More precisely, the planned developments include the following:

Three main axes of the proposed project are:
(i) Molecular flexibility and the ways of modelling it;
(ii) Analysis of polynomial translation theorems for integrative structural biology; (iii) Optimization will be performed using the algorithmic foundation to the upcoming single-molecule experimental techniques. In particular, the methods for small-angle (both SAXS and SANS) scattering experiments and to provide the reconstruction of model shapes. Many efforts have been spent to reduce the running time of these tools without degrading the quality of their approximations, most prominently the ATSAS package developed at EMBL Hamburg. Particularly, the Crysol program calculates a model SAXS profile to test new experimental data. The dependence of the scattering of the solvent to match the scattering of the model is a challenging issue that has been overcome by using the Pepsi-SAXS / Pepsi-SANS engines with the NOLB NMA method for the large-scale flexible structure optimization along the lowest-frequency normal modes.

The overall research topic of the current project is to extend the state-of-the-art computational tools without degrading the quality of their approximations, most prominently the ATSAS package developed at EMBL Hamburg. Particularly, the Crysol program calculates a model SAXS profile to test new experimental data. The dependence of the scattering of the solvent to match the scattering of the model is a challenging issue that has been overcome by using the Pepsi-SAXS / Pepsi-SANS engines with the NOLB NMA method for the large-scale flexible structure optimization along the lowest-frequency normal modes.

Assignment

Introduction:

While crystallography has been providing atomic-resolution structures of biomolecules for over half a century, the real challenge today is to correlate molecules’ structure and dynamics in solution with their function. Small-angle scattering (SAXS) is the fundamental techniques for structural studies of biological systems in solution. Thanks to advances in instrumentation and data analysis software, Small-angle X-ray scattering (SAXS), complemented by other methods, is becoming very popular in structural biology. Over the years, a number of computational tools have been developed for the analysis of SAXS curves, calculation of theoretical profiles and low-resolution reconstruction of model shapes. Many efforts have been spent to reduce the running time of these tools without degrading the quality of their approximations, most prominently the ATSAS package developed at EMBL Hamburg. Particularly, the Crysol program calculates a model SAXS profile to test a structural hypothesis of a SAXS experiment. The number of Bio-SAXS publications exploded as a result of this effort. Comparatively, the lack of user-friendly analysis tools has hindered the development of Small Angle Neutron Scattering (SANS), more complex but providing more information.

Very recently, we developed a computational tool called Pepsi-SAXS that outperforms all the competitors for atomistic modelling in both speed (5 – 50 times faster on average) and accuracy. While crystallography has been providing atomic-resolution structures of biomolecules for over half a century, the real challenge today is to correlate molecules’ structure and dynamics in solution with their function. Small-angle scattering (SAXS) is the fundamental techniques for structural studies of biological systems in solution. Thanks to advances in instrumentation and data analysis software, Small-angle X-ray scattering (SAXS), complemented by other methods, is becoming very popular in structural biology. Over the years, a number of computational tools have been developed for the analysis of SAXS curves, calculation of theoretical profiles and low-resolution reconstruction of model shapes. Many efforts have been spent to reduce the running time of these tools without degrading the quality of their approximations, most prominently the ATSAS package developed at EMBL Hamburg. Particularly, the Crysol program calculates a model SAXS profile to test a structural hypothesis of a SAXS experiment. The number of Bio-SAXS publications exploded as a result of this effort. Comparatively, the lack of user-friendly analysis tools has hindered the development of Small Angle Neutron Scattering (SANS), more complex but providing more information.

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Challenges and tasks:

The overall research topic of the current project is to extend the state-of-the-art computational methods for small-angle (both SAXS and SANS) scattering experiments and to provide the algorithmic foundation to the upcoming single-molecule experimental techniques. In particular, the practical aim of the project is the development of software tools with intuitive user-interaction feedback.

Mathematically, the project will rely on the very efficient representation of the scattering profiles based on polynomial expansions of the scattering amplitudes using spherical harmonics. Structural optimization will be performed using the fast Fourier transform–accelerated techniques and the polynomial translation theorems and large collective structural motions using Normal Mode Analysis.

Three main axes of the proposed project are:

(i) Molecular flexibility and the ways of modelling it;
(ii) Algorithmic developments for new experimental setups, particularly combining SAXS and SANS data, and 2D scattering;
(iii) Practical software development with intuitive user interfaces. Below we detail the main research axes of the project.

More precisely, the planned developments include the following:

- Modelling thermal fluctuations (see review by Hub, 2018) using an analytical model of the scattering profile resulting from thermal molecular vibrations. This will be possible thanks to the (i) expression of normal modes in spherical coordinates expanded in the Spherical Harmonics basis, and (ii) computing molecular fluctuations in this basis analytically using the algebra for triple and quadruple overlap integrals of spherical harmonics.
- Flexible fitting of atomistic structures into small-angle scattering profiles. This includes combination the Pepsi-SAXS / Pepsi-SANS engines with the NOLB NMA method for the large-scale flexible structure optimization along the lowest-frequency normal modes.
- Numerical schemes and simulations
- Scientific computing (BAP E)
- Town/City: Grenoble
- Inria Center: CRI Grenoble - Rhône-Alpes
- Starting date: 2018-10-01
- Duration of contract: 3 years
- Deadline to apply: 2018-08-15

Contacts

- Inria Team: NANO-D
- Recruiter: Grudinin Sergey / sergei.grudinin@inria.fr

About Inria

Inria, the French National Institute for computer science and applied mathematics, promotes “scientific excellence for technology transfer and society”. Graduates from the world’s top universities, Inria’s 2,700 employees rise to the challenges of digital sciences. With its open, agile model, Inria is able to explore original approaches with its partners in industry and academia and provide an efficient response to the multidisciplinary and application challenges of the digital transformation. Inria is the source of many innovations that add value and create jobs.

