG. Moreover, the PEGylation site may determine the band migration of the PEGylated Sak. Similarly, Sak-ald20k (Lane 6) shows a single band with slower migration than Sak-ald5k and slightly faster than Sak-mal20k (Lane 5). Thus, SDS-PAGE analysis indicates the high purity of the four PEGylated Saks and further confirms the results of SEC analysis.

**Structural characterization of PEGylated Saks**

CD analysis was used to investigate the secondary structure of Sak upon PEGylation. As shown in Fig. 4A and 4B, the far-UV
CD spectra (200–260 nm) of Sak show a single band with a maximum at 208 nm, indicating the rich β-sheet of Sak. The CD spectra of the four PEGylated Saks are approximately superimposed on that of Sak. This indicates that the secondary structure of Sak is essentially not influenced by PEGylation.

Intrinsic fluorescence was applied to detect the conformational changes of Sak upon PEGylation. When excited at 280 nm, Sak shows maximum fluorescence intensity at 350 nm (Fig. 4 C, D). The emission fluorescence intensity of the four PEGylated Saks is comparable to that of Sak without shift in the maximum intensity at 350 nm. Therefore, PEGylation do essentially not perturb the conformation of Sak.

Analytical ultracentrifugation

Sedimentation velocity analysis was performed to investigate the structure of the PEGylated Sak. The sedimentation coefficient ($S_{20,w}$) and the ratio of frictional coefficient ($f/f_0$) are summarized in Table 2. PEGylation can decrease the $S_{20,w}$ of Sak, due to the lower atom density of PEG chain relative to the protein core [29]. Moreover, Sak-mal20k and Sak-ald20k show lower $S_{20,w}$ values than Sak-mal5k and Sak-ald5k. This indicates that the $S_{20,w}$ of the PEGylated Sak is a function of the conjugated PEG mass. In addition, Sak-mal20k and Sak-mal5k show $S_{20,w}$ values comparable to those of Sak-ald20k and Sak-ald5k, respectively.

The ratio of frictional coefficient ($f/f_0$) is used to evaluate hydrodynamic shape of Sak [30]. Sak shows an $f/f_0$ of 1.20, indicating an almost spherical molecular shape. In contrast, PEGylation lead to the overall shape of Sak becoming geometrically asymmetric, as reflected by the increased $f/f_0$. Moreover, the increase in $f/f_0$ is dependent on the PEG mass, indicating that the conjugated PEG may affect the hydrodynamic shape of Sak.

Figure 6. RMSDs of the PEGylated Saks during the simulated annealing period. (A) Sak-mal5k; (B) Sak-ald5k; (C) Sak-mal20k, (D) Sak-ald20k.

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Figure 7. The conformations of PEGylated Saks after simulated annealing. (A) Sak-mal5k; (B) Sak-ald5k; (C) Sak-mal20k, (D) Sak-ald20k.

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comparable to those of Sak-ald20k and Sak-ald5k, respectively. Thus, the PEGylation sites do affect the hydrodynamic shape of the PEGylated protein.

**In vitro bioactivity**

The *in vitro* bioactivity of the products was measured by fibrin plate assay (Fig. 5). The relative bioactivities of the PEGylated Sak are lower than the unmodified Sak, presumably due to the steric shielding effect of PEG. Furthermore, Sak-mal20k and Sak-ald20k show lower bioactivities than Sak-mal5k and Sak-ald5k, respectively. This indicates that the steric shielding effect of PEG is dependent on the PEG mass. In addition, Sak-mal20k and Sak-mal5k show higher bioactivities than Sak-ald20k and Sak-ald5k, respectively. Presumably, N-terminus is close to the bioactive domain and C-terminus is far from it. Thus, the PEGylation site far from the bioactive domain may facilitate maintenance of the bioactivity of the PEGylated Sak. Atomic level investigation was conducted with the subsequent simulation research.

**Molecular dynamics simulation**

**Equilibration analyses.** Fig. 6 shows that the PEGylated Sak system reaches the equilibrium state at the end of molecular dynamics simulation. In addition, the RMSDs (root mean square deviation) of Sak-mal20k and Sak-ald20k show larger fluctuations than those of Sak-mal5k and Sak-ald5k, indicating the movement of the flexible PEG chain. Fig. S6 shows snapshots of PEGylated Saks at different time points. Fig. S7 shows that Sak’s secondary structures are apparently not altered with PEGylation.

Fig. 7 shows that 5 kDa PEG chains partially cover the Sak surface (Fig. 7 A, B) while 20 kDa PEGs almost completely cover the Sak surface (Fig. 7 C, D). In addition, the PEG chain is not found to wrap around Sak tightly or with a regular pattern, rather it loosely folds on the surface of the protein in an irregular form.

