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AquaSAXS: a web server for computation and fitting of SAXS profiles with non-uniformly hydrated atomic models

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ABSTRACT

Small Angle X-ray Scattering (SAXS) techniques are becoming more and more useful for structural biologists and biochemists, thanks to better access to dedicated synchrotron beamlines, better detectors and the relative easiness of sample preparation. The ability to compute the theoretical SAXS profile of a given structural model, and to compare this profile with the measured scattering intensity, yields crucial structural informations about the macromolecule under study and/or its complexes in solution. An important contribution to the profile, besides the macromolecule itself and its solvent-excluded volume, is the excess density due to the hydration layer. AquaSAXS takes advantage of recently developed methods, such as AquaSol, that give the equilibrium solvent density map around macromolecules, to compute an accurate SAXS/WAXS profile of a given structure and to compare it to the experimental one. Here, we describe the interface architecture and capabilities of the AquaSAXS web server (http://lorentz.dynstr.pasteur.fr/aquasaxs.php).

INTRODUCTION

Small Angle X-ray Scattering (SAXS) is a technique that allows the study of the structure and interactions of biological molecules in solution. It can be used to probe proteins, nucleic acids, and their complexes under a variety of conditions, from near physiological to highly denaturing, without the need to crystallize the sample and without the molecular weight limitations inherent in other methods such as NMR spectroscopy.

The increasing availability of high-flux, third-generation synchrotron sources, improvements in detector technology and algorithmic developments for data analysis have made SAXS a technique of choice for a range of biological applications (1).

The basic principle of SAXS is to scatter X-ray photons elastically off molecules in solution, and to record the scattering intensity as a function of the scattering angle. The intensity profile of the buffer is subtracted from the profile of the macromolecule in the buffer, yielding an excess intensity profile, related to the excess electronic density of the molecule and its environment.

The SAXS profile provides information about the global structure and conformation of the studied molecule(s). Several reviews on the physical principles and theory of SAXS describe in detail how the scattering data can be analyzed and how different parameters can be fit and interpreted (2–6). Recent developments and novel applications of SAXS are described in (7–9).

Existing computational approaches for modeling a macromolecular structure based on its SAXS profile can be separated into two classes: profile-to-model (ab initio methods) and model-to-profile approaches. The former aims at proposing coarse shapes represented by dummy beads that fit the experimental profile (10–16), while the latter aims at comparing the theoretical profile of a given atomic or coarse grained model to the experimental one (17,18).

The model-to-profile approach consists in computing the theoretical profile of a given atomic structure and providing a measure of the goodness-of-fit to the experimental profile. Here, we describe a web server

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AquaSAXS (AquaSAXS) that performs this task. It is useful for many applications where one needs to decide whether the proposed model is in agreement with the experiment, and to make assumptions about why they differ, if they do.

Several tools have been designed for that purpose, following various methods (19–24). To our knowledge, the most accurate method to date for computing SAXS profiles of a given macromolecule has been achieved by treating the solvent (excluded and hydrating) explicitly (24). Through this method, even higher resolution profiles of a given macromolecule have been achieved by following various methods (19–24). To our knowledge, the most accurate method to date for computing SAXS intensity ΔI of this macromolecule at a given value of the wavevector norm q is the spherical average of the scattering intensity on the sphere of radius q in reciprocal space (Equation 1).

\[ \Delta I(q) = \int d\Omega A(q) A^*(q) \]

\[ A(q) = F_{\text{solute}}(q) + \rho_s F_{\text{sev}}(q) + \rho_s F_{\text{shs}}(q) \]

where A is the excess form factor of the system and \( F_{\text{solute}}, F_{\text{sev}} \) are respectively the form factor of the solute in vacuo and of the solvent-excluded-volume. \( F_{\text{shs}} \) is the excess form factor of the hydration shell.

We perform the spherical averaging using the cubature formulae (30). Following the effective-atomic-scattering-form-factor method (25), the solute and the solvent-excluded-volume’s form factors are computed as a sum over all N non-Hydrogen atoms of the solute (Equations 2 and 3).

\[ F_{\text{solute}}(q) = \sum_{j=1}^{N} f_j(q) e^{-q r_j} \]

\[ F_{\text{sev}}(q) = G(q, C_1) \sum_{j=1}^{N} (V_j \exp -\pi V_j^{2/3} q^2) e^{-q r_j} \]

where \( f_j \) is the atomic form factor in vacuo (19), computed as a sum of a constant and four gaussians whose parameters depend on the atom type, \( V_j \) is the solvent volume displaced by atom \( j \). \( G(q, C_1) \) is an overall expansion factor, as defined in (19), with \( C_1 \) being the ratio of the adjusted and computed average atomic radii (default value = 1.0).

