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ARTICLE





Structural insights into Acyl-coenzyme A binding domain containing 3 (ACBD3) protein hijacking by picornaviruses

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1 | INTRODUCTION

Positive-sense single-stranded ribonucleic acid (+ RNA) viruses replicate at the surface of intracellular membranes where they establish their so-called viral replication organelles (ROs), a membranous platform for viral replication with complex membrane topology.¹ ROs provide shelter from innate immunity and also serve as a rendezvous point for viral and host proteins and for the assembly of new virions. A key step in the biogenesis of ROs is the recruitment of a phosphatidylinositol 4-kinase (PI4K).² Enteroviruses and kobuviruses, both members of the Picornaviridae family, use their small nonstructural 3A protein to recruit the phosphatidylinositol 4-kinase B (PI4KB) kinase. 3A is a small (<100 AAs) protein with a predicted C-terminal transmembrane helix.³ Its cytoplasmic part tightly interacts with the Acyl-coenzyme A binding domain containing 3 (ACBD3) protein.4,5

ACBD3 protein is a Golgi-resident multifunctional, multidomain protein involved in maintenance of the Golgi

Many picornaviruses hijack the Golgi resident Acyl-coenzyme A binding domain containing 3 (ACBD3) protein in order to recruit the phosphatidylinositol 4-kinase B (PI4KB) to viral replication organelles (ROs). PI4KB, once recruited and activated by ACBD3 protein, produces the lipid phosphatidylinositol 4-phosphate (PI4P), which is a key step in the biogenesis of viral ROs. To do so, picornaviruses use their small nonstructural protein 3A that binds the Golgi dynamics domain of the ACBD3

small nonstructural protein 3A that binds the Golgi dynamics domain of the ACBD3 protein. Here, we present the analysis of the highly flexible ACBD3 proteins and the viral 3A protein in solution using small-angle X-ray scattering and computer simulations. Our analysis revealed that both the ACBD3 protein and the 3A:ACBD3 protein complex have an extended and flexible conformation in solution.

KEYWORDS

Abstract

ACBD3, coarse-grained simulations, host factor, intrinsically disordered regions, picornavirus, RNA virus, small-angle X-ray scattering (SAXS)

apparatus.⁶ One of its key functions is the recruitment of the lipid kinase PI4KB,⁷ which together with the PI4K2A enzyme is responsible for the synthesis of the lipid hallmark of the Golgi—the phosphatidylinositol 4-phosphate (PI4P) lipid.^{8,9} ACBD3 consists of several domains connected by flexible linkers (Figure 1). Besides the N-terminal ACBD domain, the protein also contains a Q (glutamine-rich) domain that forms a helical bundle, which binds the most proximal N-terminal helix of PI4KB with submicromolar affinity.⁷ The Golgi dynamics (GOLD) domain is localized at the C terminus and serves as a protein-binding site. It is worth noting that most ACBD3 interacting proteins (giantin, TSPO, Rhes, PKARIα, and Golgin-160) have been reported to bind to the GOLD domain.^{6,10}

Recently, insights into interactions of the 3A protein from several picornaviruses, including the Aichi virus (genus *Kobuvirus*) and four representative enteroviruses (poliovirus, enterovirus A71, enterovirus D68, and rhinovirus B14), with the GOLD domain was gained crystallographically.^{11,12} This structural analysis revealed that the viral 3A acts as a molecular harness—it wraps around the



FIGURE 1 Schematic representation of the ACBD3 and Aichi 3A proteins with the binding sites for 3A and ACBD3 proteins highlighted

GOLD domain of ACBD3 and pins it down to the target membrane, which in turn leads to recruitment of the PI4KB lipid kinase and to the production of the PI4P. This lipid can be exchanged for other lipids such as cholesterol in order to build membranes of required properties for viral replication¹³ or to directly recruit viral RNA polymerase 3D^{pol}.^{14,15}

Crystallographic analysis of the full-length ACBD3 and 3A was not possible because the ACBD3 protein contains large intrinsically disordered regions that connect the well-defined domains. However, small-angle X-ray scattering (SAXS) combined with computer simulations is well suited for the analysis of large flexible proteins or protein complexes.^{16,17} We used a combination of SAXS experiments and coarse-grained molecular simulations to analyze the ACBD3 protein and its complex with the Aichi virus 3A protein.

