Combination of coarse-grained molecular dynamics simulations and small-angle X-ray scattering experiments

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Received June 21, 2019; accepted August 11, 2019

The combination of molecular dynamics (MD) simulations and small-angle X-ray scattering (SAXS), called the MD-SAXS method, is efficient for investigating protein dynamics. To overcome the time-scale limitation of all-atom MD simulations, coarse-grained (CG) representations are often utilized for biomolecular simulations. In this study, we propose a method to combine CG MD simulations with SAXS, termed the CG-MD-SAXS method. In the CG-MD-SAXS method, the scattering factors of CG particles for proteins and nucleic acids are evaluated using high-resolution structural data in the Protein Data Bank, and the excluded volume and the hydration shell are modeled using two adjustable parameters to incorporate solvent effects. To avoid overfitting, only the two parameters are adjusted for an entire structure ensemble. To verify the developed method, theoretical SAXS profiles for various proteins, DNA/RNA, and a protein-RNA complex are compared with both experimental profiles and theoretical profiles obtained by the all-atom representation. In the present study, we applied the CG-MD-SAXS method to the Swi5-Sfr1 complex and three types of nucleosomes to obtain reliable ensemble models consistent with the experimental SAXS data.

Key words: MD simulation, SAXS, protein solution structure, structural ensemble, nucleosome

Molecular dynamics (MD) are crucial to the structure and function of proteins [1]. To investigate the dynamic features of proteins, computational methods, such as MD simulations and normal mode analysis, are highly useful [1,2]. Simulation methods have also been utilized in combination with experimental data. Nuclear magnetic resonance (NMR) data are often incorporated into molecular simulations [3–6]. Small-angle X-ray scattering (SAXS) is another experimental tool for measuring macromolecular structures in solution. In contrast to NMR, SAXS has no limitation on the macromolecular size. However, the SAXS data are low in resolution, because they are an averaged quantity over rotational and conformational variants of solution structures. Therefore, a combination of simulation methods and SAXS has demonstrated successful modeling of detailed solution structures.

Protein dynamics are important for protein functions. To investigate protein dynamics, the combination of molecular dynamics (MD) simulations and small-angle X-ray scattering (SAXS) experiments, known as the MD-SAXS method, has been shown to be useful. However, all-atom MD simulations are limited to the microsecond time scale, which is significantly shorter than the typical time scale of large-scale protein motions. To overcome the time-scale limitation in the all-atom MD simulations, we propose a combined method of coarse-grained (CG) MD simulations and SAXS, termed the CG-MD-SAXS method, which was successfully applied to flexible and large biomolecules such as nucleosomes.
and a structure ensemble of proteins [7, 8]. To compare simulation results with SAXS data, theoretical SAXS profiles are calculated from 3D structures of proteins. Several methods (e.g., CRYSSOL [9], FoXS [10], AquaSAXS [11], the Zernike polynomials-based method [12], Fast-SAXS [13] and Fast-SAXS-pro [14]) have been proposed for the estimation of theoretical SAXS profiles from all-atom and CG models. In most of these methods, a theoretical SAXS profile is obtained from a single structure. For highly flexible proteins, a single structure is not sufficient to obtain theoretical SAXS profiles consistent with experimental profiles, because the experimental SAXS profile is an average of scattering from various conformations in solution. To handle a set of the structural ensemble of proteins, several methods (e.g., EOM [15], MES [16], EROS [17] and BSS-SAXS [18]) have been proposed. However, the unique determination of the structural ensemble is still difficult solely from SAXS data due to the limited information content. To overcome the difficulty originating from the flexibility of proteins, the incorporation of physicochemical methods such as MD simulations into the SAXS analysis is useful.

Water molecules contribute significantly to the SAXS profile. In the calculation of theoretical SAXS profiles, the following two major effects of water molecules should be considered: solvent-excluded volume and hydration water. In SAXS experiments, scattering from only the buffer solution is measured in addition to that from the protein solution. The experimental SAXS profile is obtained as the difference between two scattering intensities of the protein and buffer solutions. Thus, the excess electron density of proteins from the solvent molecules excluded by proteins becomes the source of the experimental SAXS profiles. Therefore, the X-ray scattering from the excluded solvent molecules should be considered while calculating theoretical SAXS profiles; this effect is referred to as the excluded-volume effect in this study. Another important effect of water on the SAXS profile is scattering from hydration water. Because the electron density of hydration water is higher than that of bulk water, the increment of scattering from hydration water significantly affects the SAXS profile. In the combination of all-atom MD simulations and SAXS [19–22] (e.g. the MD-SAXS method [19]), the MD simulation for the pure solvent is performed in addition to the simulation for the protein solution, and their theoretical scattering intensities are obtained. The theoretical SAXS profile is then calculated as the difference between the two scattering intensities, as done in actual experiments. Thus, two major solvent effects on SAXS profiles, i.e., the solvent-excluded volume of proteins and hydration water, are considered at the atomic level. In addition, since the electron density of bulk solvent depends on the ion concentration, ions in bulk significantly affect the SAXS profiles. In the all-atom MD-SAXS method, ion effects are also considered [23]. As an alternative approach of the explicit solvent model, the implicit solvent model has also been employed for the calculation of the theoretical SAXS profile [4, 9]. In CRYSSOL, the hydration water is modeled as the continuum hydration shell of proteins. The solvent-excluded volume is modeled as Gaussian spheres with effective radii. However, the estimation of the increment of electron density in the hydration shell and the determination of the effective radius of Gaussian spheres are difficult because they depend on the properties of protein surfaces, packing of protein interiors, and solvent compositions. Therefore, in CRYSSOL, two parameters, i.e., the increment of electron density in the hydration shell and the effective radius of Gaussian spheres, are adjusted for fitting to experimental SAXS profiles.