Programs such as AquaSol (28) [or 3D-RISM/Amber (29)] compute solvent density maps around the solute, based on the physical interactions within the system. The method used in AquaSol is based on the Poisson–Boltzmann formalism, where the solvent is no longer described as a continuum dielectric medium but rather as an assembly of self-orienting dipoles of variable density on a grid. It was shown that the resulting water distribution is in good agreement with experimental data and with the chemical nature of the atoms exposed to the solvent, both at the atomic and residue-level (26). These maps are typically cubic grids of given size and resolution (a), where each grid point \( r \) is associated to a given density value \( \rho_{\text{solute}}(r) \). Basically, in such maps, one expects a density of 0 inside the solute, and 1 (in units of bulk density \( \rho_s \)) in the bulk region of the solvent, i.e. far from the solute. At the boundary between the solute and bulk region, the density is determined by the physico-chemical nature of the environment.

We compute the form factor for the hydration shell as in Equation 4.

\[ F_{\text{shs}}(q) = C_2 \sum_{j} \rho_{\text{solute}}(r_j) e^{-|q r_j|} \]

The sum runs over all points with nonzero density. In practice, to reduce computation time, grid points with
a density close to 1 (i.e. typically within \( \pm 1 \times 10^{-4} \)) are removed from the sum. On Urate Oxidase (example mentioned below), allowing a tolerance of \( 1 \times 10^{-6} \) slowed down the computation by a factor of three and did not affect the resulting profile: the same fitting parameters were found, and the goodness-of-fit \( \chi \) (cf Equation 6) was similar (1.688 versus 1.691).

Besides the solute and solvent, another possible contributor to the SAXS profile is the ion atmosphere surrounding the solute. AquaSol (28) computes the density maps of free cations and anions, and, in principle, these maps could be used to compute the excess form factors of ions. However, at physiological concentrations (200 mM NaCl) the ratio of the fugacities of ions and water is \(< 0.5\%\). At this stage, the contribution of ions was not implemented into AquaSAXS, except in the form of explicitly bound and fixed ions. Nevertheless, the presence of free ions can indirectly affect the solvent density in the hydration shell (screening effect), so the user is prompted for the ionic strength of the solution.

The computed profile \( \Delta I \) is fitted to a given experimental SAXS profile \( \Delta I_{\text{exp}} \) (with experimental error \( \sigma \)) by minimizing the goodness-of-fit function \( \chi \) with respect to three adjustable parameters: \( C_1 \), \( C_2 \) and \( C \) (Equation 6).

\[
\chi = \left( \frac{1}{N_q} \sum_{q_i} \left( \frac{\Delta I_{\text{exp}}(q_i) - C \Delta I_{\text{fits}}(C_1, C_2)}{\sigma(q_i)} \right)^2 \right)^{\frac{1}{2}}
\]

where \( s_j \) is the fraction of solvent accessible surface of the atom \( j \) (31) and \( f_w \) is the water form factor. \( C_2 \) is a scale factor used to adjust the hydration shell’s contribution (default value = 1.0).

The solvation method, and whether a comparison is made with an experimental SAXS profile or not.

There are three options for the solvation method: either a solvation map is provided as input, or a solvation map is computed using AquaSol (28), or a hydration layer is defined using the accessible surface of each atom, following the method introduced in FoXS (20).

For the Solvent-map solvation option, for maps with 65 grid points per edge (about 2 \( \text{Å} \) grid size), the calculation typically takes from less than a minute to a few minutes for systems of a few thousands atoms to dozens of thousands of atoms (with \( N_q \approx 60 \)), which is a few times slower than CRYSTOL and FoXS. For a WAXS spectrum, \( N_q \approx 300 \). For the Surface-Accessible solvation option, computation is more efficient.

**AQUASAXS WEB SERVER**

Figure 1 shows the flowchart of a typical AquaSAXS calculation. Starting with a PDB file or a PQR file, the user is expected to make two decisions, namely the selection of the solvation method, and whether a comparison is made with an experimental SAXS profile or not.

AquaSAXS was successfully tested with all PDB (32) structures that have an experimental SAXS profile in the open access BioIsis database (33), and gave results similar to CRYSTOL and FoXS. It was also tested on Urate Oxidase (see Figure 2).