2 | MATERIALS AND METHODS

2.1 | Protein expression and purification

Expression and purification of the full-length ACBD3 and 3A protein from the human Aichi virus were described earlier.¹¹ Briefly, the sequence encoding the full-length ACBD3 protein was subcloned into the pRSFD plasmid with 6xHis tag fused to GB1 solubilization tag and TEV protease cleavage site.¹⁸ For the expression of the 3A protein, the sequence encoding for Aichi virus 3A (amino acids 2-58) was also inserted to the pRSFD plasmid with 6xHis tag fused to GB1 solubilization tag and TEV protease cleavage site and coexpressed with the GOLD domain from ACBD3 (ACBD3 amino acids 364-528) from the same plasmid because interaction of these partners stabilizes the 3A protein and protects it from degradation in bacteria. Proteins were expressed in E. coli BL21 Star in autoinduction ZY medium. Cells were lysed by sonication in lysis buffer (50 mM Tris, pH 8.0, 300 mM NaCl, 20 mM imidazole, 3 mM β-mercaptoethanol, 10% glycerol), centrifuged, and the supernatant was used for the first step of purification by affinity chromatography (Ni-NTA, Macherey-Nagel). The 6xHis-GB1

solubilization tag was cleaved by the TEV protease, and the proteins were further purified by size exclusion chromatography using Superdex 200 HiLoad 16/60 column (GE Healthcare) in 10 mM Tris, pH 8.0, 200 mM NaCl, and 3 mM β -mercaptoethanol. For the preparation of samples for SAXS measurements, ACBD3 alone or incubated with 6× molar excess of the 3A protein was repurified on the Superdex 200 Increase 10/300 GL column (GE Healthcare) in 10 mM Tris at pH 7.4, 150 mM NaCl, and 2 mM tris(2-carboxyethyl)phosphine. Purified proteins were concentrated, aliquoted, flash frozen in liquid nitrogen and stored in -80° C until needed.

2.2 | SAXS data processing

The SAXS data were collected using BioSAXS Beamline BM29 (ESRF Grenoble) equipped with the Pilatus 1M detector (Dectris) at the ACBD3 protein concentrations of 5.8, 3.4, and 2.6 mg/mL. For each of the three concentrations, the SAXS data were frame averaged and buffer subtracted using PRIMUS.¹⁹ The resulting three data sets were then extrapolated to the dilute limit using the same software (Figure S1). The extrapolated SAXS data set was next used to model conformations of the ACBD3 protein, as detailed in the following section. In parallel, SAXS data were collected for the ACBD3:3A/AiV protein complex at concentrations of 4.7, 2.7, 1.8, and 0.9 mg/mL. These SAXS data were frame averaged, buffer subtracted, and subsequently extrapolated to the dilute limit in the same way that the ACBD3 SAXS data were processed (Figure S2). The extrapolated SAXS data set was used to model conformations of the ACBD3:3A/AiV protein complex in solution.

2.3 | Molecular modeling and simulations

To efficiently sample conformations of the full-length ACBD3 protein in solution, we used a coarse-grained simulation model introduced by Kim and Hummer (KH).²⁰ The KH model is equipped with a transferable energy function and devised to simulate conformational ensembles of large multidomain proteins and multiprotein complexes. It has been successfully applied to protein systems ranging from the endosomal sorting complexes required for transport membrane-binding trafficking machinery²¹⁻²³ to multidomain protein kinases²⁴ and kinases in dynamic complexes with phosphatases.^{25,26} Most recently, the KH model has been also used to study multidomain carbohydrate-active enzymes,^{27,28} flexible protein complexes containing lipid kinases,^{29,30} and complexes of cell adhesion proteins involved in immunological responses.³¹ In the framework of the KM model, the solvent is implicit and amino acid residues are represented as single beads centered at α -C atoms. The interactions between the pairs of residue beads are quantified by Lennard-Jones (LJ) potentials and long-range electrostatic potentials. The electrostatic interactions are described by the Debye-Hückel equation at a screening length of 1 nm, which corresponds to physiological salt concentrations. The LJ interaction parameters are adapted from knowledge-based statistical contact potentials and scaled relative to the electrostatic interaction energy. Importantly, folded-protein domains are treated as rigid bodies, whereas disordered segments (e.g., flexible loops, linkers, and termini) are represented by bead polymers. In addition to the LJ potentials and the electrostatic potentials, the potential energy of the disordered segments contains contributions from bond-length potentials, bond-angle potentials, and torsion-angle potentials.