Although the all-atom treatment in MD simulations precisely describes protein dynamics and solvent effects, sufficient sampling is computationally expensive particularly for very large and/or very flexible proteins due to the long timescale of their dynamics. For such targets, coarse-grained (CG) models are often utilized to reduce computational burden [24]. For example, the CG-MD software CafeMol [25] employs a CG model in which one amino-acid residue is represented by one CG particle, and the effect of water molecules is implicitly incorporated by a continuum dielectric model. Owing to the CG representation, CG-MD simulations can be performed in a substantially fast computation; the combination of CG-MD simulations and SAXS will thus be efficient, particularly for the sampling of large and flexible proteins. Therefore, in this study, we propose a method of combining CG-MD and SAXS, referred to as the CG-MD-SAXS method.

To develop the CG-MD-SAXS method, due to the implicit treatment of water molecules, solvent effects on SAXS profiles should be modeled in a CG manner. As in CRYSSOL, two parameters, i.e., the increment of electron density in the hydration shell and the effective radius of Gaussian spheres, are adjusted in the CG-MD-SAXS method. However, if the two parameters are adjusted for every snapshot in MD trajectories, a large number of parameters are required in total with respect to one experimental SAXS profile. Owing to limited SAXS resolution, a large number of adjustable parameters should be avoided. In this study, we propose a method to overcome this problem. First, in our method, six terms comprising the CG-MD-SAXS profile are independently evaluated from MD trajectories. Second, the six terms are combined into one CG-MD-SAXS profile using the abovementioned two adjustable parameters. Therefore, only these two adjustable parameters of solvent effects are used for an entire MD trajectory. In this study, it is shown that these two parameters are sufficient for solvent modeling in the calculation of CG-MD-SAXS profiles for trajectories.

For CG-MD simulations, the software CafeMol was employed. In CafeMol, one amino acid is represented as one CG particle for proteins, as described above. For nucleic acids, the base, sugar and phosphate of nucleotides are represented as three different CG particles, respectively. First, the scattering factors of the CG particles for 20 amino acids,
5 bases, 2 sugars (ribose/deoxyribose), and 1 phosphoric acid were evaluated from atomic scattering factors by averaging structural data in the Protein Data Bank (PDB). Next, the hydration model in a CG manner was developed for the CG-MD-SAXS analysis. In the hydration model, only the two adjustable parameters were employed for an entire trajectory. The performance of the CG-MD-SAXS method was examined by comparisons with the all-atom MD-SAXS analysis and experimental data. Additionally, the position dependencies of CG particles for theoretical SAXS profiles were examined. Four positions representing an amino acid were evaluated from atomic scattering factors by averaging structural data because the excluded volume depends on the packing of atoms.

Methods

Scattering factor of CG particles

The excess scattering intensity of a CG-model protein in dilute solution, $I_C^{\text{CG}}(q)$, is defined by the Debye formula as

$$I_C^{\text{CG}}(q) = \sum_{i=1}^{N-M} f_i^{\text{CG}}(q) f_j^{\text{CG}}(q) \frac{\sin(qr_{ij})}{qr_{ij}},$$  \hspace{1cm} (1)

where $q$ is the wave vector transfer defined as $q = 4\pi \sin(\theta)/\lambda$ (2$\theta$ is the scattering angle, and $\lambda$ is the X-ray wave length), $N$ is the number of CG particles for proteins, $M$ is the number of grid points in the hydration shell, $f_i^{\text{CG}}(q)$ is a scattering factor for the $i$-th CG particle or grid, and $r_{ij}$ is the distance between the $i$-th and $j$-th particle or grid. The scattering factor $f_i^{\text{CG}}(q)$ can be expressed by

$$f_i^{\text{CG}}(q) = f_i^{\text{CG}, \text{vac}}(q) - f_i^{\text{CG}, \text{ex}}(q) \text{ for } i = 1, ..., N,$$  \hspace{1cm} (2)

where $f_i^{\text{CG}, \text{vac}}(q)$ is the scattering factor of CG particle $i$ in vacuo, and $f_i^{\text{CG}, \text{ex}}(q)$ is the factor of the solvent-excluded volume. The scattering factor for the hydration shell can be expressed by

$$f_i^{\text{CG}}(q) = f_i^{\text{CG}, \text{hyd}}(q) \text{ for } i = N+1, ..., M,$$  \hspace{1cm} (3)

where $f_i^{\text{CG}, \text{hyd}}(q)$ is the factor of the contrast of electron densities between the hydration shell and the buffer solution.

Scattering factors of CG particles in vacuo and solvent-excluded volume

The scattering factor of CG particle $k$ representing an amino acid, base, sugar or phosphate in vacuo, $f_k^{\text{CG}, \text{vac}}(q)$, was calculated from the averaged scattering intensity $I_k^{\text{CG}, \text{vac}}(q)$ of the corresponding moiety in the PDB data as follows:

$$f_k^{\text{CG}, \text{vac}}(q) = \left\langle f_k^{\text{CG}, \text{vac}}(q) \right\rangle_{\text{PDB}}^{1/2} = \left( \frac{\sum_{j=1}^{N} f_j^{\text{vac}}(q) f_j^{\text{vac}}(q) \frac{\sin(qr_{jk})}{qr_{jk}}}{\sum_{j=1}^{N} f_j^{\text{vac}}(q) f_j^{\text{vac}}(q) \frac{\sin(qr_{jk})}{qr_{jk}}} \right)^{1/2},$$  \hspace{1cm} (4)

where $f_j^{\text{vac}}(q)$ is an atomic scattering factor in vacuo of the $i$-th atom belonging to CG particle $k$. In this calculation, we used a set of high-resolution crystal structures including hydrogen atoms as described in the last paragraph of this section.

