

Computational Biology

CHARMM-GUI *Glycan Modeler* for modeling and simulation of carbohydrates and glycoconjugates

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Abstract

Characterizing glycans and glycoconjugates in the context of three-dimensional structures is important in understanding their biological roles and developing efficient therapeutic agents. Computational modeling and molecular simulation have become an essential tool complementary to experimental methods. Here, we present a computational tool, *Glycan Modeler* for in silico *N/O*-glycosylation of the target protein and generation of carbohydrate-only systems. In our previous study, we developed *Glycan Reader*, a web-based tool for detecting carbohydrate molecules from a PDB structure and generation of simulation system and input files. As integrated into *Glycan Reader* in CHARMM-GUI, *Glycan Modeler* (*Glycan Reader & Modeler*) enables to generate the structures of glycans and glycoconjugates for given glycan sequences and glycosylation sites using PDB glycan template structures from Glycan Fragment Database (<http://glycanstructure.org/fragment-db>). Our benchmark tests demonstrate the universal applicability of *Glycan Reader & Modeler* to various glycan sequences and target proteins. We also investigated the structural properties of modeled glycan structures by running 2- μ s molecular dynamics simulations of HIV envelope protein. The simulations show that the modeled glycan structures built by *Glycan Reader & Modeler* have the similar structural features compared to the ones solved by X-ray crystallography. We also describe the representative examples of glycoconjugate modeling with video demos to illustrate the practical applications of *Glycan Reader & Modeler*. *Glycan Reader & Modeler* is freely available at <http://charmm-gui.org/input/glycan>.

Key words: Glycan Reader, molecular dynamics simulation, *N*-linked glycosylation, *O*-linked glycosylation, protein–glycan interaction, template-based glycan modeling

Introduction

Carbohydrate moieties, also referred to as glycans, are one of the most abundant cell components (Ohtsubo and Marth 2006). They can be highly branched and covalently attached to protein (glycoprotein) or lipid (glycolipid), and can exist as free ligands (Dwek 1996). The covalent attachment of glycans to proteins (protein glycosylation) is one of the most important and abundant post-translational protein modifications. More than half of all proteins are expected to be glycosylated based on SWISS-PROT database (Apweiler et al. 1999). Glycolipids and unbranched polysaccharides such as glycosaminoglycans (GAGs) have been found in many different bacterial species and their structural forms have the enormous diversity (Curatolo 1987; Muthana et al. 2012). Therefore, it is not surprising that glycans play a critical role in living organisms. Indeed, numerous studies have shown that glycans are associated with a variety of important biological processes, such as immune recognition and response (Rabinovich and Toscano 2009), tumor growth and metastasis (El Ghazal et al. 2016), protein quality control and trafficking (Trombetta 2003), and cell–cell communications (Collins and Paulson 2004).

To understand how glycosylation is functionally involved in these physiological and pathological processes, knowledge of the three-dimensional structures of glycans, their dynamic properties, and their interactions with proteins is essential (Malik and Ahmad 2007; Nagae and Yamaguchi 2012; Malik et al. 2014; Jo et al. 2016; Qi et al. 2016; Dong et al. 2018). Experimental methods such as X-ray crystallography and nuclear magnetic resonance (NMR) spectroscopy have been widely used to provide the atomic details of biomolecular structural information (Wormald et al. 2002). During last decades, X-ray crystallography has been improved substantially with recent advances such as the synchrotron radiation (Perez and de Sanctis 2017). However, despite these technical advances, solving the complex carbohydrate structures remains nontrivial. For example, during the crystallization of glycoproteins, heterogeneity of glycans on the protein surface hinders the crystal packing, and the electron density maps of these glycans are not often fully resolved because of their high flexibility (Perez and de Sanctis 2017). NMR spectroscopy also has been advanced in terms of the labeling strategies such as labeled glycans with stable NMR-active isotopes. For instance, the use of ^{13}C -labeled glycans provides the large signal dispersion and high sensitivity and also makes it possible to observe the fast exchanges of ligand complexes in the NMR chemical shift timescales (Arda and Jimenez-Barbero 2018). Nevertheless, some limitations are still remained in terms of the relatively low intrinsic sensitivity and the dependency on the availability of isotopic labeling or specific chemical tags required in the case of large biomolecules (Marchetti et al. 2016).

In order to alleviate these difficulties, combination of experimental techniques with molecular modeling and simulation methodologies such as Monte Carlo and molecular dynamics (MD) simulations has emerged as an indispensable tool to investigate glycan conformations in a biological environment (Imberty and Perez 2000; Marchetti et al. 2016; Hamark et al. 2017). Over the years, several carbohydrate force fields such as GLYCAM06 (Kirschner et al. 2008), CHARMM36 (Guvench et al. 2009), GROMOS 53A6_{GLYC} (Pol-Fachin et al. 2012), OPLS-AA-SEI (Kony et al. 2002), and MM4 (Allinger et al. 2003) have been developed. However, building the simulation systems containing glycans and glycoconjugates still remains nontrivial due to the structural complexity of glycans (i.e., branches and various types of glycosidic

linkages (Campbell et al. 2015)). To facilitate the computational modeling of glycans, standalone programs such as POLYS (Engelsen et al. 2014), *doGlycans* (Danne et al. 2017), CarbBuilder (Kuttel et al. 2016), and RosettaCarbohydrate (Labonte et al. 2017) have been developed. Several web-based tools, such as GlyProt (Bohne-Lang and von der Lieth 2005) and SWEET-II (Bohne et al. 1999) at GLYCOSCIENCES.de (Lutteke et al. 2006), GLYCAM Carbohydrate Builder (Woods-Group 2005–2018) at GLYCAM-web, Glycan Fragment Database (GFDB) (Jo and Im 2013) and GS-align (glycan structure alignment) (Lee and Jo et al. 2015) at GlycanStructure.ORG (Im-Group 2011–2018), *Glycan Reader* in CHARMM-GUI (Jo et al. 2011), and PROCARB (Malik et al. 2010) provide glycan-related applications and databases, allowing users to model glycan structures with experimentally determined glycan structures as a template without any installation of program in user-side. However, except CHARMM-GUI and GLYCAM-web, these tools require significant post-processing efforts to prepare simulation systems with the generated glycan structure. Currently, CHARMM-GUI is the only available tool for the valid input setup of major MD simulation programs (Lee et al. 2016) such as CHARMM (Brooks et al. 2009), NAMD (Phillips et al. 2005), GROMACS (Hess et al. 2008), AMBER (Case et al. 2005), GENESIS (Jung et al. 2015), LAMMPS (Plimpton 1995), Desmond (Bowers et al. 2006), OpenMM (Eastman et al. 2013) and CHARMM/OpenMM (Arthur and Brooks 2016) (using the CHARMM force field).