As in recent coarse-grained simulations of the ACBD3: PI4KB complex,³⁰ here the structural model, that is, the fulllength ACBD3 protein comprises four rigid domains: (a) the crystallized portion of the GOLD domain with the PDB code 5LZ1 (residue numbers ranging 364-448 and 473-528); (b) the glutamine-rich domain (Q domain, residue numbers ranging 241-308) given by the nuclear magnetic resonance structure with the PDB code 2N73; (c) the charged amino acid region (CAR, residue numbers ranging 180-240) represented by a single alpha-helix, as proposed in Reference 32; and (d) a homology model of the ACBD domain (residue numbers ranging 82-179) based on the PDB entry 3FLV (29% sequence identity, 96% sequence coverage). All of the missing loops, disordered linkers, and terminal segments were built into the ACBD3 model using ModLoop³³ and simulated as chains of amino acid beads with appropriate stretching, bending, and torsional potentials.²⁰

Within the framework of the KH model, as described above, we performed replica exchange Monte Carlo (REMC) simulations with 16 replicas at temperatures ranging from 300 to 750 K. The basic MC moves were translations and rotations of the rigid domains. For flexible segments, in addition to local MC moves on each of the residue beads, crankshaft moves were employed to enhance sampling. After the initial 10^5 MC sweeps for equilibration, the REMC production run comprised 5×10^6 MC sweeps. The ACBD3 protein conformations were recorded every 10^3 MC sweeps. We thus obtained a pool of 8×10^4 conformations for further analysis.

The scattering intensity profile was computed for each of the simulated conformations individually using an algorithm codeveloped with the ensemble refinement of SAXS (EROS) method.²³ The scattering intensity profile corresponding to a set *S* of conformations was then obtained from the scattering intensity profiles $I_k(q)$ of the individual conformations belonging to set *S*, that is,

$$I_S(q) = \sum_{k \in S} I_k(q)$$

The discrepancy between the experimental SAXS data, $I_{exp}(q)$, and the scattering intensity profile $I_S(q)$ of a given set of simulated conformations was quantified by

$$\chi_{S}^{2} = \sum_{i=1}^{N_{q}} \frac{(I_{\exp}(q_{i}) - aI_{S}(q_{i}) - b)^{2}}{\sigma^{2}(q_{i})}.$$

Here, N_q is the number of SAXS data points, $\sigma(q)$ is the statistical error of the extrapolated SAXS intensity $I_{exp}(q)$, whereas the scale factor *a* and offset *b* result from the conditions $\partial \chi^2 / \partial a = 0$ and $\partial \chi^2 / \partial b = 0$. The offset parameter *b* accounts for inaccuracies in the buffer subtraction and/or uncertainties in the electron density contrast. We applied a minimum-ensemble method^{21,34} to identify the smallest set of conformations consistent with the experimental SAXS data. We treated the electron density contrast of the protein hydration shell as a fitting parameter and varied it in the range between 0.01 and 0.05 $e/Å^3$.

We performed analogous simulations and analyses for the ACBD3:3A/AiV protein complex. In the coarse-grained simulations, the GOLD domain in complex with the N-terminal part of the nonstructural 3A protein was treated as one rigid body based on the crystal structure with the PDB code 5LZ3. The C-terminal tail of the 3A protein was simulated as a chain of amino acid beads. Other portions of the ACBD3 protein were represented in the manner as previously in the coarse-grained simulations of the full-length ACBD3 protein.