Next, the factors of the solvent-excluded volume of CG particles are examined. First, the treatment of the solvent-excluded volume in an all-atom model is briefly described. The modified atomic scattering factor [26], $f_k^{\text{vac}, \text{ex}}(q)$, in which the effect of the solvent-excluded volume is incorporated, is defined as

$$f_k^{\text{vac}, \text{ex}}(q) = f_k^{\text{vac}}(q) - v_i \rho_s \exp \left( \frac{-q^2 r_{ij}^2}{4\pi} \right),$$  \hspace{1cm} (5)

where the second term on the right-hand side represents the effect of the excluded volume by a Gaussian sphere, $v_i$ is the atomic volume of the $i$-th atom, and $\rho_s$ is the electron density of the bulk solvent. Using the modified atomic scattering factor, the excess scattering intensity, $I_k^{\text{vac}, \text{ex}}(q)$, of a protein is then calculated by

$$I_k^{\text{vac}, \text{ex}}(q) = \sum_{i,j} f_i^{\text{vac}, \text{ex}}(q) f_j^{\text{vac}, \text{ex}}(q) \frac{\sin(qr_{ij})}{qr_{ij}}.$$  \hspace{1cm} (6)

This formula is used in CRYSSOL, and $v_i$ is scaled so that the theoretical SAXS data are fitted to the experimental SAXS data because the excluded volume depends on the packing density of the protein interior.

Yang, S., et al. extended Eq. (5) to obtain the modified CG scattering factor, $f_k^{\text{CG}, \text{vac}, \text{ex}}(q)$ including the excluded volume [13]. The modified CG scattering factor is defined by a knowledge-based formula as

$$f_k^{\text{CG}, \text{vac}, \text{ex}}(q) = \left\langle f_k^{\text{CG}, \text{vac}, \text{ex}}(q) \right\rangle_{\text{PDB}}^{1/2} = \left( \frac{\sum_{i,j \in k} f_i^{\text{ex}}(q) f_j^{\text{ex}}(q) \frac{\sin(qr_{ij})}{qr_{ij}}}{\sum_{i,j \in k} f_i^{\text{ex}}(q) f_j^{\text{ex}}(q) \frac{\sin(qr_{ij})}{qr_{ij}}} \right)^{1/2}.$$  \hspace{1cm} (7)

In this approach, the parameters for excluded volumes such as $v_i$ and $\rho_s$ must be determined before the evaluation of $f_k^{\text{CG}, \text{vac}, \text{ex}}(q)$, and these parameters cannot be modified after evaluation of $f_k^{\text{CG}, \text{vac}, \text{ex}}(q)$ because the excluded-volume term is already averaged over many conformations in the PDB. Therefore, we introduced another definition of the modified CG scattering factor including the solvent-excluded volume.

In our method, to modify the parameters for the excluded volume after evaluating the atomic scattering factors in vacuo and that of the excluded volume, we defined a CG scattering factor $f_k^{\text{CG}, \text{vac}, \text{ex}}(q)$ by introducing the excluded volume of CG particles explicitly as
where the second term on the right-hand side represents the excluded volume of i-th CG particle, \( v_i^{CG} \) is the volume of i-th CG particle defined as the total volume of atoms in an amino acid, base, sugar or phosphate represented by a CG particle. The radius of the Gaussian sphere, \( \alpha_i \), is determined by minimization of \( S_i \):

\[
S_i = \frac{1}{N} \sum_{j} \left( v_i^{CG} - v_i^{CG, \text{ex}}(q,j) \right)^2.
\]

To control the amount of the excluded-volume effect sampled by CG-MD simulations, \( v_i^{CG, \mu} \) was scaled after the evaluation of \( f_i^{CG}(q) \).

The determination of \( f_i^{CG, \text{ex}}(q) \) was conducted using high-resolution crystal structures of the following 5 proteins, 2 DNA structures, and 2 RNA structures. The proteins were serine protease (PDB ID: 1GCI [27], 0.78 Å resolution), xylose isomerase (PDB ID: 1MUW, 0.86 Å resolution), trypsin (PDB ID: 1PQ7 [28], 0.8 Å resolution), \( P. \) abyssi rebredoxin (PDB ID: 2PYA [29], 0.86 Å resolution) and HEW lysozyme (PDB ID: 2VB1 [30], 0.65 Å resolution). The nucleic acids were DNA (PDB ID: 1BNA [31], 1.9 Å resolution), RNA (PDB ID: 1ION [32], 0.99 Å resolution), and a CAG RNA repeat (PDB ID: 3NJ6 [34], 0.95 Å resolution). Using Eq. (4), \( f_i^{CG, \text{ex}}(q) \) for each CG particle corresponding to 20 types of the amino acid was determined. For nucleotides, the 3SPN.1 model [35,36] was used as the CG representation. In the model, the sugar, phosphate, and base were represented as three different CG particles, and the determination of \( f_i^{CG, \text{ex}}(q) \) for the CG particles was performed using guanine, adenine, cytosine, thymine, uracil, ribose, deoxyribose and phosphate in DNA/RNA. Using Eq. (9), the radius of the Gaussian sphere, \( \alpha_i \), for each type of CG particle was determined.

### Scattering from the hydration shell

Next, scattering from the hydration shell was examined. The water density of the hydration shell of proteins is higher than that of the bulk solvent. The contrast of the electron densities between the hydration shell and the bulk solvent were incorporated in the calculation of the excess scattering intensity. In our method, the effects of the hydration shell were incorporated by generating grids of the high-density region around the protein, as follows: A cubic lattice with a spacing of \( \Delta D \) was generated in the vicinity of proteins. A unit cell of the lattice (\( \Delta D^3 \)) was set such that a water molecule can be fit in a cell. Then, the grid points of the hydration shell were selected as the points that were satisfied with the range of \( r_g + r < r < r_g + r_i \) over all the CG particles. In this instance, \( r \) is the distance of the grid point from a CG particle, \( r_g \) is the radius of the CG particle, \( r_w \) is the radius of a water molecule, and \( r_i \) is the thickness of the hydration shell.