CHARMM-GUI, <http://www.charmm-gui.org>, is a web-based graphical user interface and provides various functional modules to prepare complex biomolecular systems and input files for molecular simulations. During the 13 years of successful services, CHARMM-GUI has contributed to facilitate molecular modeling and simulation of glycans and glycoconjugates by developing *Glycolipid Modeler* (Lee et al. 2019), *LPS Modeler* (Lee et al. 2019) and *Glycan Reader* (Jo et al. 2011). In particular, *Glycan Reader* has greatly simplified the reading of PDB and PDBx/mmCIF structure files containing glycans by automatic detection of carbohydrate molecules and glycosidic linkage information. In the recent update (Park et al. 2017), *Glycan Reader* not only detects most sugar types and chemical modifications in the PDB, but also allows users to edit the glycan sequences through addition/deletion/change of sugar types, chemical modifications, glycosidic linkages, and anomeric states. However, CHARMM-GUI *Glycan Reader* did not support in silico glycosylation and addition of a sugar at the reducing end of an existing glycan chain.

In this work, we present *Glycan Modeler*, a new CHARMM-GUI module for in silico *N/O*-glycosylation and generation of carbohydrate-only systems. CHARMM-GUI *Glycan Modeler* can generate the most appropriate representative glycan structures through GFDB database searches and determines proper orientations relative to target protein. Although GFDB stores all available complex carbohydrate structures in the PDB after the validation processes such as reannotation of carbohydrate residue names based on their Cartesian coordinates using *Glycan Reader*, not every possible carbohydrate sequence is available in GFDB. In the absence of target glycan sequence in GFDB, our module generates the structures by using the valid internal coordinate information in the CHARMM force field (see Materials and methods for details). *Glycan Reader & Modeler* (as *Glycan Modeler* is integrated into *Glycan Reader*) also allows addition of a sugar at the reducing end of a pre-existing glycan chain. We performed benchmark analysis and MD simulations to validate the reliability of modeled *N*-glycan structures. In addition, we also describe the representative examples

of glycoconjugate modeling with video demos to illustrate the practical applications of *Glycan Reader & Modeler* (<http://www.charmm-gui.org/demo>).

Results and discussion

Glycan Modeler can build *N/O*-glycans at multiple glycosylation sites with or without pre-existing glycan(s) in a protein structure. The inputs for *N/O*-glycan modeling are a protein structure, glycosylation site(s), and glycan sequence for each glycosylation site (see Materials and methods for details). In addition to *N/O*-glycans, *Glycan Modeler* can build a carbohydrate-only structure with a user-specified glycan sequence.

Benchmark test of *Glycan Modeler* against PDB non-redundant *N*-glycan set

To evaluate the quality of glycan structure models generated by *Glycan Modeler*, we measured the structural similarities of glycan models to their native PDB structures for a set of target *N*-glycans. We first prepared a set of target proteins and their *N*-glycan structures by collecting non-redundant (unique) *N*-glycan sequences containing at least five carbohydrates from the PDB and randomly selecting one PDB entry for each unique sequence. We used the PDB glycan sequence and glycosylation site information to generate *N*-glycans structure models onto their parent protein structure (see Materials and methods for how we set up the benchmark set in detail).

Similarities between the *N*-glycan models and their target PDB glycan structures were measured by the root-mean-squared deviation (RMSD) of glycans without superposition between the modeled and PDB glycan structures; i.e., the measured RMSD represents structural deviation of modeled glycan from its native one, resulting from differences in their conformation and orientation with respect to the target protein. We also examined if pre-existing glycan can improve the quality of the modeled glycan structures and orientations, as a small orientational difference of first one or two *N*-glycan residues could lead a big difference of the entire *N*-glycan orientation. We tested three approaches by building glycan models from (1) Asn residue (“from Asn”), (2) first sugar unit from pre-existing glycan in the PDB file (“sugar 1”) or (3) first two sugar units from pre-existing glycan in the PDB file (“sugar 2”). Figure 1 shows RMSD comparisons of the three approaches. The RMSD values of some *N*-glycan models are as high as 20 Å due to their orientational differences with respect to proteins, while the RMSD values after superposition between modeled and crystal *N*-glycan structure (i.e., only

for glycan conformational differences) are mainly within 1–3 Å (Figure S1). Most RMSD values of “sugar 1” and “sugar 2” are much smaller than those of “from Asn”, indicating that the use of a pre-existing partial glycan can provide better orientation in the modeled glycans. This is also in agreement with our previous PDB glycan structure survey study (Jo et al. 2013) that the difficulties in finding the global orientation of the glycan with respect to the protein, even when the homologous *N*-glycan templates are present, can be significantly alleviated when a partial glycan structure is available.

Benchmark test of *Glycan Modeler* against PDB deglycosylated protein set

In our previous work, Lee et al. measured the local and global structural similarities between glycosylated and deglycosylated forms of identical proteins (GP/P pairs) in the PDB to investigate the impact of *N*-glycosylation on protein structure (Lee and Qi et al. 2015). To examine if *Glycan Modeler* is applicable to in silico glycosylation on a deglycosylated form of proteins, we performed another benchmark test using the same GP/P pairs. We discarded the pairs containing only monosaccharide, resulting in a total of 1,543 GP/P pairs. Using *Glycan Modeler*, we generated *N*-glycan models on the deglycosylated proteins based on the glycan sequence and glycosylation site information from the paired glycoproteins. Since the pairs often did not have the exactly same sequences, the glycosylation sites were determined based on the aligned structures of GP/P pairs by TM-align (Zhang and Skolnick 2005). Figure 2 shows the examples of predicted *N*-glycans on the crystal structures of a target deglycosylated protein (human alpha 1-antitrypsin (Engh et al. 1989)). In the deglycosylated protein structures, the side chains of Asn residues at the glycosylation sites have different orientations even though the structures come from an identical protein (Figure 2A). Because *Glycan Modeler* determines the orientations by randomly searching the torsional angle space of Asn side chain with an energy cutoff, predicted χ_1 and χ_2 of target Asn residues can be diverse, as shown in Figure 2B. Nonetheless, for the 1,543 GP/P pairs, all *N*-glycan chains were successfully generated without structural/chirality errors and bad contacts with their parent proteins, demonstrating the universal applicability of *Glycan Modeler* in generating glycan structures regardless of pre-existing glycans.

Simulation study of HIV envelope protein with glycans from PDB and *Glycan Modeler*

To further examine the reliability of modeled glycan structures, we performed MD simulations of two fully glycosylated HIV envelope

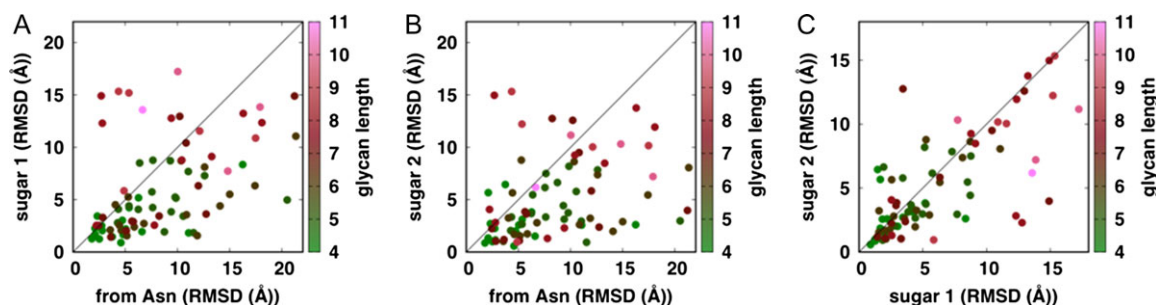


Fig. 1. Benchmark result of *Glycan Modeler*. For 82 non-redundant *N*-glycan structures, glycan RMSDs were measured against actual crystal *N*-glycan structures. The length of each target *N*-glycan is represented by color. Note that the RMSD was calculated without superposition between crystal and modeled *N*-glycan structures, so the RMSD value includes the glycan conformational and orientational differences on the protein surface.

