Research Article

Predicted 3D Model of the Rabies Virus Glycoprotein Trimer

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Received 8 December 2015; Revised 21 February 2016; Accepted 6 March 2016

Academic Editor: Serdar Kuyucak

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The RABVG ectodomain is a homotrimer, and trimers are often called spikes. They are responsible for the attachment of the virus through the interaction with nicotinic acetylcholine receptors, neural cell adhesion molecule (NCAM), and the p75 neurotrophin receptor (p75NTR). This makes them relevant in viral pathogenesis. The antigenic structure differs significantly between the trimers and monomers. Surfaces rich in hydrophobic amino acids are important for trimer stabilization in which the C-terminal of the ectodomain plays an important role; to understand these interactions between the G proteins, a mechanistic study of their function was performed with a molecular model of G protein in its trimeric form. This verified its 3D conformation. The molecular modeling of G protein was performed by a I-TASSER server and was evaluated via a Rachamandran plot and ERRAT program obtained 84.64% and 89.9% of the residues in the favorable regions and overall quality factor, respectively. The molecular dynamics simulations were carried out on RABVG trimer at 310 K. From these theoretical studies, we retrieved the RMSD values from Cα atoms to assess stability. Preliminary model of G protein of rabies virus stable at 12 ns with molecular dynamics was obtained.

1. Introduction

Rabies is a 100% fatal disease caused by the rabies virus (RABV) that affects the central nervous system [1]. Rabies virus belongs to the order Mononegavirales, classified in the Rhabdoviridae family, which includes at least three genera Lyssavirus, Ephemerovirus, and Vesiculovirus. The genus Lyssavirus includes rabies virus. The viral genome consists in a single and negative-stranded nonsegmented RNA, which encodes five proteins: nucleoprotein, matrix protein, phosphoprotein, glycoprotein, and the viral-dependent RNA polymerase [2, 3].

The glycoprotein (RABVG) rabies virus is comprised of four domains: signal peptide (SP), ectodomain (ED), transmembrane (TM), and a cytoplasmic domain (CD) [4, 5].

The RABVG is 65 kDa and has 524 amino acids. This is due to the presence of its signal peptide (SP) that is located on the N-terminal. It spans 19 residues. The SP is responsible for anchoring the protein to the ER-Golgi Apparatus (AP) membrane. This promotes subsequent transport of the nascent protein to the membrane before it is cleaved from the N-terminus in the AP [6, 7].

The RABVG in each peak is anchored in the plasma membrane and the lipid envelope by the transmembrane domain of 22 amino acids from 439 to 461 residues [8]. The C-terminal with the final 44 amino acids is the cytoplasmic domain. It extends into the cytoplasm of the infected cell where it interacts with M to complete the viral assembly [9].

On the other hand, the RABVG ectodomain is a homotrimer that contains a transmembrane domain. Each
2. Material and Methods

2.1. Sequence Analysis, Modeling, and Stereochemical Analysis. This study was designed to predict the 3D model of RABV protein under an iterative threading assembly refinement algorithm implemented in I-TASSER [19]. This was performed because the experimental 3D structure of RABV protein was not available at Protein Data Bank (PDB) (http://www.rcsb.org). Various physical and chemical parameters of primary structure analysis were computed using the ProtParam online tool [20]. The secondary structure of the protein was computed using J-PRED servers [21]. The DIANNA tool [22] was used to check the system classification and disulfide connectivity. This knowledge can be helpful in understanding the secondary structure of the protein because the disulfide bond bridges are important in protein fold stabilization. The transmembrane topology of the RABV was checked using TMHMM [23], MEMSAT3, and MEMSAT-SVM [24].

Finally, the 3D model of RABV was generated using the I-TASSER online server [25]. This generated 3D models along with their confidence score (C-score). After generating the 3D model, structure and stereochemical analysis were performed using different evaluations and validation tools. The Psi/Phi Ramachandran plot was obtained using PROCHECK [26]. This assisted in the evaluation of backbone conformation. The Ramachandran plot was used to check the non-Gly residues in the disallowed regions. Structural quality of the model was assured using Z-scores, which indicate the overall model quality and confirm that the predicted structure is within the range of scores as found in the native proteins. The ProSA web tool [27] was used to determine the Z-scores. Furthermore, the generated model was submitted in the protein model database (PMDB) (https://bioinformatics.cineca.it/PMDB/) with PMDB identifier PM0079619.