From the REMC simulations, we obtained a pool of 1.6×10^5 conformations of the ACBD3:3A/AiV protein complex. We postprocessed and analyzed these conformations in the same way as the simulated conformations of the full-length ACBD3 protein in the absence of the 3A/AiV protein. Finally, we identified the smallest set of conformations consistent with the experimental SAXS data using the minimum-ensemble method.^{21,34}

3 | **RESULTS AND DISCUSSION**

We performed SAXS measurements at the ACBD3 concentrations of 5.8, 3.4, and 2.6 mg/mL. The resulting SAXS intensity profiles, I(q), were found to overlay to a great extent after rescaling (Figure S1a). However, a closer inspection revealed small systematic differences between the three I(q) data sets at the smallest scattering angles (Figure S1a–d). These systematic differences most likely reflect interactions between protein molecules in solution. Therefore, we extrapolated the three I(q) profiles to the dilute limit, that is, the formal limit of the protein concentration approaching zero (Figure S1e,f, and Figure 2a). The Guinier plot (Figure 2a, inset) of the extrapolated SAXS data yields a radius of gyration of about 69 Å, which suggest that the full-length ACBD3 protein can attain relatively extended conformations in solution.



FIGURE 2 SAXS analysis and modeling of the full-length ACBD3 protein. (a) The points in red constitute the SAXS intensity profile that results from extrapolating to the dilute limit three data sets collected at protein concentrations of 5.8, 3.4, and 2.6 mg/mL. The curve in black represents the theoretical SAXS intensity profile corresponding to an ACBD3 conformational ensemble that satisfactorily fits the experimental SAXS data ($\chi^2 = 1.7$). The logarithmic scale is used on both the vertical and horizontal axes. The inset shows the Guinier plot, that is, $\log(I(q)/I(0))$ versus q^2 . The value range on the horizontal axis corresponds to the region $qR_g < 1.3$ where the Guinier approximation is valid for globular proteins. The dashed line corresponds to the Guinier approximation with $R_g = 69$ Å. (b) The difference between the experimental and simulated SAXS intensity profiles as computed from the data shown in panel A. (c) The Kratky plot, that is, $q^2I(q)$ versus q, corresponding to the data shown in panel A. (d) Two structural models that constitute the minimal representation of the ACBD3 conformational ensemble. These two models jointly yield the SAXS intensity profile shown in panel A in black and, thus, fit the experimental SAXS data with $\chi^2 = 1.7$. The ACBD3 protein domains are indicated

We used the extrapolated SAXS intensity profile in combination with computer simulations to identify representative conformations of the full-length ACBD3 protein in solution. To this end, we performed coarse-grained simulations of the full-length ACBD3 and quantitatively compared the SAXS intensity profiles of the simulated conformations to the experimental SAXS intensity profile. It turned out that none of the individual conformations that were obtained from the simulations could be fitted to the extrapolated SAXS data with $\chi^2 < 2.9$. Since the full-length ACBD3 protein contains disordered linkers, loops, and termini, we expected it to exhibit conformational flexibility and diversity. Therefore, we applied an EROS method,^{21,34} and thus identified two conformations that jointly could account for the extrapolated SAXS intensity profile (Figure 2a). We found that the differences between the experimental and simulated SAXS intensity profiles were very small in the whole range of scattering angles (Figure 2b), yielding $\chi^2 = 1.7$. Importantly, the Kratky plot (Figure 2c) shows the typical features of partially disordered proteins, which additionally supports our conclusion that the full-length ACBD3 protein is flexible in solution. We found out that the minimum ensemble consistent with the experimental SAXS data was represented by two conformations, namely, one "extended" conformation with $R_g = 74$ Å and $D_{max} = 235$ Å, and one "compact" conformation with $R_g = 64$ Å and $D_{max} = 190$ Å (Figure 2d). These two conformations taken together with equal statistical weights fit the extrapolated SAXS intensity profile with $\chi^2 = 1.7$ (Figure 2a,b). Interestingly, the individual domains of the ACBD3 protein do not make any extensive contacts in neither of these conformations.

