Characterization of conformational deformation-coupled interaction between immunoglobulin G1 Fc glycoprotein and a low-affinity Fcγ receptor by deuteration-assisted small-angle neutron scattering

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\textbf{A B S T R A C T}

A recently developed integrative approach combining varied types of experimental data has been successfully applied to three-dimensional modelling of larger biomacromolecular complexes. Deuteration-assisted small-angle neutron scattering (SANS) plays a unique role in this approach by making it possible to observe selected components in the complex. It enables integrative modelling of biomolecular complexes based on building-block structures typically provided by X-ray crystallography. In this integrative approach, it is important to be aware of the flexible properties of the individual building blocks. Here we examine the ability of SANS to detect a subtle conformational change of a multidomain protein using the Fc portion of human immunoglobulin G (IgG) interacting with a soluble form of the low-affinity Fcγ receptor IIIb (sFcγRIIIb) as a model system. The IgG-Fc glycoprotein was subjected to SANS in the absence and presence of 75%-deuterated sFcγRIIIb, which was matched out in D\textsubscript{2}O solution. This inverse contrast-matching technique enabled selective observation of SANS from IgG-Fc, thereby detecting its subtle structural deformation induced by the receptor binding. The SANS data were successfully interpreted by considering previously reported crystallographic data and an equilibrium between free and sFcγRIIIb-bound forms. Our SANS data thus demonstrate the applicability of SANS in the integrative approach dealing with biomacromolecular complexes composed of weakly associated building blocks with conformational plasticity.

\textbf{1. Introduction}

Recent progress in X-ray crystallography has yielded more than hundred thousand protein structures deposited in the Protein Data Bank. As such, current trends in structural biology are increasingly focusing on more complicated macromolecular complexes, which remain difficult to analyze using traditional approaches. A recently developed approach for dealing with such complex and often huge systems is integrating the atomic coordinates of individual building blocks determined by X-ray crystallography and/or NMR spectroscopy into a low-resolution three-dimensional (3D) map as a template provided typically by cryo-electron microscopy [1]. This integrative approach has successfully resulted in 3D structure models of biomolecular assemblies such as HIV capsid assemblies and the DNA telomerase complex [2,3]. Solution scattering also provides information on gross shapes of biomacromolecular complexes complemented by crystal structures. In particular, small-angle neutron scattering (SANS) enables selective observation of specific components in the complexes by selective

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deuteration combined with contrast-matching methods. This is because the neutron scattering length is different between hydrogen and deuterium. This technique was pioneeringly applied to determine the spatial arrangements of 30 S ribosomal subunits [4] and recently sophisticated as exemplified by the application to quaternary structure analyses of a variety of protein complexes [5–8]. Deuteration-assisted SANS data in conjunction with crystallographic data has thus facilitated 3D modelling of quaternary structure views of protein complexes like the circadian clock protein complex, KaiB and KaiC [9], and the large dodecameric aminopeptidase TET [10].

In this integrative approach, it is important to correctly assess the conformational alteration of each component upon complex formation, when the overall complex structures are fabricated with the individual building-block structures. It is generally difficult to integrate flexible parts of the building blocks, whose electron densities are often missing in the crystal structure [6,11–13]. This is particularly problematic if the subunits are modified with flexible oligosaccharide moieties. Furthermore, if some building blocks were assembled through weak interactions, dynamic equilibrium between complexed and uncomplexed states should be considered to accurately interpret the data.

In view of this situation, we examined the capability of SANS to deal with protein complexes involving glycoproteins that undergo conformational alterations upon their weak association. As a model system, we used an interaction between the Fc region of human IgG1 (IgG1-Fc) and a low-affinity Fcγ receptor (FcγRIIIb, FcγRIIIb). IgG1-Fc consists of two identical 26 kDa-polypeptide chains each divided into Cγ2 and Cγ3 domains. Each Cγ2 domain possesses a conserved glycosylation site at Asn297 displaying a complex-type oligosaccharide (with a molecular mass of 21.4 kDa), indicating a conformational change of Fc upon formation of their 1:1 complex (Fig. 1). This allows us to assess the capability of the deuteration-assisted SANS method to detect conformational alterations of the multidomain glycoprotein in solution upon interacting with its low-affinity binding partner.