The calculation scales linearly with the number of points at which the profile is sampled \( (N_q) \), as well as with the number of non-Hydrogen atoms of the solute. It scales to the cube of the number of points per grid edge, although this expensive cost is attenuated by a preliminary compressing process discarding all points with an excess density close to zero.
The structure file in PDB or PQR format is mandatory. Each line of the file starting with the label ‘ATOM’ or ‘HETATM’ is stored in memory. If the corresponding atom is a hydrogen or belongs to a water molecule, it is discarded. Otherwise, the residue and atom’s names are parsed among the standard PDB protein, nucleic acids and ligands library (32). Thirteen atom types are currently recognized: Carbons with zero, one, two or three bound hydrogens, nitrogen with zero, one, two or three bound hydrogens, oxygen with zero or one bound hydrogen, sulfur with zero or one bound hydrogen, and phosphorus. Once the atomic type of atom \( j \) has been recognized, its position \( r_j \) is stored and it is assigned the corresponding form factor \( f_j \), excluded volume \( V_j \) and radius \( r_j \). On the ‘Flowchart’ web-page, the user is given the possibility to check whether the atomic types of the residues in the provided PDB/PQR file can be recognized. If the user wants to define other atomic types than those listed above, two optional files can be given as input to the program: one listing the atom types of a given residue, the other listing the atomic parameters of new atom types.

Several other options/parameters can be set: the maximum \( q \)-value considered, the sampling resolution of the profile, the bulk average electron density (in eÅ\(^{-3}\)), the subset chains in the structure to be considered, as well as the values of \( C_1 \) and \( C_2 \) in the nonfitting mode.

The computation is performed in real time and the browser is redirected to the result’s page when the calculation has finished. If an email address is provided, an email will be sent to the user. Depending on the system’s size and server’s queue load, the typical running time ranges from less than a minute to a few minutes. The result’s page displays a plot of the computed profile (see Figure 2), superimposed to the experimental profile, if provided, as well as the run logfile. Links to three output files are displayed, to retrieve the logfile, the profile file, and a PDB file listing all the atoms that have been considered in the calculation. Possibly, links toward the output files of AquaSol (28) are displayed too (computed solvent map and logfile).

### ADDITIONAL FEATURES

In addition to single conformation fitting of experimental SAXS profiles, AquaSAXS provides two methods to deal with multiple structure files.

The first one, called ‘Sequential fit’, provides a way to compute the SAXS profile of a set of PDB/PQR files in a single run. That way, if an experimental profile is provided, the user can readily compare the relevance of provided models, without having to rerun the server several times.

The second method, called ‘Ensemble fit’, aims at plugging in all possible models and refining their population, and is directly inspired from (35). This method can prove useful when the macromolecule visits several conformational states in solution. Another interesting way to take profit of this approach would be the following: starting from a given PDB structure, several models are built along a given deformation parameter (e.g. a normal mode); the best value for this deformation parameter would likely be detected by the refinement process.

In practice, this method couples a mean-field optimization of the model’s populations with a simulated annealing protocol. Let us consider an ensemble of \( M \) models, each model \( m \) being associated the scattered intensity \( I_m \). Noting \( p_m \) the associated probability (or population) of model \( m \), the total scattered intensity (in the limit of infinite dilution) is then:

\[
\Delta I = \sum_{m=1}^{M} p_m \Delta I_m
\] (7)
If one defines a free energy $F$ of the form:

$$F = \chi - TS$$

$$S = -\sum_{m=1}^{M} p_m \log p_m$$  \hspace{1cm} (8)

where $\chi$ is defined as in Equation 6, $S$ is the entropy, and $T$ is the temperature, then minimizing the free energy with respect to $p_m$ gives:

$$p_m = (1/Z) \exp(-\beta E_m)$$  \hspace{1cm} (9)

where $\beta$ is the inverse of $T$, $E_m = \delta \chi / \delta p_m$ and $Z$ is a normalization constant given by $1 = \sum_m p_m$.

The refinement starts with uniform values of the probabilities, which are updated at each cycle of the refinement until a self-consistent solution is obtained; at each cycle the derivative are evaluated at the current solution, i.e. the current set of $p_m$ values. The temperature governs the contrast between the different populations, the contrast being higher as the temperature is lower. After convergence at a given temperature, the set of populations found is used as initial guess for a new mean-field refinement at lower temperature, until the final temperature is reached.

Results of both methods applied on synthetic data are presented in Supplementary Materials.

**CONCLUSION**

We have described a program that allows structural biologists to compare their SAXS data to the theoretical one for a model given as a PDB or PQR file. Its major novelty resides in the possibility to better model the hydration layer through a physically sound representation of the solvent density map, combined with the use of the cubature method for spherical averaging. The user-friendly interface allows to modify (or add new entries to) the list of scatterers and their parameters.

The possibility to fit the data with multiple models, either independently, or through population refinement has also been implemented.

Future developments will allow for the possibility to refine the coordinates of the model against the experimental data. In that case, care must be taken to use as few degrees of freedom as possible. One possibility is to restrict the deformation of the model along a small number of 'essential' normal modes within the framework of the Elastic Network Model (36).

Finally, the possibility to compute the theoretical anomalous SAXS profile of a solute containing atoms with anomalous contribution (e.g. Bromide or Cesium ions) will be made available soon.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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