In the present calculations, \( \Delta D, r_g \) and \( r_w \) were set at 3 Å, 1.5 Å and 4.5 Å, respectively. For each grid point in the hydration shell, the scattering factor \( f_i^{CG, \text{hyd}}(q) \) is assigned. The scattering factor represents the contrast of the electron densities between the hydration shell and the buffer solution as

\[
f_i^{CG, \text{hyd}}(q) = w \Delta D f_w^{CG}(q),
\]

where \( i \) represents the i-th grid point, \( w \) is the ratio of the electron-density increase in the hydration shell from the buffer solution, and \( f_w^{CG}(q) \) is the scattering factor of a water molecule in an all-atom representation (one oxygen atom and two hydrogen atoms) given by

\[
f_w^{CG}(q) = \left(\frac{\sum_{i} f_i^{CG}(q) \sin(qr_i)}{qr_i}\right)^{1/2}.
\]

To incorporate the amount of the contrast of the electron-density in a CG manner, the parameter \( w \) is scaled after evaluation of \( f_i^{CG}(q) \).

### Computational details of the scattering intensity from CG models

The excess scattering intensity of a CG model, \( f_i^{CG}(q) \), including effects of the excluded volume and the hydration shell was developed by Eqs. (1)–(4), (8) and (10). In the CG-MD-SAXS method, \( f_i^{CG}(q) \) is averaged over a large number of snapshot structures in CG-MD trajectories. The two parameters of \( v_i^{CG, \mu} \) and \( w \) defined in Eqs. (8) and (10) are adjustable parameters. For computational convenience, we introduced a scaling factor \( c_w \) by replacing \( v_i^{CG, \mu} \) in Eq. (8). Taken together, Eq. (1) can be rewritten as

\[
F_i^{CG}(q) = I_{i}^{CG}(q) + c_w I_{w}^{CG}(q) + c_{w} I_{w}^{CG}(q) + w I_{w}^{CG}(q).
\]

In this equation, \( I_{i}^{CG}(q) \) is expressed as

\[
I_{i}^{CG}(q) = \sum_{i \neq j} (\mathbf{2} - \mathbf{\delta}_{ij}) I_{i}^{CG}(q) f_{j}^{CG}(q) \sin(qr_{ij}) \frac{1}{qr_{ij}},
\]

\[
I_{w}^{CG}(q) = \sum_{i} -2 \mu f^{CG}(q) v_i^{CG} \exp(-\frac{\alpha_i q^2}{4\pi}) \sin(qr_{i}),
\]

\[
I_{w}^{CG}(q) = \sum_{i} (\mathbf{2} - \mathbf{\delta}_{ij}) \mu^2 v_i^{CG} \exp(-\frac{\alpha_i q^2}{4\pi}) \sin(qr_{i}) \frac{1}{qr_{i}},
\]

\[
I_{w}^{CG}(q) = \sum_{i} \sum_{j} 2 I_{i}^{CG}(q) \Delta D f_{w}^{CG}(q) \sin(qr_{ij}) \frac{1}{qr_{ij}},
\]

\[
I_{w}^{CG}(q) = \sum_{i} \sum_{j} -2 \mu v_i^{CG} \exp(-\frac{\alpha_i q^2}{4\pi}) \Delta D f_{w}^{CG}(q) \sin(qr_{ij}) \frac{1}{qr_{ij}}.
\]
\[ I_{\text{CG}}(q) - I_{\text{sol}}(q) \] can be averaged over MD trajectories as

\[ I_{\text{CG-MD}}(q) = (I_{\text{CG}}(q))_{\text{MD}} \]

\[ = I_{1_{\text{CG}}}(q)_{\text{MD}} + c_{\text{w}} I_{2_{\text{CG}}}(q)_{\text{MD}} + c_{\text{w}} I_{3_{\text{CG}}}(q)_{\text{MD}} + w I_{4_{\text{CG}}}(q)_{\text{MD}}. \]

(14)

Next, \( c_{\text{w}} \) and \( w \) were searched through a specified range to minimize \( \chi^2 \) defined by

\[ \chi^2 = \frac{1}{N_{\mu}} \sum_{\mu=1}^{N_{\mu}} \frac{(I_{\text{exp}}(q) - I_{\text{calc}}(q))^2}{\sigma_{\text{exp}}(q)} \]

\[ c = \left( \frac{1}{N_{\mu}} \sum_{\mu=1}^{N_{\mu}} \frac{I_{\text{exp}}(q) I_{\text{calc}}(q)}{\sigma_{\text{exp}}(q)^2} \right)^{-1} \]

(16)

where \( N_{\mu} \) is the number of \( q \), \( I_{\text{exp}}(q) \) is the experimental scattering intensity, \( I_{\text{calc}}(q) \) is a trial intensity of Eq. (14) with fixed parameters in the search, and \( \sigma_{\text{exp}}(q) \) is the experimental error. The search ranges of \( c_{\text{w}} \) and \( w \) were set at \( 0.9 \leq c_{\text{w}} \leq 1.1 \), and \( 0.0 \leq w \leq 0.01 \), respectively. These ranges are similar to those used in CRYSTOL; the parameter of the excluded volume is searched up to a 10% decrement/increment, and the contrast in the hydration shell is searched up to a 27% corresponding to \( w = 0.01 \).