With the monomer structure in hand, we attempted to make the trimer interact with protein-protein docking studies to predict the protein complex formed in a protein-protein interaction. These docking studies used the ClusPro server [28–31] that is the first fully automated web-based program for docking proteins. It was one of the top performers at CAPRI (Critical Assessment of Predicted Interactions) rounds 1–12—a community-wide experiment devoted to protein docking [28].

We used the PDBeSum Generate server [32] to understand the interactions between—and assembly of—the five subunits of CRP. This server helps us analyze the interfaces between the subunits and summarizes the interactions across any selected interface. This server also provides information about the residues that actually interact across the interface.

2.2. Molecular Dynamics Simulations. These MD simulations employed NAMD 2.6 (Nanoscale Molecular Dynamics) [33] by applying the CHARMM27 force field for lipids and proteins [34]. This first neutralized the model RABVG trimer with 24 sodium ions along with the TIP3P model for the water box containing 58,566 waters molecules. Structural energy minimization was done using 10,000 steps. Multiple time-stepping algorithms were used with an integration time step of 2 fs. Various interactions were computed in 1, 2, and 4 time steps for covalent bonds, as well as short-range nonbonded interactions and long-range electrostatic forces, respectively. For every ten time steps, the nonbonded interactions had a pair list distance of 13.5 Å. The Van-der-Waals and electrostatic interactions were defined as interactions between short-range nonbonded interactions between particles within 12 Å. A smoothing function was employed for 10 Å Van-der-Waals interactions. Simulations were performed on the equilibrated system for 80 ns under constant pressure and a temperature of 1 atm and 310 K, respectively. The structure with the least energy and converged root mean square deviation was used for subsequent exercises. The final structure was analyzed with CARMA [35] and visualized with the VMD [36] program using 100 frames.

3. Results

3.1. Structural Description of the RABVG 3D Model. This study was initiated to perform structure-based sequence analysis studies on RABV. The protein sequence was retrieved using accession AGN94258.1 from the NCBI protein. The primary structure analysis showed that the RABVG protein had a molecular weight of 58487.3 Daltons and a theoretical isoelectric point (pI) of 7.83. The instability index (II) is computed to be 38.05. This classifies the protein as stable. The negative grand average of hydropathicity (GRAVY) shows a value of −0.173 indicating that the protein was hydrophilic according to other reports [37].

Sequence and secondary structure analyses of RABVG revealed that it has 6 β-sheets, 7 beta hairpins, 3 beta bulges, 21 strands, 5 α-helices, 10 helix-helix interac, 38 beta turns, and 10 γ-turns. Secondary structural features are shown in Figure 1. Disulfide bonds predicted by DiANNA are shown in Table 1.

Disulfide connectivity was predicted to be within 1–8, 2–7, 3–12, 5–11, 6–10, 9–19, 13–17, 14–18, and 15-16. The
**Figure 1:** Predicted secondary structure of RABVG using the J-prep server.

**Figure 2:** A transmembrane motif is revealed along with a 23-amino acid signal peptide at the extracellular N-terminus.

TMHMM, MEMSAT3, and MEMSAT-SVM programs identify a transmembrane region between amino acids 460 and 480. The C-terminal is possibly located in the cytoplasm. The N-terminal—including 19 amino acids from the signaling peptide—is located in the extracellular region (Figure 2).