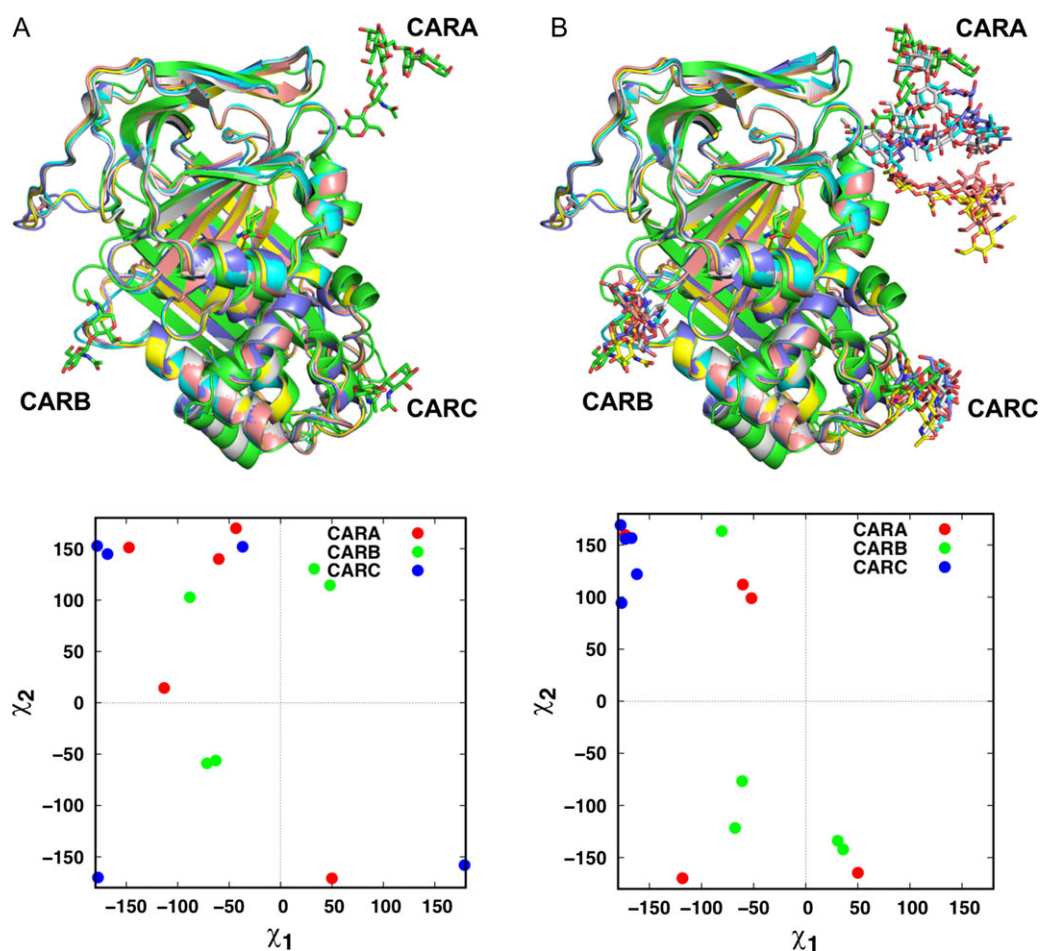


Fig. 2. An example of in silico glycosylation on deglycosylated protein targets. (A) Green is a glycosylated protein (PDB: 7API) and others are deglycosylated proteins (PDB: 1QLP, 2QUG, 3CWL, 3CWM, 3NE4) in the PDB. On the left bottom, the scatter plot shows the distribution of χ_1 and χ_2 of target Asn residues of deglycosylated proteins. (B) The *N*-glycan models were generated on deglycosylated proteins with the identical glycan sequences and glycosylation sites. The χ_1 and χ_2 of target Asn residues after the modeling are displayed as scatter plots.

trimer systems. One system (S^{cryst}) was built based on all 66 glycan structures directly taken from the crystal structure (PDB:5FYL (Stewart-Jones et al. 2016)) (see the next section for system building in detail). In the other system (S^{glymod}), all glycan structures were removed and modeled by *Glycan Modeler* using the same sequence as those in the crystal structure. For a sufficient sampling of glycan conformations, each system was simulated for 2 μs using Anton 2 (Shaw et al. 2014) (see Materials and methods for simulation details; see also Supporting Movies, [HIVenv_cryst.mpg](#) and [HIVenv_glymod.mpg](#)). As shown in Figure 3A, the RMSDs of the envelope trimers reached a comparable value after 2 μs (i.e., the average RMSDs during the last 1 μs are $4.59 \pm 0.24 \text{ \AA}$ (S^{cryst}) and $5.42 \pm 0.40 \text{ \AA}$ (S^{glymod})), indicating the protein is stable in both systems.

To compare the conformations of the glycans, we calculated the φ , ψ dihedrals of all 243 glycosidic linkages in each system. The distributions of these dihedrals for the same glycosidic linkage in both systems were compared using the Pearson correlation coefficient, i.e., if the two systems sample very similar dihedral distributions, the coefficient would be close to 1. Only two out of the 486 dihedrals have correlations less than 0.9 (Figure 3B), and both are the ψ dihedral in the Man β (1 \rightarrow 4)GlcNAc linkage. The distributions of these two dihedrals show that the deviations mostly come from minor

populations at -60° (Figure 3C, D). To examine the preference of this dihedral, we searched for all Man β (1 \rightarrow 4)GlcNAc linkages in the PDB using GFDB (Jo and Im 2013) and found that ψ indeed has a minor population ($\sim 2.5\%$) around -60° (Figure S2), which may require much longer simulations to reach sampling convergence in the context of glycoprotein. Taken together, these results demonstrate that there is no significant difference in the torsion angle distributions of glycosidic linkages between crystal and modeled glycans.