Comparison of the intensities calculated from CG-MD-SAXS, CRYSTOL, and all-atom MD-SAXS

To examine the performance of the CG-MD-SAXS method, the scattering intensity of a crystal structure calculated using the CG-MD-SAXS method was compared to that calculated using CRYSTOL in an all-atom manner and that calculated using the all-atom MD-SAXS for HEW lysozyme (PDB ID: 6LYZ [37]) with explicit water. The experimental profile was included in the CRYSTOL package. When the CG-MD-SAXS method was applied to the crystal structure, an amino acid was represented as one CG particle placed at the center of electrons. The calculation of CRYSTOL was performed using the default setting. When the all-atom MD-SAXS was applied, the excess scattering intensity of lysozyme was calculated as in the following protocol. In this report, the procedure is described only briefly. See reference [19] for details. Two all-atom MD simulations of both a protein solution and a pure solvent were carried out independently. From the trajectory of the protein solution, a spherical region including the protein and hydration shell was extracted, and the region was used in the calculation of the scattering intensities. In the present calculation, a radius of 36 Å from the center of mass was employed. Next, the scattering intensities of snapshots in the trajectory were averaged. To extract the scattering intensity of the pure solvent, a spherical region of the same size region determined in the protein solution was extracted from the trajectory of the pure-solvent MD simulation, and the scattering intensities for snapshots were averaged. The excess scattering intensity was then obtained by subtracting the two scattering intensities.

All-atom MD simulations were carried out with the MD program MARBLE [38] using the CHARMM27/CMAP force field [39,40] and the TIP3P water model [41]. The solution system was prepared by placing the crystal structure at a cubic cell and adding a 20 Å water molecules. No counterions were added. A 1-ns production run was performed in the isothermal-isobaric (NPT) ensemble at 1 atm and 293 K. Before the production run, a 100 ps heating simulation from 10 K to 293 K with harmonic constraints on the heavy atoms of the solute in the NVT ensemble and a 100 ps simulation for gradually removing the constraints in the NPT ensemble were sequentially performed. In addition, the MD simulations of the pure solvent were performed under the same procedure as the protein solution.

Comparison of CG-SAXS profiles for various structures and CG models

For the performance test of the CG-MD-SAXS method, the scattering intensities of various proteins, DNA/RNA, and a protein-RNA complex were calculated with their fixed structures. The intensities were compared with those calculated by CRYSTOL and their experimental profiles. The experimental SAXS data were taken from BIOISIS.net (http://www.bioisis.net/). The examined proteins are Immunoglobulin-like domains 1 and 2 of the protein tyrosine phosphatase LAR3 (BIOISIS ID: LAR12P [42], PDB ID: 3PKJ), Superoxide dismutase (BIOISIS ID: APSODP [43], PDB ID: 1HL5 [44]), Ubiquitin-like modifier-activating enzyme ATG7 C-terminal domain (BIOISIS ID: ATG7CP [45], PDB ID: 3T7E [45]), DNA double-strand break repair protein MRE11 (BIOISIS ID: MRERAP [46], PDB ID: 3AV0), glucose isomerase (BIOISIS ID: GISRUP, PDB ID: 2G4J [47]), and Complement C3b+Efb-C (BIOISIS ID: C3BEFP [48]). The examined DNA/RNA and complexes are 28 base-pairs of DNA (BIOISIS ID: 28BPDD), the P4–P6 RNA ribozyme domain (BIOISIS ID: 1P4P6R [49], PDB ID: 1GID [50]), and A. aeolicus MnmG bound to tRNA (BIOISIS ID: MNMG2X [51]).

To examine the choice of CG particle placement, four CG models for proteins and two CG models for nucleic acids were compared. For proteins, a CG particle represents an amino acid, and there are four different variations in terms of the locations of the CG particle: the center of electrons (CME model), Cα position (CA model), Cβ position (CB model) and the center of the side chain of each residue (CMS model). For nucleic acids, three different CG particles represent a base, sugar and phosphate, respectively, and there are two different variations about the locations. The first is that the CG particles are located at the centers of electrons of each moiety (CME model). The other model is the 3SPN.1 model employed in CafeMol. In the 3SPN.1 model, the CG particles for the sugar and phosphate are placed at the center
of mass for each moiety. The CG particle for purine bases (adenine and guanine) is located at the N1 position, and that for pyrimidine bases (cytosine and thymine) is located at the N3 position. Using various proteins, DNA/RNA, and the protein-RNA complex described above, CG-SAXS profiles for the different models were calculated and were compared to the experimental SAXS data and the all-atom SAXS profile calculated using CRYSOL.

CG-MD-SAXS analysis for a flexible Swi5-Sfr1 protein complex

The Swi5-Sfr1 protein complex was employed in the CG-MD-SAXS analysis. The crystal structure of Swi5 and the C-terminal domain of the Sfr1 complex, referred to as Swi5-Sfr1C, (PDB ID: 3VIQ [52]) was used. The mutated residues 170-180 of Sfr1 and the missing side chain of residue 299 of Sfr1 were replaced with wild-type residues using the program MODELLER [53].

All the CG-MD simulations were performed using the program CafeMol with the atomic interaction-based CG2+ (AICG2+) model [54] for the energy potential of proteins. One residue was coarse-grained as one CG particle, located at the Ca position (CA model). For disordered regions, the flexible local potential (FLP) [54] was used. The electrostatic interaction was handled by the Debye-Hückel type interactions. The ion concentration was set at 200 mM corresponding to the same conditions used in the SAXS experiments [52]. The time evolution was handled by Langevin dynamics at 300 K. Changing the range of the disordered region modulated by the FLP, a 107-step CG simulation was independently repeated 10 times in each calculation condition. Snapshots were saved at every 1,000th step in one simulation, the latter half of the simulation was used in the CG-MD-SAXS analysis (5,000 snapshots in one simulation), and the snapshots of 10 different simulations were merged (50,000 snapshots in total). Using the merged snapshots, a theoretical SAXS profile was calculated. The experimental SAXS profile from q=0.015 to 0.25 was used.