**Solvent accessible surface area (SASA) analyses.** Computed SASA of different entities are shown in Fig. 8. SASA is calculated between 20 and 25 ns. Sak-mal5k and Sak-ald5k both show SASAs larger than Sak and much lower than Sak-mal20k and Sak-ald20k. This indicates that the PEG chain can induce a large hydrated layer. C-terminally PEGylated Saks show larger SASAs than the N-terminal ones with identical PEG mass (Fig. 8A), indicating a larger hydrodynamic volume of C-terminally PEGylated Sak.

To further investigate the steric shielding effect of PEG on the interaction of Sak and miro-plg, we computed SASA of the eight amino acids (Lys1, Asp3, Tyr4, Met26, Asn28, Tyr33, Glu46 and Ile128) in the binding domain of Sak [31]. Figure S8 shows active site SASA fluctuation from 20 to 25 ns. The PEGylated Saks show decreased SASAs compared with the free Sak (Fig. 8B), indicating the presence of the steric shielding effect of PEG. Moreover, SASAs of C-terminally PEGylated Saks (7.56 nm² for Sak-mal5k and 6.02 nm² for Sak-mal20k) are larger than that of N-terminal products (5.20 nm² for Sak-ald5k and 3.67 nm² for Sak-ald20k). Lower SASA of the eight amino acids indicates an intensified steric shielding effect of PEG, which would retard the binding ability of micro-plg to Sak. Computed SASA suggests that the bioactivity of Sak is decreased by the steric shielding effect of PEG. Moreover, the steric shielding effect of PEG conjugated at N-terminus is stronger than that at C-terminus, in spite of the larger hydrodynamic volume of C-terminally PEGylated Sak. Thus, computational result agrees well with results of *in vitro* bioactivity assay (Fig. 5).

**Molecular size.** Dynamic light scattering was used to measure the molecular radii of the PEGylated Saks. The radii of Sak, Sak-mal5k, Sak-ald5k, Sak-mal20k and Sak-ald20k are 2.77, 3.71, 3.34, 5.43 and 5.12 nm, respectively. The molecular volumes (Ve) of the PEGylated Saks were thus calculated, assuming spherical shape of the PEGylated Saks (Table 3). The volumes at the end of simulation (Vs) of PEGylated Saks were also calculated

| Sample       | Ve (nm³) a  | Vs (nm³) b |
|--------------|-------------|------------|
| Sak          | 89          | 25.9       |
| Sak-ald5k    | 156         | 33.8       |
| Sak-mal5k    | 213.8       | 35.4       |
| Sak-ald20k   | 562.1       | 65.8       |
| Sak-mal20k   | 670.6       | 67.3       |

aVe: Molecular volumes measured by dynamic light scattering.
bVs: Molecular volumes calculated according to conformations of PEGylated Saks at the end of simulations.

Table 3. Molecular volumes of the PEGylated products.
by rolling a probe with a radius of 1.4 Å over the molecular surface (Table 3). Clearly, \( V_s \) values of the PEGylated Saks are smaller than their corresponding \( V_e \) values. This is due to the fact that \( V_e \) of the PEGylated Sak consisted of the volumes from Sak, PEG itself and the hydrated layer of PEG, whereas \( V_s \) of the PEGylated Sak lacks the volume from the hydrated layer of PEG. \( V_e \) and \( V_s \) are found to be strongly correlated with each other \( (R^2 = 0.9821) \), revealing the fact that MD simulation results correspond well to experimental data.

**Docking analyses.** Figure 9 shows final docked poses of PEGylated Saks and micro-plg. Table 4 shows the energy results of docking PEGylated Saks to micro-plg. The E-value represents the interaction energy between PEGylated Sak and its receptor (micro-plg). Lower E-value means higher stability of Sak-micro-plg complex and thus higher binding affinity of Sak to micro-plg. PEGylated Saks show higher E-values than free Sak, indicating that PEG hinders the binding of plasminogen to Sak. Moreover, Sak-mal5k and Sak-mal20k show lower E-values than Sak-ald5k and Sak-20k, respectively. This suggests that C-terminally PEGylated Saks show higher binding affinity to plasminogen than N-terminally PEGylated Saks, consistent to the experimentally assayed bioactivity of PEGylated Saks (Fig. 5).

### Discussion

PEGylation of therapeutic proteins (e.g., cytokines) often leads to substantial loss of bioactivity, presumably due to the steric shielding effect of PEG which interferes with the interaction of the protein and its high \( M_w \) receptor. However, the steric shielding effect of PEG is not adequately understood so far, particularly at the molecular level. Our present study aimed to reveal the steric shielding effect of PEG on the therapeutic protein by experimental and molecular simulation analyses.