Examples of practical applications of *Glycan Modeler*

In this section, we describe the representative examples (Figure 4) to illustrate practical applications of *Glycan Reader & Modeler* for glycoconjugate modeling and simulation with the following four video demos in CHARMM-GUI (<http://www.charmm-gui.org/demo>): (1) solution system building of the HIV envelope protein, (2) modeling *N*-glycans on immunoglobulin G1 Fc, (3) membrane system building of a cholera toxin B and ganglioside GM1 complex, and (4) solution system building of a heparin molecule. It should be noted that *Glycan Reader & Modeler* is embedded in *PDB Reader*, which makes *Glycan Reader & Modeler* available in most other CHARMM-GUI modules such as *Solution Builder*, *Membrane*

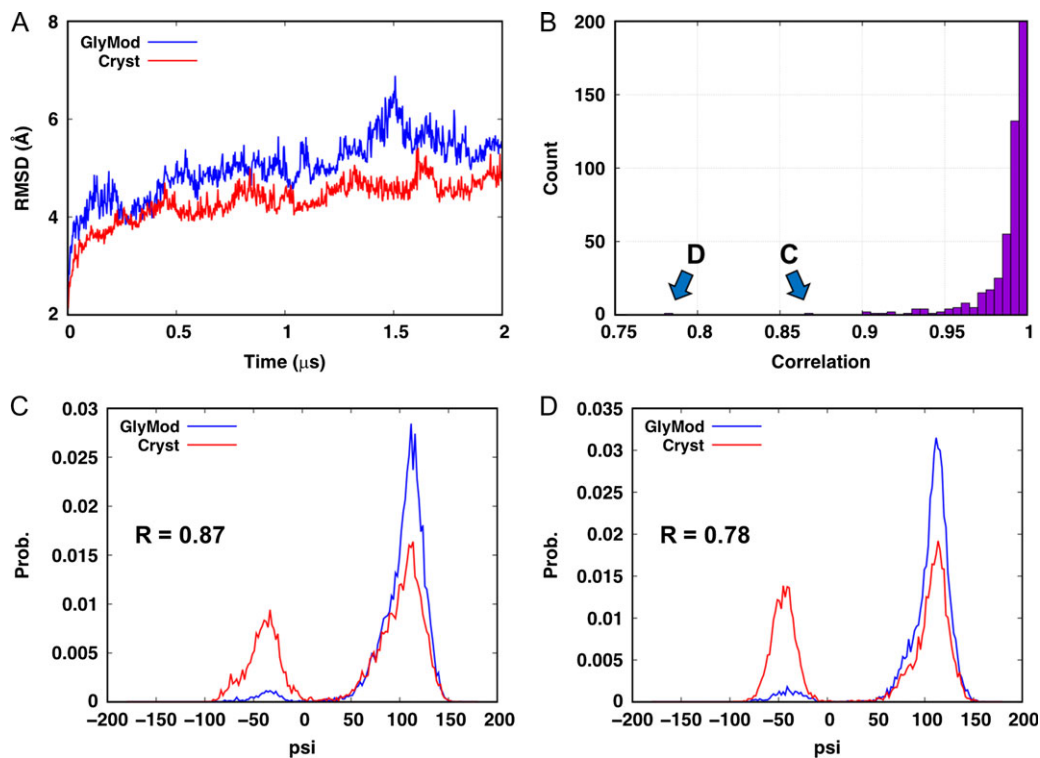


Fig. 3. Simulations of the HIV envelope trimer. (A) Backbone RMSDs of the protein. (B) Distribution of the linear correlation between the glycosidic dihedrals in the two MD systems. (C, D) Distributions of the ψ dihedrals that have a correlation of less than 0.9.

Builder (Jo et al. 2007; 2009; Wu et al. 2014; Lee et al. 2019), *HMMM Builder* (Qi et al. 2015), *Nanodisc Builder* (Qi et al. 2019), *Micelle Builder* (Cheng et al. 2013), and *PBEQ-Solver* (Im et al. 1998; Jo et al. 2008).

Solution system building of the HIV envelope protein: To build a solution system of a glycoprotein like the HIV envelope protein (PDB:5FYL (Stewart-Jones et al. 2016)), users can use *Solution Builder*. The PDB contains a closed prefusion structure of the HIV envelope protein (Figure 4A). The envelope protein is synthesized as a gp160 precursor and processed into a trimer of heterodimer consisting of glycoproteins gp120 and gp41 that are heavily glycosylated and mediate receptor binding and membrane fusion. Note that the gp41 transmembrane domain is truncated in this structure and thus not included in this example. In addition to the envelope protein, this PDB file contains antibodies PGT122 and 35O22 that interact with the envelope protein. In this system building example, these antibodies are excluded for convenience. PDB:5FYL contains only one copy of the trimeric envelope protein, so users need to generate the remaining two copies during the PDB manipulation step. One can follow the step-by-step procedure in Section 1 of Supplementary Data to build a solution system of the HIV envelope protein.

Modeling N-glycans immunoglobulin G1 Fc Immunoglobulin G1 (IgG1) is a subclass of human serum antibodies and is the most widely used platform for developing therapeutic monoclonal antibodies (Wei et al. 2017). The antibody-mediated effector functions require binding of the crystallizable fragment (Fc) of IgG1 to Fc γ receptors that are expressed on the surface of recruited cells. IgG1 Fc is a symmetric homodimer and an N-glycan is attached to Asn297 in each chain. Our recent simulation study reports that changes in N-glycan composition alter the conformational ensembles of C'E loop and C γ 2- C γ 3 orientation in Fc chains, eventually affecting the binding affinity to its Fc receptor (Lee and

Im 2017). The results also suggest that computational optimization of Fc N-glycans can guide engineering of IgG1-based antibodies for their better efficiency. In this example, a modified PDB file (see Section 2 of Supplementary Data) that contains only the first N-acetylglucosamine (GlcNAc) on chain B is used to build the original N-glycans in both chains using *Glycan Reader & Modeler* (Figure 4B).

Membrane system of a cholera toxin B and ganglioside GM1 complex: Cholera toxin (CT) produced by *Vibrio cholerae* is an AB₅ toxin with a single enzymatic A subunit and a pentameric B subunit (CTB). CTB can bind to ganglioside GM1 receptors in the plasma membrane for CT's entry to the cell. PDB:3CHB (Merritt et al. 1998) has five CTB subunits and associated GM1 carbohydrates with no ceramide lipid part. In addition, two GM1 glycan structures do not have β -glucose (Glc) at the reducing end. This example illustrates how to build a complete CTB:GM1 pentameric complex using *Glycan Reader & Modeler* and to insert the ceramide lipid portions into a membrane bilayer using *Membrane Builder* (Jo et al. 2009; Wu et al. 2014) (Figure 4C); see Section 3 of Supplementary Data for the detailed steps. Note that it is relatively easy to move one acyl chain of GM1 initially located outside the membrane into the bilayer during the equilibration.

Solution system of a heparin molecule: PDB:1AXM (DiGabriele et al. 1998) contains a heparin molecule complexed with a fibroblast growth factor. Heparin is a linear polysaccharide sulphated on alternating IdoA (L-iduronic acid) and GlcNS (6-O, N-sulfonated glucosamine), which can modulate endothelial cell proliferation and migration (Giroux et al. 1998). One can build a solution system only with this heparin molecule using *Solution Builder*. However, this example illustrates how to use *Glycan Reader & Modeler* to build a carbohydrate-only solution system (Figure 4D); see Section 4 of Supplementary Data for the detailed steps.