CG-MD-SAXS analysis for large and flexible nucleosomes

Three types of nucleosomes were employed: the canonical nucleosome, the CENP-A nucleosome, and the H2A.B nucleosome. The initial structures of the three nucleosomes were prepared by modeling processes, as follows. The crystal structure of the canonical nucleosome and the CENP-A nucleosome were used; their PDB IDs were 3AFA [55] and 3AN2 [56], respectively. All missing histone tails were added by MODELLER. In the canonical nucleosome, 1 base pair was added to the edge of DNA (11, J292 in 3AFA), and then the DNA sequence was changed to that of the SAXS experiment (147 base pairs) [56] using the MMTSB tool [57]. In the crystal structure of the CENP-A nucleosome, both ends of DNA (26 base pairs in total) were short compared with the DNA used in the SAXS experiment. Because the DNA sequence used in the SAXS experiment for the CENP-A nucleosome is the same as that of the SAXS experiment for the canonical nucleosome, the missing part of DNA was transplanted from those of the modeled canonical nucleosome. The modeled canonical nucleosome was superimposed into the CENP-A nucleosome such that the root mean square deviation (RMSD) values of H2A/B and H4 were minimized, and then the superimposed coordinates of both ends of DNA of the canonical nucleosome were added to the missing part of DNA of the CENP-A nucleosome. Because no crystal structures of the H2A.B nucleosome have been reported to date, homology modeling was performed using the modeled canonical nucleosome as a reference structure: H2A in the canonical nucleosome was changed to H2A.B. Since the DNA sequence of the H2A.B nucleosome used in SAXS experiments [58] was 145 base pairs, each 1 base pair at both ends of DNA was erased from the modeled nucleosome, and the DNA sequence (145 base pairs) was changed to those used in SAXS experiments [58] using the MMTSB tool. After the modeling processes described above, all modeled structures of canonical, CENP-A, and H2A.B nucleosomes were relaxed in solution by a 2000 step minimization (steepest descent method) by MARBLE with the CHARMM27/CMAP force field and the TIP3P water model. Counterions were added to keep the system charge-neutral.

The CG-MD simulations for the nucleosomes were performed using the program CafeMol. One residue in proteins was represented as one CG particle by the CA model, and one base in DNA was represented as three CG particles by the 3SPN.1 model. The AICG2+ model was used as a potential for proteins, and the FLP was used for the CG particles in the histone tails. For the energy potential of DNA, both Gō potential and physical interactions of base pairing were used, and the interaction between DNA and proteins was handled by the Gō potential. In the simulations of the CENP-A nucleosome, the interaction between both ends of DNA and proteins was not incorporated since both ends of DNA were missing in the crystal structure and might be flexible [58]. The electrostatic interaction was handled by the Debye-Hückel type interactions. For the structural investigation, CG-MD simulations were performed by changing both the strength of the Gō potential of the interactions among proteins and DNA, and the ion strength. The strength of the Gō potential was changed by introducing the scaling coefficient, and the range of the scaling was 0.4, 0.6, 0.8, and 1.0. The range of the ion strength was 0, 0.5, 1, 2.5, 5, 10, 25, 50, 100, 200, 300, 400, 500, 600, 700, 800, 900 and 1000 (unit is mM). In each calculation condition, a 107-step simulation at 300 K was performed and was independently repeated 100 times. Snapshots were saved at every 10,000th step in one simulation, the latter half of the simulation was used in the CG-MD-SAXS analysis (500 snapshots in one simulation), and the snapshots of 100 different simulations were merged (50,000 snapshots in total). Using the merged snapshots, a theoretical SAXS profile was obtained. The experimental
Comparison of intensities calculated by the all-atom model and the CG model

To examine the accuracy of our method, the theoretical scattering intensities calculated from the CG-MD-SAXS method, CRYSOl and the all-atom MD-SAXS method, as well as the experimental profile were compared for the HEW lysozyme crystal structure (PDB ID: 6LYZ), which was not used in the determination of the scattering factor of CG particles. In the CG-MD-SAXS method, each amino acid was represented as one CG particle placed at the center of electrons, and the theoretical profile was calculated from the CG model using the CG-MD-SAXS method. For comparison, the theoretical profile of the all-atom model of the crystal structure was calculated using CRYSOl. In addition, a 1-ns trajectory of the all-atom MD simulation with explicit water was prepared, and the theoretical profile was calculated from the MD trajectory using the all-atom MD-SAXS method.

Four SAXS profiles are shown in Figure 1A. The theoretical scattering intensities are in good agreement with each other. In addition, the discrepancy between the theoretical and experimental curves, $\chi$, showed a similar value in the three methods. These results indicate that the CG-MD-SAXS method is accurate and that the CG-MD-SAXS profile is largely consistent with those calculated in an all-atom manner. Because the solvent effect is explicitly incorporated in the all-atom MD-SAXS method, the agreement of the profiles between the all-atom MD-SAXS and CG-MD-SAXS methods indicates that the current CG treatment of the solvent effects is appropriate to calculate SAXS profiles.

Components of the scattering intensity were compared between CG and all-atom models at a fixed structure. The comparison was performed using the intensity calculated from CRYSOl since the solvent effect was implicitly incorporated in CRYSOl, as was the case in the CG-MD-SAXS method. The scattering intensities for amino acids and bases, $I_{\text{CG, vac, ex}}^{\text{CG-MD-SAXS}}(q)$ and $I_{\text{CG, vac, ex}}^{\text{CRYSOl}}(q)$, are in good agreement with each other, suggesting that the scattering factors of the CG particles were successfully determined. In our formulation, the term for the excluded volume can be scalable through $v_i^{\text{ex}}\rho$, after the determination of the atomic scattering factors. This property is advantageous over the formulation of Yang, S., et al. The scattering factor of a water molecule given by Eq. (11) was also determined (Supplementary Fig. S1C).