2. Materials and methods

2.1. Protein preparation

The Fc fragments were prepared by papain digestion from human IgG1. The extracellular regions of human FcγRIIIb was expressed using Escherichia coli strain BL21(DE3) as inclusion body. For preparation of deuterated proteins, the bacterial cells were grown in an M9 minimal medium containing 2 g/l of glucose as a 1–3 mixture of isotopically natural and fully deuterated glucose (1,2,3,4,5,6-D6-, 98%, Cambridge Isotope Laboratories Inc.), along with a 25:75 ratio of H2O and D2O as previously described [6]. The inclusion body was suspended in 50 mM Tris-HCl, pH 8.0, containing 150 mM NaCl. After sonication, the homogenate was centrifuged at 8000 g for 15 min at 4 °C and the precipitate was denatured by 8 M urea. The denatured protein solution was diluted with refolding buffer (50 mM Tris-HCl, pH 8.0, 0.5 mM arginine, 5 mM CaCl2, 5 mM reduced form of glutathione, and 0.5 mM oxidized form of glutathione). After two days of incubation at 4 °C, the diluted protein solution was applied to aComplete His-Tag Purification Resin (Roche) and further purified on the HiLoad 26/60 Superdex 75 pg column (GE Healthcare).

2.2. Solution scattering measurements

SANS experiments were performed using the D22 instrument installed at the Institut Laue-Langevin (ILL), Grenoble, France (Table S1). The SANS intensities were observed with 6.0 Å neutrons and two sample-to-detector distances of 5.6 m and 1.4 m: the covered q-ranges are 0.01–0.55 Å−1. The temperature was maintained at 25 °C in the irradiation. The observed SANS intensity was corrected for background, empty cell and buffer scatterings, and transmission factors and subsequently converted to the absolute scale by GRASP software using incident beam flux (http://www.ill.eu/instruments-support/instruments-groups/groups/lss/grasp/home/).

For all SANS measurements, we used 99.8% D2O (ISOtec) containing 50 mM Tris-HCl, pH 8.0, and 150 mM NaCl as the solvent to reduce the incoherent scattering and obtain high statistical data in the higher q-range. We measured the SANS of non-deuterated Fc (14 μM, 0.73 mg/ml) and 75%-deuterated sFcγRIIIb (31 μM, 0.70 mg/ml) as well as their 1:1 mixture at each concentration of 16 μM (0.83 mg/ml of Fc plus 0.36 mg/ml of sFcγRIIIb). As reference, 75%-deuterated sFcγRIIIb (31 μM) under identical solution conditions was also subjected to small-angle X-ray scattering (SAXS) measurement with BL15A2 at Photon Factory, KEK, Tsukuba, Japan.

SANS profiles were calculated from atomic coordinates deposited in the Protein Data Bank (PDB) with Debye functions, taking into account the deuteration ratios of both solvent-exchangeable and nonexchangeable hydrogens, as well as the excluding volume of the protein(s) (Table S2) (for more detail, refer to Supplementary Method in Supplementary Materials).

3. Results and discussions

In this study, we performed SANS characterization of human IgG1-Fc in solution in both the presence and absence of sFcγRIIIb. Firstly, we observed a SANS profile of human IgG-Fc alone, which was compared with theoretical profiles computed from its crystal structure (Fig. 2). The observed profile was well reproduced by the simulated curve only when the carbohydrate moieties attached to Fc were considered in the calculation.

Because the total contrast and scattering intensity of the carbohydrate part are much smaller than those of the polypeptide part in IgG1-Fc (Fig. S1 (a) and (b)), it is expected that the glycans contribute only very slightly to the scattering profiles. However, our simulation based on the crystal structure indicates that the glycans specifically affected the scattering profile in the q-range of 0.12–0.20 Å−1, where the interference term between the glycans and the polypeptides in Fc was specifically negative (Fig. S1 (c)). This q-dependent effect can be attributed to the unique horseshoe-shaped structure of Fc accommodating the glycans in its cavity (Fig. 1). The experimentally obtained SANS profile can be thus interpreted, confirming that the pair of biantennary complex-type oligosaccharides are packed between the two Cγ2 domains as observed in the crystal structure.