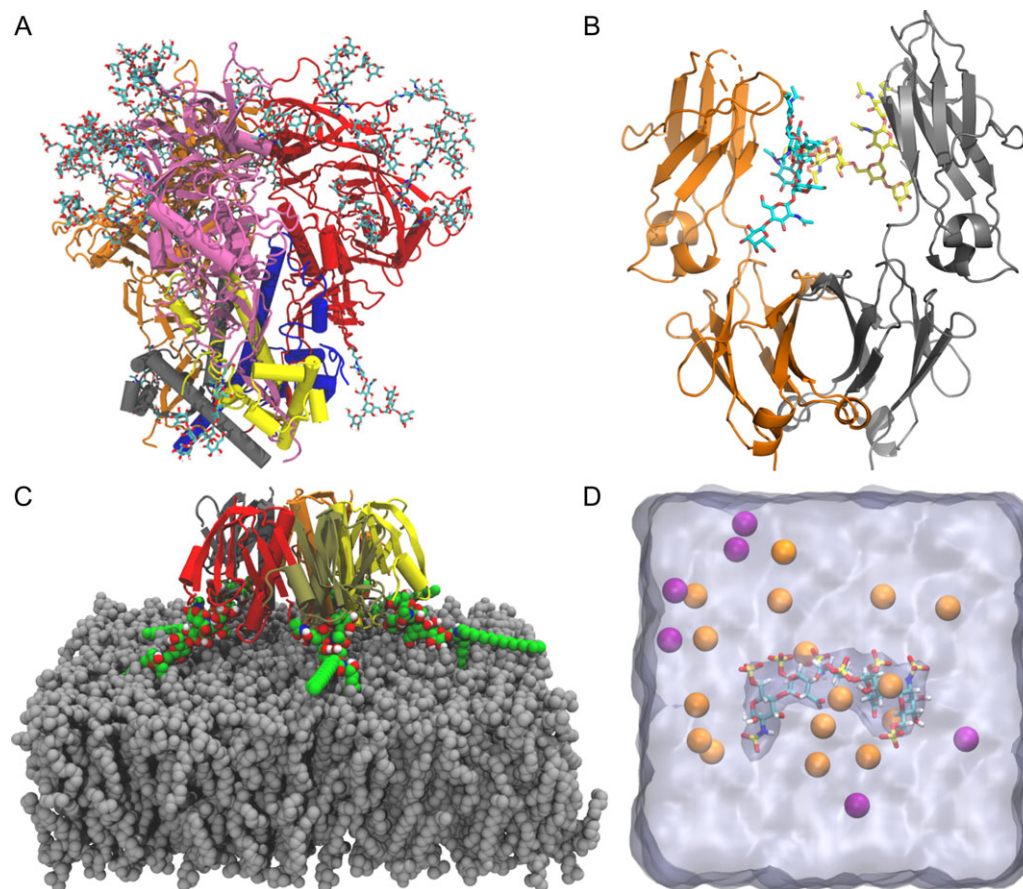


Fig. 4. Practical applications of *Glycan Reader & Modeler*. (A) HIV envelope protein trimer of heterodimer with *N*-glycans. Each heterodimer is composed of gp120 (gray, blue, and yellow) paired with gp41 (orange, red, and pink, respectively). (B) Immunoglobulin G Fc dimer (orange and gray) with *N*-glycans. (C) Cholera toxin B and ganglioside GM1 pentameric complex shown with each protein segment colored differently. GM1 glycolipids are shown as green sphere. A membrane bilayer (gray) is composed of DMPC. (D) Heparan sulfate pentasaccharide surrounded by K⁺ (orange) and Cl⁻ ions (purple). Water box is shown as a translucent surface.

Conclusions

We have described the development of a new CHARMM-GUI module, *Glycan Modeler* that provides the functionality of in silico *N/O*-glycosylation onto the target protein and carbohydrate-only system building. Our benchmark tests show that the *Glycan Modeler* framework can be efficiently used to model multiple glycan structures onto the target protein with or without pre-existing glycan structures. We also carried out the 2- μ s MD simulations of a fully glycosylated HIV envelope protein to compare the structural properties of glycan structures modeled by *Glycan Modeler* to those from the crystal structure. The simulation results show no significant difference in glycan conformations, demonstrating the reliability of the glycan structure models generated by *Glycan Modeler* for molecular simulation studies. In addition to the benchmark evaluations, we described practical representative examples with video demos (<http://www.charmm-gui.org/demo>) to illustrate how to build complex glycoconjugate systems using *Glycan Reader & Modeler*.

Glycan Reader & Modeler can handle the glycan modeling with all kinds of glycan residues described in Symbol Nomenclature for Glycans (SNFG) (Varki et al. 2015) through addition/deletion/changes of sugar types, chemical modifications, glycosidic linkages, and anomeric states. Furthermore, the method allows the generation of simulation systems and inputs of user-defined glycoconjugate

systems as it is integrated into other CHARMM-GUI modules such as *Solution Builder* or *Membrane Builder*. We hope that *Glycan Reader & Modeler* can become a useful tool enabling researchers to easily carry out innovative glycan modeling and simulation studies to acquire novel insight into structures, dynamics, and underlying mechanisms of complex glycan-related biological systems.

Materials and methods

N/O-glycan modeling

Glycan Modeler uses PDB glycan structures as templates for glycan structure modeling. To identify the relevant template structures, *Glycan Modeler* starts with Glycan Fragment Database (GFDB) searches between a query glycan sequence (for each glycosylation site, Figure 5A) and the PDB glycan structures in the database (Jo and Im 2013). PDB glycan structures containing the query glycan sequence are all searched in this step (Figure 5B). Glycosidic torsion angle based clustering (Jo and Im 2013) is performed to generate five representative glycan structures among the searched glycans (Figure 5C). Briefly, the clustering algorithm is composed of the following steps: (1) construction of the pairwise distance matrix where a distance is defined as the torsion angle difference between the same glycosidic linkages for all-to-all template structure pairs, (2)

identification of the first cluster determined by the maximum number of neighbors within a cutoff radius of 30° , (3) identification of the second cluster after excluding the first cluster members, and (4) repeat of the clustering analysis for third, fourth and fifth clusters. After the clustering analysis, representative glycan models are generated by CHARMM using average glycosidic torsion angles from

each cluster (Figure 5C). If an uploaded protein structure contains pre-existing glycan structure(s) and the pre-existing glycan is the part of a query glycan sequence, *Glycan Modeler* maps the pre-existing glycan onto the glycan models to conserve the coordinates of the pre-existing glycan structures. If there is no match in GFDB searches for a given glycan sequence, its structure is generated based