The scattering intensities for various structures and CG models

To examine the dependence of CG models, theoretical profiles for various proteins, DNA/RNA, and a protein-RNA complex were calculated using the various CG models. In the CG models for proteins, the locations of the CG-particles were different: the center of electrons (CME model), Cα position (CA model), Cβ position (CB model), and the center of the side chain of each residue (CMS model). For nucleic acids, the three CG particles represent a base, sugar, and phosphate, respectively, and the CME model and the 3SPN.1 model were examined. The $\chi$ values for all the
CG-MD-SAXS analysis of the Swi5-Sfr1C complex

The Swi5-Sfr1 complex is an activator of Rad51-mediated strand exchange in homologous recombination [59,60]. The Swi5-Sfr1 complex is bound directly to the Rad51 filament and stabilizes it [60]. According to Kuwahara, N., et al. [52], a truncated Sfr1 lacking the N-terminal intrinsically disordered region forms a complex with Swi5, and the complex, known as Swi5-Sfr1C, still has the ability to stimulate strand exchange. X-ray crystallography and SAXS measurements of Swi5-Sfr1C [52] showed that the envelope of the dummy atoms calculated from the SAXS data was essentially similar to the crystal structure (PDB ID: 3VIQ [52]). However, two regions protruded from the envelope (Fig. 3A). One region is the β-sheet region of Sfr1 (250–270 residues), while the other is the helical structure of the C-terminal region of Swi5 (70–85 residues); the regions were termed βSfr1 and CSwi5, respectively. In the crystal structure, βSfr1 interacts with the...
ensemble including disordered $\beta_{Sfr1}$ was close to that observed in the experiment (Fig. 5). The $\chi$ value of the profile of $\beta_{DCO}$ was the smallest, indicating good agreement with the experimental profile. Therefore, $\beta_{Sfr1}$ was presumably disordered in solution unlike the crystal structure. Indeed, $\beta_{Sfr1}$ interacts with other molecules in the crystal (Fig. 3B), and the $\beta$-sheet structure may be an artifact of crystal packing. Interestingly, the $\chi$ value of the theoretical CG-SAXS profile obtained from the structural ensemble was smaller than that obtained from the crystal structure by CRYSOL. This result indicates the importance of structural flexibility in solution. The CG-MD-SAXS analysis can provide an explicit view of the structural ensemble for flexible proteins including disordered regions.

CG-MD-SAXS analysis of three types of nucleosomes

The nucleosome is the elemental unit of chromatin in eukaryotes [55]. The nucleosome is a complex of core histones and DNA. In the canonical nucleosome, the core histones are H2A, H2B, H3, and H4, and two of the H2A-H2B other dimer (Fig. 3B). In contrast, C$^{\text{Swi5}}$ interacts with its own core-domain.

To examine how conformations of $\beta_{Sfr1}$ and C$^{\text{Swi5}}$ adopt in solution, the CG-MD-SAXS analysis was performed. To simulate ordered/disordered conformations of the regions, the flexible local potential (FLP) was employed in CG-MD simulations. When the FLP is activated, the residues in the region move freely as disordered conformations. Using the FLP, three different simulations were carried out: both ordered $\beta_{O CO}$ and C$^{\text{Swi5}}$ ($\beta_{O CO}$), disordered $\beta_{O CO}$/ordered C$^{\text{Swi5}}$ ($\beta_{DCO}$) and ordered $\beta_{O CO}$/disordered C$^{\text{Swi5}}$ ($\beta_{O CD}$). The RMSD of $\beta_{O CO}$ and C$^{\text{Swi5}}$ of the three simulations are illustrated in Figure 4A and 4B, respectively. The final structures in the simulations are shown in Figure 4C. The RMSD of $\beta_{O CO}$/C$^{\text{Swi5}}$ with the FLP is larger than those without the FLP, suggesting that the difference between ordered/disordered conformations was sufficiently large. Theoretical SAXS profiles were then calculated from the ensembles using the CG-MD-SAXS method.

The theoretical SAXS profiles showed that the structural ensemble including disordered $\beta_{O CO}$ was close to that observed in the experiment (Fig. 5). The $\chi$ value of the profile of $\beta_{O CO}$ was the smallest, indicating good agreement with the experimental profile. Therefore, $\beta_{O CO}$ was presumably disordered in solution unlike the crystal structure. Indeed, $\beta_{O CO}$ interacts with other molecules in the crystal (Fig. 3B), and the $\beta$-sheet structure may be an artifact of crystal packing. Interestingly, the $\chi$ value of the theoretical CG-SAXS profile obtained from the structural ensemble was smaller than that obtained from the crystal structure by CRYSOL. This result indicates the importance of structural flexibility in solution. The CG-MD-SAXS analysis can provide an explicit view of the structural ensemble for flexible proteins including disordered regions.

Figure 3 Crystal structure and the dummy residue model of the Swi5-Sfr1C complex. (A) A dummy residue model (magenta) and the crystal structure of the Swi5 (cyan) and Sfr1C (green) complex. (B) Two heterodimers (cyan-green and yellow-magenta) of the Swi5-Sfr1C complex in the asymmetric unit.

Figure 4 RMSD of $\beta_{O CO}$ and C$^{\text{Swi5}}$ in three CG-MD simulations, and final structures. (A) RMSD of $\beta_{O CO}$ in three simulations, $\beta_{O CO}$ (red), $\beta_{O CD}$ (blue) and $\beta_{O CD}$ (orange). (B) RMSD of C$^{\text{Swi5}}$ in three simulations, $\beta_{O CO}$ (red), $\beta_{O CD}$ (blue), and $\beta_{O CD}$ (orange). (C) Final structures of three simulations, $\beta_{O CO}$ (red), $\beta_{O CD}$ (blue), and $\beta_{O CD}$ (orange).