The keys to success

We are looking for a motivated person willing to advance the state of the art in the field of integrative structural biology. The essential skills of the candidate include algorithm development and validation, C++ coding and proficiency in computational physics and math.

Conditions for application

Defence Security:

This position is likely to be situated in a restricted area (ZRR), as defined in Decree No. 2011-1425 relating to the protection of national scientific and technical potential (PPST). Authorisation to enter an area is granted by the director of the unit, following a favourable Ministerial decision, as defined in the decree of 3 July 2012 relating to the PPST. An unfavourable Ministerial decision in respect of a position situated in a ZRR would result in the cancellation of the appointment.

Recruitment Policy:

As part of its diversity policy, all Inria positions are accessible to people with disabilities.

Warning: you must enter your e-mail address in order to save your application to Inria. Applications must be submitted online on the Inria website. Processing of applications sent from other channels is not guaranteed.
working prototype exists, however, a proper analytical optimization technique with respect to the goodness of fit should be added. Also, the method needs to be extended to SANS data. Finally, the method needs to be made more robust and user-friendly, possibly, with a GUI inside the SAMSON modelling platform.

- Application of motion planning to adapt the sampling techniques for flexible unstructured regions of the molecules.
- In some cases it is impossible to represent the molecular structural heterogeneity continuously. This situation will require fitting the experimental SAS curves with an ensemble of structures. Therefore, we plan to extend the Pepsi-SAXS / Pepsi-SANS methods for the cases of multiple models, whose contributions will be adjusted with stochastic optimization techniques.
- We will consider the case when the molecular system can be represented with a set of rigid domains. Here we can compute scattering profiles for supra-molecular assemblies based on the precomputed scattering amplitudes of individual rigid bodies with subsequent rigid-body transforms applied to the amplitudes in the Fourier space.

- Fitting of atom-resolved structures into multiple profiles obtained in different experimental configurations from the same sample. Indeed, the current modelling pipelines merge different experimental measurements into a single scattering profile. This approach has a significant drawback since the scaling factors for individual scattering components are generally unknown and can be erroneously guessed upon merging. A much more general, and also more computationally elegant approach would be to adjust the scaling factors of individual scattering profiles upon fitting atomistic models into multiple scattering curves. This approach would also have an obvious advantage of mixing different experimental information, e.g. SAXS and SANS experiments.
- Novel developments for 2D scattering based on the spherical harmonics and cylindrical harmonics representation. Many elongated or polymeric molecules can be aligned in space along the magnetic field. Thanks to this spatial alignment, their scattering profiles wouldn’t be axially symmetric any longer and will contain much more information compared to profiles from spherically averaged particles. We plan to extend the current method for analytical cylindrical averaging of spherical harmonics with multiple advantages over the standard approaches.

References:

Collaboration:
The recruited person will be in connection with

1. computational expert Sergei Grudinin from Inria / CNRS Grenoble
2. bioSAXS experts Anne Martel and Sylvain Prevost from ILL Grenoble
3. 2D fiber diffraction experts Trevor Forsyth, and Estelle Mossou from ILL Grenoble
4. an instrument scientist at BioSAXS Martha Brennich from EMBL Grenoble

Main activities
This is a very challenging project whose aim is to change the modelling paradigm in the small-angle community. It has many ambitious goals both on the algorithmic side and also considering software applications. The main activities of the candidate will be algorithms and code developments including the following:

- Combination of the existing software tools that model molecular flexibility (deformations along the slowest normal modes, rigid-body perturbations of rigid domains, modeller of flexible loops using robotics techniques) into a unified library and link it with the Pepsi-SAXS library. Then, multiple tests should be performed, and probably novel benchmarks should be created.
- Adaptation of the developed methods for SAXS scattering and flexible fitting of molecular models into scattering profiles for SANS experiments. Computationallly, it is an easy and straightforward task, however, numerous tests should be performed and very convenient software released.
- Formulation of the local refinement problem in the basis of Spherical Harmonics (SH). This is a challenging task, since no scattering methods that use analytical gradients in the SH basis have been developed.
- Developments for scattering of aligned objects and single-particle experiments
- Creation of practical interactive GUI interfaces with force feedback computed from the experimental scattering profiles. This is a challenging task, however, we have already done attempts for the integration of the NOUB NMA and Pepsi-SAXS methods inside the SAMSON modular simulation platform (see https://www.samson-connect.net). Again, the most difficult part will be to create convenient and interactive software tools.

Skills
Technical skills and level required: the candidate must be proficient in computer science, computational physics and applied math. Excellent knowledge of C++ and linear algebra are strictly required.

Languages: excellent oral, written and interpersonal communication skills are essential (working language will be English – knowledge of French is a plus).

Relational skills: the candidate will have to communicate with the software developers at the side of the Inria partner, instrument scientists at the side of EMBL-ILL partners, and also with numerous users of the small-angle instruments at ILL and ESRF facilities.

Other valued appreciated: we are looking for creative, passionate and hard-working individuals with exceptional talent for computer science and mathematics.

Benefits package
Subsidised catering service
Partially-reimbursed public transport
Social security
Paid leave
Flexible working hours
Sports facilities

Remuneration
Salary: 1768.55 € gross/month paid by university