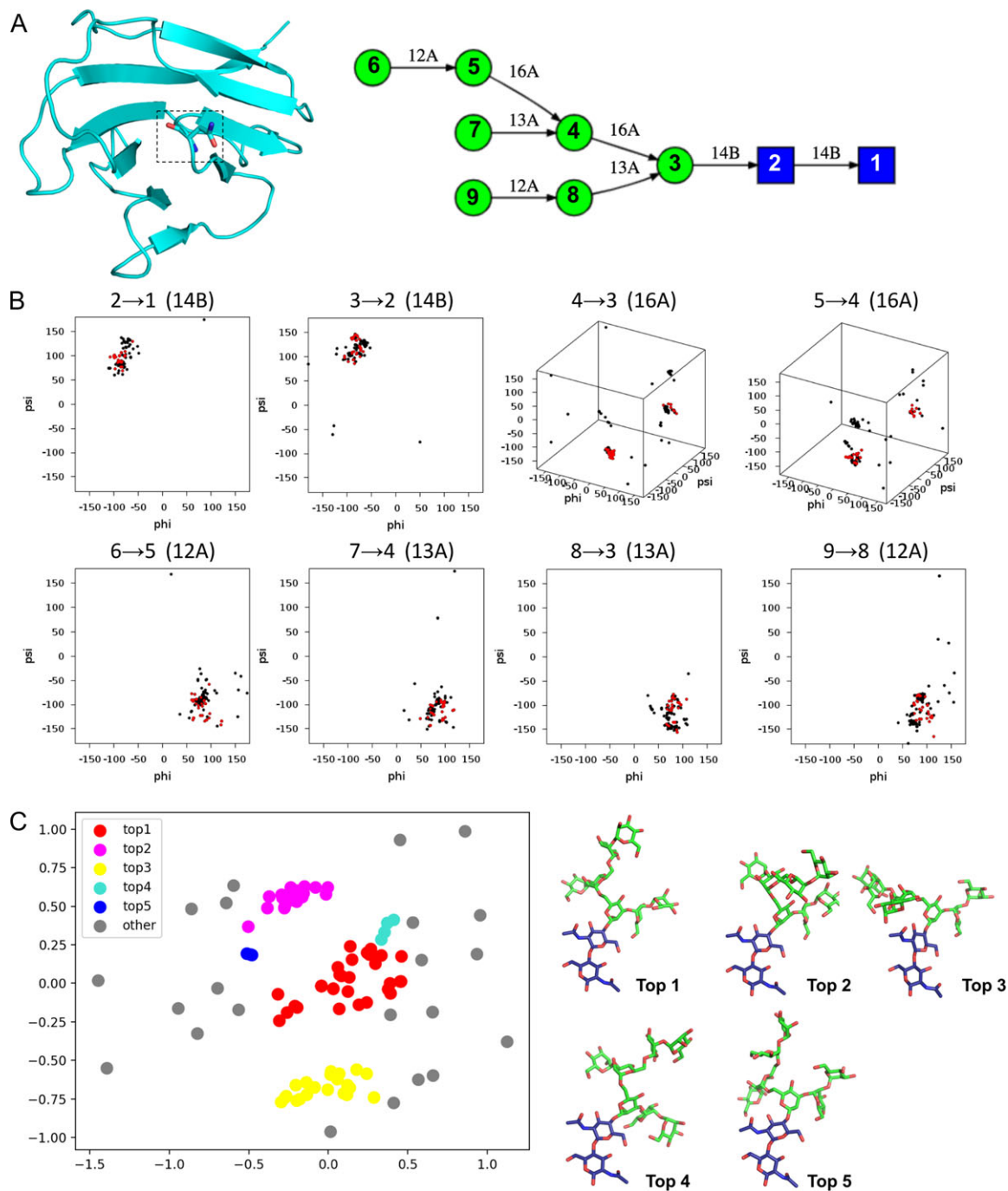


Fig. 5. Illustration of glycan modeling in *Glycan Modeler*. (A) An example of a query glycan sequence and uploaded protein structure (PDB ID: 1CDB). Target glycosylation site (stick) and query glycan sequence were obtained from the identical protein with a glycan structure, PDB ID:1GYA (human CD2). (B) GFDB database search for glycan fragments that contain the query glycan sequence. For the glycan sequence in (A), 112 glycan fragments were identified. Torsion angles of all glycosidic linkages (ϕ , ψ , and ω) are used for the cluster determination. Red dots belong to the largest cluster based on a clustering analysis that is briefly described in Method. (C) Visualization of top five clusters of glycan fragments through a multi-dimensional scaling algorithm in a scikit-learn python module (Pedregosa et al. 2011).

on the internal coordinate (IC) information in the CHARMM carbohydrate force field (Guvench et al. 2009).

Glycan conformation sampling algorithm

Glycan Modeler aims to build glycan models onto glycosylation sites of a target protein within a reasonable computational time without structural errors and bad contacts between glycan models and the protein. To achieve this goal, *Glycan Modeler* conducts conformational sampling by rigid-body rotations of each glycan chain and evaluates the protein–glycan interaction energy using the CHARMM force field.

Figure S3 shows the overall procedure of the conformation sampling. (1) The protein structure is fixed except glycosylated Asn residue(s). The all missing atoms in the protein structures are generated using the IC information in the CHARMM force field and remains to be flexible during the orientation search. (2) The glycan structure from the first GFDB cluster is attached to each glycosylation site (or onto pre-existing glycans) and is restrained by BESTFIT harmonic restraints (to maintain the overall glycan structure) and improper restraints (to maintain proper chirality of each sugar). (3) 500 iterations of orientation search are performed for each glycan chain to find an initial orientation. The orientation search is accomplished by changing torsional angles (ϕ and ψ angles) of a rotatable glycosidic linkage; the selection of a rotatable glycosidic linkage is elaborated

below. (4) The number of bad contacts between the glycan and the protein is counted using a heavy atom distance cutoff of 2.5 Å. If there are less than five bad contacts, a short energy-minimization (10 steepest descent steps) is performed to further avoid bad contacts. Otherwise, the orientation trial is rejected and proceeds to the next iteration. Note that, to reduce the computational time, a bad contact checkup is performed before the minimization. (5) After the energy minimization in (4), if the glycan–protein interaction energy is lower than the previous interaction energy, the glycan orientation is accepted, and IC information of the accepted glycan orientation is stored. (6) After the initial orientation search in (3)–(5), another iteration of orientation search is performed to find lower energy orientation with the same methods described in (4) and (5). In any step, if the glycan–protein interaction energy is lower than a cutoff value (60 kcal/mol per glycan chain, which was empirically determined in this study), the orientation search is terminated and *Glycan Modeler* provides the accepted structure. Otherwise, the procedure goes to (6). The maximum number of cycles for a given glycosylation site is 500. If the procedure fails to find a glycan orientation that satisfies the given criteria, the procedure goes to (6) with the glycan structure from the second GFDB cluster. If the procedure fails to find a glycan orientation that satisfies the given criteria even with all five glycan cluster structures, the minimum-energy glycan orientation among all the searched glycan orientations is provided. (7) The accepted (or minimum-energy) glycan structure(s) in all glycosylation sites are