Since the full-length ACBD3 protein contains flexible linkers, loops and termini, it can attain multiple conformations, ranging from extended to compact ones. The two conformations identified in our SAXS analysis (Figure 2d) cannot fully represent the entire conformational ensemble of the ACBD3 protein in solution. However, the vast differences between the two conformations (Figure 2d) provide a glimpse of the conformational diversity of this highly flexible protein.

We also performed SAXS experiments on the ACBD3:3A/AiV complex at protein concentrations of 4.7, 2.7, 1.8, and 0.9 mg/mL. These protein concentrations are more than an order of magnitude larger than the dissociation constant of the ACBD3:3A/AiV protein complex (K_d in the low micromolar range).¹¹ We extrapolated the four I(q) profiles to the dilute limit (Figure S2 and Figure 3a). The Guinier plot (Figure 3a, inset) of the extrapolated SAXS data shows that the radius of gyration is about 70.5 Å, which is 1.5 Å more than the radius of gyration of the ACBD3 protein in the absence of the 3A protein, and which indicates that the ACBD3:3A/AiV protein complex attains extended conformations in solution.

The extrapolated SAXS intensity profile for ACBD3:3A/ AiV (Figure 3a) looks very similar to the extrapolated SAXS intensity profile for ACBD3 (Figure 2a). Since each of the two SAXS intensity profiles contains the same number of data points ($N_q = 1,034$), one way to quantify their similarity is to calculate χ^2 . It turns out that $\chi^2 = 3.1$ for these two SAXS curves, which indicates that the two SAXS profiles are clearly distinguishable within the experimental error despite their apparent similarity.

We next used coarse-grained molecular simulations to structurally interpret the extrapolated SAXS intensity profile. The coarse-grained simulations of the ACBD3:3A/AiV protein complex provided a large and diverse pool of conformations,

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but no single conformation from this pool was fully consistent with the experimental SAXS data ($\gamma^2 > 3.4$ for any of the simulated conformations). However, we found that a set of two conformations with equal statistical weights fitted the SAXS data with $\gamma^2 = 1.7$ (Figure 3a). We checked that the differences between the experimental and simulated SAXS intensity profiles were very small in the whole range of scattering angles (Figure 3b). The Kratky plot (Figure 3c) shows the typical features of partially disordered proteins, which supports our conclusion that the ACBD3:3A protein complex is flexible in solution. The two representative conformations of the ACBD3:3A protein complex are quite different, namely, the "extended" conformation has $R_{\rm g} = 79$ Å and $D_{\rm max} = 250$ Å, whereas the "compact" conformation has $R_{\rm g} = 63$ Å and $D_{\text{max}} = 185 \text{ Å}$ (Figure 3d). However, neither of these conformations involves any extensive interdomain contacts. These observations indicate that the ACBD3:3A protein complex exhibits large conformational fluctuations in solution.

Taken together, our data clearly show that the ACBD3 protein is extended and flexible in solution—in both the presence and absence of the Aichi virus 3A protein. We did not observe any extensive contacts between the individual domains of the ACBD3 protein alone or when in complex with viral 3A protein, which implies that the binding of the Aichi virus 3A protein to the ACBD3 protein does not significantly affect the conformations of ACBD3 in solution. Our recent SAXS studies³⁰ show that the ACBD3:PI4KB protein complex exhibits large conformational fluctuations



FIGURE 3 Analogous to Figure 2 but for the full-length ACBD3:3A protein complex

and little interdomain contacts. Therefore, we also predict that the binding of ACBD3 to PI4KB should not significantly affect the conformational diversity of ACBD3 in solution.

ACBD3 is essential for the replication of all enteroviruses and kobuviruses, which have been tested in this respect.^{4,35–38} These results imply that viruses do not interfere with the conformation of ACBD3, but are consistent with the hypothesis that viruses use ACBD3 to recruit the lipid kinase PI4KB. This is also consistent with the fact that these viruses are sensitive to chemical inhibition of PI4KB.^{39–42} However, because the 3A recruitment of ACBD3 also changes the cellular localization of ACBD3, it could also interfere with other, Golgi related, functions of ACBD3 such as intracellular transport for the benefit of the virus.

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CONFLICT OF INTEREST

The authors declare no conflict of interest.

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SUPPORTING INFORMATION

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