Next, we observed SANS profiles of IgG1-Fc in complex with...
For selective observation of SANS originating from the Fc glycoprotein, we prepared bacterially expressed sFcγRIIIb with 75% deuteration. This sFcγRIIIb preparation was subjected to SANS as well as SAXS in 99.8% D2O solution (Fig. S2). The results confirmed that the SANS of 75%-deuterated sFcγRIIIb was matched out with the D2O solvent. The SANS profile of IgG1-Fc in complex with 75%-deuterated sFcγRIIIb was significantly different from that observed for Fc-sFcγRIIIb complex without deuteration, demonstrating the successful elimination of SANS contribution of sFcγRIIIb (Fig. 3).

The inversely contrast-matched profile, which originated selectively from the Fc part in the complex, exhibited marked deviation from the profile of IgG1-Fc alone in the high-angle region, although their Rg values calculated from Guinier plots were almost identical, i.e. 26.3 ± 0.4 Å and 26.5 ± 0.3 Å for Fc in the presence and absence of 75%-deuterated sFcγRIIIb, respectively (Fig. S3 and Table S3). This indicates that Fc undergoes subtle conformational rearrangement upon binding to sFcγRIIIb in solution, presumably as indicated by the crystallographic data (Fig. 1). Because the scattering curve in the q-range of 0.12–0.2 Å⁻¹ sensitively reflects the structural perturbations on the horseshoe-shape of IgG1-Fc (vide supra), the quaternary structural change induced by sFcγRIIIb binding is expected to affect SANS profile in this range. Hence, we compared the observed SANS profile of Fc complexed with 75%-deuterated sFcγRIIIb with the profile simulated from the Fc part of the previously reported crystal structure of the Fc-

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**Fig. 2.** (a) SANS profile for Fc alone (open circles) shown with theoretical profiles computed from the crystal structure of human IgG1-Fc in an uncomplexed state with (green) and without (orange) the glycan parts. (b) The enlarged view of (a).

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**Fig. 3.** SANS profiles for Fc in the presence of non-deuterated sFcγRIIIb (gray circles) and Fc in the presence of 75%-deuterated sFcγRIIIb (black circles) in 99.8% D2O. Graphical models illustrate deuteration (colored yellow) of solvent and protein based on the crystal structure of the Fc-sFcγRIIIb complex (PDB code 1T89).

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**Fig. 4.** (a) SANS profile for Fc in complex with 75%-deuterated sFcγRIIIb (closed circles) shown with theoretical profiles computed from the crystal structures of human IgG1-Fc in an uncomplexed state green), and in a sFcγRIIIb-bound state (blue). The theoretical profile of 72% sFcγRIIIb-bound form of Fc was reconstructed from the crystal structures of human IgG1-Fc in the free and the sFcγRIIIb-bound forms (red). (b) The enlarged view of (a).
sFcRIIib complex [14,15]. The simulated curve is closer to the experimentally observed profile than that of free Fc, but still exhibits significant deviation from the contrast-matched SANS profile of the complex (Fig. 4).

Because the interaction between human IgG1-Fc and sFcRIIib is characterized by their low affinity, the remaining discrepancy might be attributed to partial dissociation of the complex. Taking into account the possible partial dissociation of the complex, the residual sum of squares (RSS) between the experimental and simulated data was calculated for varying ratios of sFcRIIib-bound Fc (Fig. S4). The RSS plotted as a function of the bound Fc ratio exhibited a minimum of 72% sFcRIIib-bound form of Fc. Indeed, the simulated curve computed taking into account this equilibrium was in excellent agreement with the experimentally observed SANS profile (Fig. 4). Because the dissociation constant of the Fc-sFcRIIib interaction has been reported to be 1.4 × 10^{-6} M [16], the mixture of Fc and sFcRIIib subjected to the SANS measurement (at each concentration of 16 μM) is supposed to be under equilibrium between 26% free and 74% sFcRIIib-bound form. Thus, the partial dissociation of the low-affinity protein complex could be successfully treated in this approach.

The present study demonstrated the utility of the SANS approach with inverse contrast-matching technique for structural characterization of a protein with glycosylation that undergoes conformational rearrangement upon formation of a weakly associated biomolecular complex. This is because the use of approximately 100% D2O as solvent for inverse matching enables us to obtain high S/N data even in the higher q-range. Thus, the deuteration-assisted SANS method plays a key role in the integrative structural biology approach to deal with complicated biomacromolecular complexes composed of weakly associated building blocks with conformational plasticity.

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Appendix A. Transparency document

Supplementary data associated with this article can be found in the online version at http://dx.doi.org/10.1016/j.bbrep.2017.08.004.

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