The steric shielding effect of PEG on Sak was investigated on the aspects of PEG chain length and PEGylation site. Accordingly, four PEGylated Saks were prepared by site-specific conjugation of 5 kDa/20 kDa PEG to the N-terminus and the C-terminus of Sak, respectively. Aldehyde chemistry was used for N-terminal PEGylation of Sak [32,33] and site-specific PEGylation at the N-terminus was confirmed by tryptic peptide mapping analysis as described previously [18]. Maleimide chemistry was used for C-terminal PEGylation of Sak, which was achieved by PEGylation of the thiol group of the C-terminal Cys residue.

Structural characterizations suggest that the native conformation of Sak that underpins its bioactivity is essentially not altered upon PEGylation (Figs. 4 and S7). In contrast, the PEG chain length and the PEGylation site may alter its bioactivity (Fig. 5), hydrodynamic volume (Table 3) and molecular symmetry of the PEGylated Saks (Table 2).

Molecular dynamics simulation of the PEGylated Saks suggests that the PEG chain remains flexible at the equilibrium state. PEG can form a distinctive hydrated layer, which protects Sak, maintains its native conformation and increases its hydrodynamic volume. However, C-terminally PEGylated Saks show higher hydrodynamic volume than N-terminally PEGylated Saks, indicating higher flexibility of PEG in C-terminally PEGylated Sak that results in a larger hydrated layer. Presumably, the flexible PEG chain conjugated at the C-terminal domain of Sak is more loosely assembled on the Sak surface.

Simulation results (Fig. 7) show that PEG chain wraps around Sak and provides a steric shield around Sak. The steric shielding

![Figure 9. Final docked poses of PEGylated Saks and micro-plasminogen.](image)

| Sample     | E-value |
|------------|---------|
| Sak        | -552.0  |
| Sak-ald5k  | -334.9  |
| Sak-mal5k  | -380.6  |
| Sak-ald20k | -253.7  |
| Sak-mal20k | -340.2  |

**Table 4.** Docking results of PEGylated Saks with micro-Plasminogen.
The steric shielding effect of PEG

**Conclusions**

Wet lab assays and computational simulations were used to investigate the steric shielding effect of PEG on the conjugated Sak. Experimental study suggests that the native conformation of Sak is essentially not altered by PEGylation. In contrast, the bioactivity, the hydrodynamic volume and the molecular symmetric shape of the PEGylated Sak are altered and dependent on PEG chain length and PEGylation site. Molecular dynamics simulation of the PEGylated Saks suggests that the PEG chain remains highly flexible and forms a distinctive hydrated layer, resulting in the steric shielding effect of PEG on Sak. Docking analyses indicate that the binding affinity of Sak to its receptor also depends on PEG chain length and PEGylation site.

Computational simulation results agree well with experimental data. The present study provides clear molecular insight into the steric shielding effect of PEG on Sak at an atomic level. This type of study is essential to the rational design, fabrication and clinical application of PEGylated proteins, especially the ones involving interactions with high Mₘ substrates or receptors.

**Supporting Information**

**Figure S1** Snapshots of PEG chain (10 units) during MD. (A) Initial conformation: linear, force field: GROMOS_PEG; (B) Initial conformation: linear, force field: GROMOS_OE; (C) Initial conformation: coiled, force field: GROMOS_PEG; (D) Initial conformation: coiled, force field: GROMOS_OE.

**Figure S2** Snapshots of PEG-Sak MD simulation from different initial PEG conformations. (A) Sak-mal5k; (B) Sak-ald5k; (C) Sak-mal20k; (D) Sak-ald20k.

**Figure S3** Radii of gyration of PEG chains (10 units). (A) Sak-mal5k; (B) Sak-ald5k; (C) Sak-mal20k; (D) Sak-ald20k.

**Figure S4** Radii of gyration of PEG chains (20 units). (A) Sak-mal5k; (B) Sak-ald5k; (C) Sak-mal20k; (D) Sak-ald20k.

**Figure S5** PEG-Sak MD simulation from different PEGylated Saks during MD. (A) Sak-mal5k; (B) Sak-ald5k; (C) Sak-mal20k; (D) Sak-ald20k.

**Figure S6** Snapshots of PEG-Saks during MD. (A) Sak-mal5k; (B) Sak-ald5k; (C) Sak-mal20k; (D) Sak-ald20k.

**Figure S7** Secondary structures of PEGylated Saks during MD. (A) Sak-mal5k; (B) Sak-ald5k; (C) Sak-mal20k; (D) Sak-ald20k.

**Figure S8** Active site SASAs of PEGylated-Saks at the equilibrium state.

**File S1** PEG parameterization in GROMOS 53a6 force field.

**File S2** MD Cross-validation.

**Author Contributions**

Conceived and designed the experiments: TH JY. Performed the experiments: QM. Analyzed the data: QM TH JY. Contributed reagents/materials/analysis tools: TH JY. Wrote the paper: QM TH JY.

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