Figure 5 Experimental and theoretical scattering intensities of the Swi5-Sfr1C complex. The experimental intensity is represented by gray dots, and the theoretical intensity of the crystal structure obtained using CRYSOL is represented by a green line. Theoretical intensities obtained from $\beta_{O CO}$, $\beta_{O CD}$, and $\beta_{O CD}$ simulations are represented by red, blue, and orange lines, respectively.

Figure 6 Theoretical scattering intensities of the Swi5-Sfr1C complex. The theoretical intensity is represented by red, blue, and orange lines, respectively.
all three nucleosomes. The theoretical SAXS profiles under the condition are shown in Figure 6. All theoretical profiles are in good agreement with the experimental profiles.

The CG-MD-SAXS analysis revealed an explicit view of the solution structures of the nucleosome satisfied with the experimental SAXS profile. In CG-MD simulations, both ends of DNA moved toward the outside of the histone core and fluctuated between the distant and close positions from the histone core. To analyze how the DNA fluctuates, the separation of DNA from the histone core was characterized by a minimum distance between the end of DNA and the histone core. The free energy representation of the DNA fluctuation against two distances is shown in Figure 7. Structures of the canonical, CENP-A, and H2A.B nucleosomes at the free energy minimum are illustrated in Figures 8A, 8C and 8D, respectively. The free-energy distributions were different from each other. In the canonical nucleosome, the structures were almost distributed around the free energy minimum (20 Å, 20 Å) and showed an L-shaped distribution. By contrast, in the CENP-A nucleosome, the free-energy distribution around the minimum (43 Å, 38 Å) was wide, and the area of the distribution was larger than that of the canonical nucleosome. In the H2A.B nucleosome, the free-energy distribution around the minimum (59 Å, 67 Å) was broader than the other nucleosomes, and the structures were almost evenly distributed.

From the free-energy maps, solution structures in the structural ensembles can be described, as follows: The canonical nucleosome almost adopted a compact form as shown in Figure 8A and occasionally adopted a loose form with one separated-DNA (Fig. 8B). In contrast, the CENP-A and H2A.B nucleosomes adopted a loose form. In the CENP-A nucleosome, both edges of DNA were slightly separated from the histone core (Fig. 8C). However, in the H2A.B nucleosome, the lengths of the separated DNAs were long (Fig. 8D), and both edges of the DNA fluctuated widely. These structural characteristics are not obtained solely from X-ray crystallography or SAXS, demonstrating the effectiveness of the CG-MD-SAXS method for large and flexible molecules.
SAXS profiles calculated using the CG-MD-SAXS method were almost identical to those from all-atom models including MD simulations with explicit solvent. Furthermore, for 9 structures of proteins, DNA and RNA, the theoretical SAXS profiles using the CG-MD-SAXS method were in good agreement with experimental profiles and theoretical profiles from the all-atom model. Finally, the CG-MD-SAXS method was applied to the flexible proteins, Swi5-Sfr1C, and nucleosomes. In the Swi5-Sfr1C analysis, the disorder of the β sheet improved the agreement with the experimental SAXS profile. In the nucleosome analysis, three types of nucleosomes, i.e., the canonical, CENP-A and H2A.B nucleosomes were compared, and the CG-MD-SAXS method clearly indicated the difference in the DNA conformations.

**Conclusion**

In this study, we proposed a combined method of CG MD simulations and SAXS, termed the CG-MD-SAXS method. First, the scattering factors of CG particles representing 20 amino acids, 5 bases, ribose, deoxyribose and a phosphate were obtained from a set of high-resolution crystal structures. Two important solvent effects, i.e., the excluded volume and the hydration shell, were then incorporated into the CG-MD-SAXS method. For the excluded-volume effects, the Gaussian-sphere model was employed. For the hydration shell, the hydration grids were generated around protein structures. To avoid overfitting, only two adjustable parameters were used for the entire MD trajectories. The theoretical
bound to three types of nucleosomes. Finally, the proposed method is not limited to the CG models used in CafeMol. Because the procedure of this method is general, it will be applicable to other coarse-graining methods such as the Martini model.

Acknowledgments

The authors are grateful to Prof. Mamoru Sato, Prof. Masaaki Sugiyama, Prof. Hitoshi Kurumizaka and Dr. Takashi Oda for generously providing the SAXS data for nucleosomes. This work was financially supported by the Innovative Drug Discovery Infrastructure through Functional Control of Biomolecular Systems, Priority Issue 1 in Post-K Supercomputer Development from the Ministry of Education, Culture, Sports, Science and Technology (MEXT) to M. I. (Project ID: hp150269, hp160223, hp170255, hp180191 and hp190171); by the Basis for Supporting Innovative Drug Discovery and Life Science Research (BINDS) from the Japan Agency for Medical Research and Development (AMED) (19am0101090j0003) to M. I.; a Grant-in Aid for Scientific Research on Innovative Areas “Molecular Engine” (18H05426) to M. I.; and by a RIKEN Dynamic Structural Biology Project to M. I. This research used computational resources of the K computer provided by the RIKEN Center for Computational Science. Finally, we celebrate the 80th anniversary of Prof. Nobuhiro Gō. His excellent research on biomolecules has largely inspired us to computationally study biomolecular structures. We are honored to make a contribution to this anniversary issue.

Conflicts of Interest

None.

Author Contribution

M. I. designed the research. Y. K., T. O. and M. I. developed the theory. Y. K. performed the simulations. T. E. and Y. K. analyzed the results. T. E., Y. K., T. O. and M. I. wrote the manuscript.

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