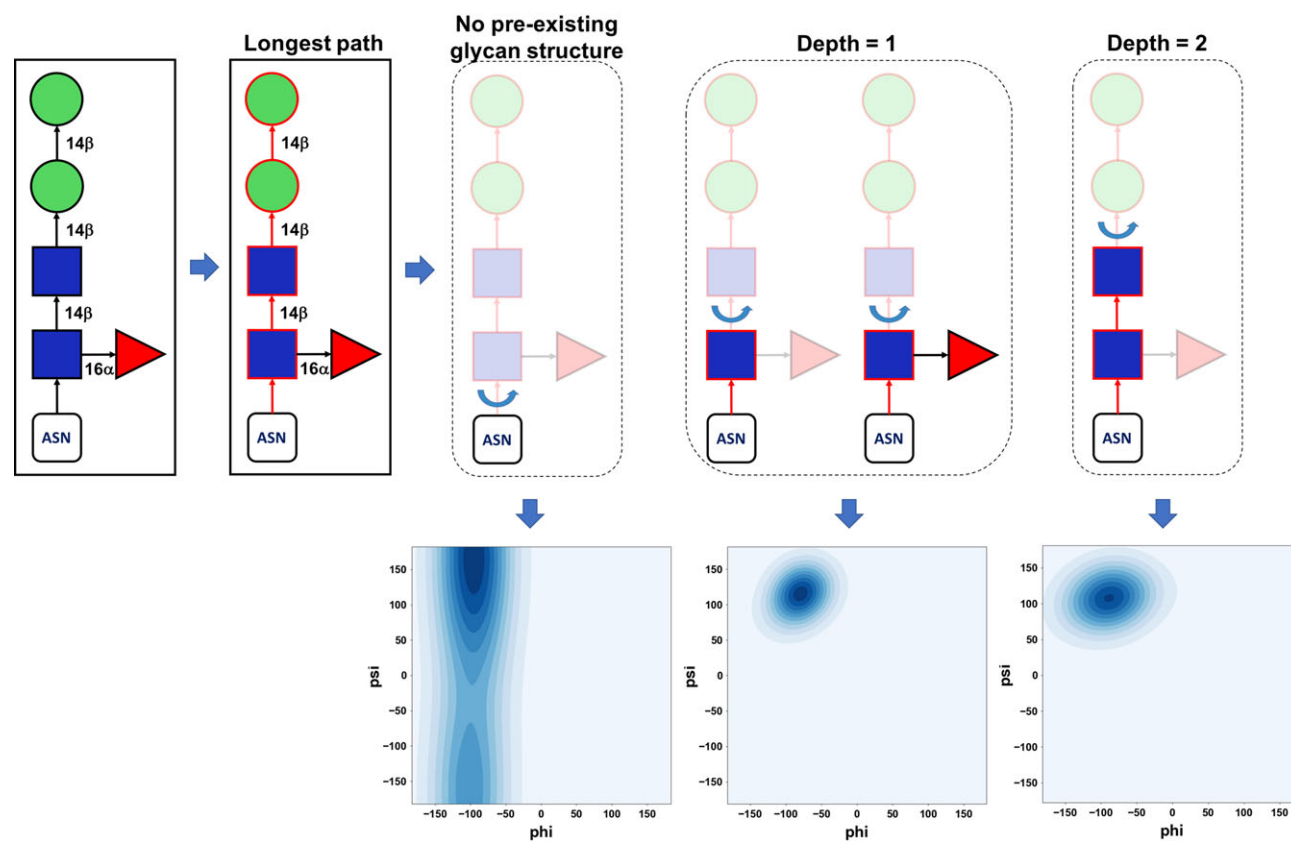


Fig. 6. Rotatable glycosidic linkage. Graphical illustration of rotatable glycosidic linkage determination depending on the extent of a pre-existing glycan structure in a protein. After the longest path on the glycan sequence at each glycosylation site is identified (red), a rotatable glycosidic linkage is defined as a linkage between residues with and without coordinates on the longest path. If there are no coordinates only for the last glycans (i.e., the depth more than two in this case), they are generated simply by IC information in the CHARMM force field and has no rotation during the modeling step. Note that all glycosidic linkages in the pre-existing glycan structure are fixed. Random glycosidic torsion angles are generated based on glycosidic torsion angle populations from GFDB by using the kernel density estimation algorithm in a scikit-learn python module (Pedregosa et al. 2011).

further minimized only with the improper dihedral restraints. In this way, inherent protein–glycan or glycan–glycan bad contacts due to the GFDB average structure are removed without BESTFIT restraints.

In (3) and (6), *Glycan Modeler* only changes ϕ and ψ torsion angles of a selected glycosidic linkage (even for 1–6 linkages) with preservation of other glycosidic linkages to maintain the glycan structures from GFDB. Figure 6 illustrates our selection scheme using an example of a query glycan sequence, Man β 4Man β 4GlcNAc β 4(Fuc α 6)GlcNAc β -Asn. When the query glycan has branches, the longest path on the glycan sequence is first determined (red in Figure 6) and then a rotatable glycosidic linkage is defined as a linkage between the residues with and without coordinates on the longest path. For the glycosylation site with no pre-existing glycan structure, the glycosidic linkage to protein (e.g., CG–ND2–C1–O5 (ϕ) and CB–CG–ND2–C1 (ψ) for N-linked glycan) is selected for torsion angle variations. In the case where the glycosylation site contains a pre-existing glycan structure, the rotatable linkage is defined as the one between the existing glycan and model glycan, as shown in Figure 6. If the depth is greater than two in this exemplary case, *Glycan Modeler* generates the structure only using IC information in the CHARMM carbohydrate force field. Note that all atomic coordinates in the pre-existing glycan structures are conserved. For each (3) and (6) step, the torsion angles are

randomly selected within the ϕ/ψ distribution in the PDB glycan structures of the selected glycosidic linkage type (Figure 6). We use a kernel density estimation algorithm (Pedregosa et al. 2011) to fit the PDB ϕ/ψ distribution and randomly generate the torsional values within the distribution.

In some cases where a protein structure does not contain a pre-existing glycan for a certain glycosylation site, the glycosylation site can be buried inside the protein and thus does not have proper side chain orientation for glycan modeling. To handle such cases, *Glycan Modeler* randomly changes the χ angle of the glycosylation site. For an N-linked glycan, the χ_2 (CA–CB–CG–OD1) angle is changed, while the χ_1 (N–CA–CB–OG/OG1) angle is changed for O-linked glycans.

Web implementation in CHARMM-GUI: glycoprotein modeling

Glycan Modeler is integrated into *Glycan Reader* in CHARMM-GUI (*Glycan Reader & Modeler*). *Glycan Reader & Modeler* displays the detected carbohydrate chains in a PDB entry or in an uploaded PDB file and allows users to select the glycan chains of interest (Park et al. 2017). The selected glycans are listed and their sequences are shown in CASPER sequence representation (Lundborg and Widmalm 2011)

A

Glycosylation / Glycan Ligand(s):

CARA n-linked bDGlcNAc(1→2)aDMan(1→6)[aDMan(1→3)]bDMan(1→4)bDGlcNAc(1→4)[bLFuc(1→6)]bDGlcNAc(1→)PROA-297

Pre-existing glycan chain

- CARB n-linked bDGlc(1→)PROA-276

B

Glycosylation / Glycan Ligand(s)

bDGlc(1→)PROA-276

Glycan Sequence:

Protein: PROA ASN 276

1 β D-glucose Chemical modification:

Sequence Graph:

Next Step:

C

Glycosylation / Glycan Ligand(s)

1 BGLCNA
2 - 14B: BGLCNA
3 - - 14B: BMAN
4 - - - 16A: AMAN
5 - - - 12B: BGLCNA
6 - - - 13A: AMAN
7 - 16B: BFUC

bDGlcNAc(1→2)aDMan(1→6)[aDMan(1→3)]bDMan(1→4)bDGlcNAc(1→4)[bLFuc(1→6)]bDGlcNAc(1→)PROA-276

Glycan Sequence:

Protein: PROA ASN 276

1 β N-acetyl-D-glucosamine Chemical modification:

2 4 \leftarrow β N-acetyl-D-glucosamine Chemical modification:

3 4 \leftarrow β D-mannose Chemical modification:

4 6 \leftarrow α D-mannose Chemical modification:

5 2 \leftarrow β N-acetyl-D-glucosamine Chemical modification:

6 3 \leftarrow α D-mannose Chemical modification:

7 6 \leftarrow β L-fucose Chemical modification:

D

Glycosylation / Glycan Ligand(s):

CARA n-linked bDGlcNAc(1→2)aDMan(1→6)[aDMan(1→3)]bDMan(1→4)bDGlcNAc(1→4)[bLFuc(1→6)]bDGlcNAc(1→)PROA-297

- CARB n-linked bDGlcNAc(1→2)aDMan(1→6)[aDMan(1→3)]bDMan(1→4)bDGlcNAc(1→4)[bLFuc(1→6)]bDGlcNAc(1→)PROA-276

Fig. 7. Snapshots of *Glycan Reader & Modeler*. (A) The glycans that exist in a PDB entry or in an uploaded PDB file are listed and their sequences are shown in CASPER sequence representation (Lundborg and Widmalm 2011) under the Glycosylation/Glycan Ligand(s) category on the PDB manipulation page. In silico glycosylation can be applied using the “Add Glycosylation” button. After clicking this, a new glycan chain is listed at the bottom of existing glycan structures. (B) A new popup window is displayed after clicking the “edit” button, so users can build their glycan model by selection of the glycosylation site, as well as additions, deletions, and changes of sugar residues. (C) Users can also use GRS (*Glycan Reader* sequence) format for a glycan sequence. (D) The CARB glycan chain in (A) is then updated based on the input GRS format in (C).

under the “Glycosylation/Glycan Ligand(s)” category on the PDB manipulation page (Figure 7A). As described in our previous work (Park et al. 2017), an edit button next to each glycan chain enables users to edit a specific glycan chain on the web interface. Specifically, CHARMM-GUI displays a new popup window that allows users to add, remove, or change sugar types and anomeric states, glycosidic linkages, glycosylation types, and chemical modifications (Figure 7B). The new features of *Glycan Reader & Modeler* are (1) availability for in silico glycosylation on any Asn/Ser/Thr residues on the protein (with or without pre-existing glycans), (2) glycan modeling using GRS (*Glycan Reader* sequence) format (Figure 7C), and (3) addition of a sugar at the reducing end of glycan chain. For example, when users click the “Add Glycosylation” button, a new glycan chain is listed at the bottom of the Glycosylation/Glycan Ligand(s) category (Figure 7A). On the new popup window displayed after clicking the “edit” button, users can specify its sequence and target glycosylation site (Figure 7B) or GRS format (Figure 7C) can be used to specify a glycan sequence. The selected glycan is updated under the Glycosylation/Glycan Ligand(s) category (Figure 7D).

Web implementation in CHARMM-GUI: carbohydrate-only structure modeling

The carbohydrate-only solution system can be generated by selecting “Glycan Only System” option in the main page of *Glycan Reader & Modeler*. In the next page, a new web interface is provided, allowing users to build a glycan sequence of their interest. Using the sequence information provided by users, *Glycan Reader & Modeler* generates CHARMM input files to model the glycan structure, and the glycan model structure is generated using the IC information in the CHARMM carbohydrate force field. The generated glycan model is then solvated with ions (upon user’s selection) following the standard building procedures in CHARMM-GUI *Solution Builder*. The glycan system, topology and parameter files, simulation inputs for various MD simulation programs (based upon user’s selection) (Lee et al. 2016), as well as all the CHARMM input files used to generate the glycan system are provided as a downloadable single tar file (“download.tgz”).

Video demos of *Glycan Reader & Modeler*

In CHARMM-GUI Video Demo (<http://www.charmm-gui.org/demo>), we created the following four *Glycan Reader & Modeler* demos: (1) solution system building of the HIV envelop protein using PDB:5FYL (Stewart-Jones et al. 2016), (2) modeling *N*-glycans on a Fc heterodimer of immunoglobulin G1 using PDB:5TPS (Wei et al. 2017), (3) membrane system building of a cholera toxin B and ganglioside GM1 complex using PDB:3CHB (Merritt et al. 1998), and (4) solution system building of a heparin molecule. These examples are further elaborated in Results and discussion.

Preparation of non-redundant *N*-glycan structure set from PDB

PDB structures having resolution better than 3 Å were obtained from the RCSB Protein Data Bank (as of December 2017). *Glycan Reader* (Jo et al. 2011) was used to identify PDB entries with at least one *N*-glycan chain, resulting in 11,844 PDB entries having 28,364 *N*-linked glycan structures in total. All the *N*-glycan structures were categorized based on the glycan sequences to prepare the unique glycan sequences. For each unique glycan sequence having more than 5

PDB entries, we randomly selected one PDB entry as a target. The unique glycan sequences with glycan length smaller than four were discarded to avoid small glycans. Consequently, the benchmark set was composed of 82 target *N*-glycans and parent glycoproteins. Each target glycoprotein has one specific target glycan to measure the performance, although target glycoproteins can have multiple glycans.

MD simulations

MD simulations of a fully glycosylated HIV envelope trimer (from PDB:5FYL or using *Glycan Modeler*) were performed using the CHARMM36m force field (Huang et al. 2017) and NAMD 2.12 (Phillips et al. 2005) with a time step of 2 fs. Proteins were solvated in TIP3P cubic boxes (~160 Å in each side) with 150 mM KCl, yielding 375,305 atoms (a system based on PDB:5FYL) and 380,451 atoms (a system based on *Glycan Modeler*). The non-bonded interactions were smoothly switched off at 10–12 Å with a force-switching function (Steinbach and Brooks 1994). Long-range electrostatic forces were calculated with the particle mesh Ewald algorithm (Essmann et al. 1995). Temperature was maintained at 303.15 K using Langevin dynamics with a friction coefficient of 1 ps⁻¹. Pressure was controlled at 1 bar using the Langevin-piston method with a piston period of 50 fs and a piston decay of 25 fs (Feller et al. 1995). After equilibration with NAMD, the MD systems were transferred to Anton2 (Shaw et al. 2014) for 2-μs production runs with a time step of 2 fs. The non-bonded interactions were cut off at 10 Å. Long-range electrostatic interactions were calculated with the u-series approach (Shaw et al. 2014). Pressure and temperature were controlled at 1 bar and 303.15 K using the semi-isotropic MTK barostats and Nosé–Hoover thermostats under the Multigrator framework (Lippert et al. 2013). Trajectory analysis was performed with CHARMM (Brooks et al. 2009) and VMD (Humphrey et al. 1996). Trajectories for all 2 μs simulation were used to calculate the backbone RMSD and torsion angle distributions.

Supplementary data

Supplementary data is available at *Glycobiology* online.

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Conflict of interest statement

None declared.

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