Knowing the 3D structure of RABVG is very important to understanding the proteins interactions, functions, and their important site localization. The model was not obtained by homology because the identities were low level (23%) with 2J6J and 2CMZ crystals; for closely related
protein sequences with identity higher than 40%, the alignment is almost always correct. Regions of low local sequence similarity become common when the overall sequence identity is below 40% [38, 39]. In this sense, the 3D structure of RABVG was predicted using the I-TASSER (http://zhanglab.ccmb.med.umich.edu/I-TASSER/) online server and the best predicted structure with the maximum confidence score (C-score = −2.18) was selected (Figure 3 of glycoprotein. The top lateral domain (D1) contains about 90 residues in two segments (1 to 17 and 312 to 383). Trimerization domain (DII) is made of three segments (18 to 35, 259 to 311, and 384 to 409), PH domain (DIII) is inserted within domain II. It is made of two segments (36 to 50 and 181 to 258) and has the fold of a pH domain. Fusion (DIV) (51 to 180) is inserted in a loop of the pH domain and is made of an extended sheet structure at the tip of which two loops are located that constitute the membrane-interacting motif of the G ectodomain. (a) Glycoprotein monomeric divided into 4 domains: D1 (red) is lateral domain, DII (blue) is trimerization domain, DIII (orange) is domain of pH, and DIV (yellow) is fusion domain. (b) Surface representation of glycoprotein monomeric colored by domains. (c) Top view of the trimer, colored by domain, shows the formation of 6 alpha helices (blue) which contribute to the stability of the structure. (d) Top view of the trimer surface representation. (e) Glycoprotein trimer (divided into different domains) does not show a significant change in the organization of the domains. (f) Surface representation of the glycoprotein trimer, showing the cavity inside the molecule.
monomer) to achieve protein-protein docking studies using the Cluspro (http://cluspro.bu.edu/) server to get a trimer complex for visualizing the protein interactions (Figure 3).

The central trimerization domain has a significant contribution to the structural stability of the G protein trimer, due to the formation of 6 alpha helices. The different conformations (monomer, dimer, and trimer) of G protein are pH-dependent, several acidic residues (Glu286, Glu293, Glu294, Glu295, Glu405, and Glu408) are brought close together in the postfusion bundle of six helices, suggesting that the acidic residues play molecular switches role in their deprotonated forms, and this should destabilize the central six-helix bundle and thus allow the unfolding of G back toward its prefusion conformation similar to VSV G protein [40].

These residues ensure a correct activation of G and show that the stability of the 6 chain helices has to be tightly regulated since both its destabilization and over-stabilization are detrimental to the virus.

The glycoprotein of rhabdovirus is the target of neutralizing antibodies; its antigenicity and antigenic sites of G have been extensively studied in VSV and RABV [41]. The antigenic site II of RABVG is located between positions 34 and 42 and positions 198 and 200. These peptides are probably linked by a disulfide bridge and held together in the tertiary structure of G antigenic site III which extends from amino acids 330 to 338. This site is associated with the virulence [42].

They have described various regions of the protein G which have an important role in membrane fusion for the internalization of the virus; the region between amino acids 118 and 139 was generally considered to represent an internal fusion peptide for VSV G. However, other studies demonstrated that amino acids 395–418 have a significant influence on fusion, and additional studies identified region 145–164, termed p2-like peptide, as being a pivotal domain in facilitating glycoprotein G-mediated membrane fusion [43]. The stability and the preservation of these areas are important in the structure of G protein for viral internalization.

The structural organization of G is very similar to that of VSV G. This similarity extends from the N-terminal part to at least the end of helix G of domain II. It includes both the PH domain and the fusion domain (109 residues of the fusion domain), as well as the trimerization domain, and reveals a clear structural homology between the two proteins.

G has an altogether different structural organization from those of both class I and class II viral fusion proteins described so far. The polypeptide chain of G folds into four distinct domains (Figure 3): a lateral domain rich in β sheet at the top of the molecule (domain I), a central, mostly α-helical domain that is involved in the trimerization of the top of the molecule (domain II), a neck domain that has the characteristic fold of pleckstrin homology (PH) domains (domain III), and the elongated fusion domain that makes the trimeric stem of the molecule (domain IV). The C-terminal part of G corresponds to AA 411 to 422 in VSV and 410 to 455 in RAVBG [44].

The top lateral domain I contains about 90 residues in two segments (I to 17 and 310 to 383 for VSV and 1 to 17 and 311 to 383 for RABV). Domain II is made of three segments (18 to 35, 259 to 309, and 384 to 405 for VSV and 18–35, 269 to 310, and 384 to 409 for RABV). Domain III is inserted within domain II. It is made of two segments (36 to 50 and 181 to 258 for VSV and RABV) and has the fold of a PH domain. Domain IV (51 to 180 for VSV and RABV) is inserted in a loop of the PH domain [44].

These residues ensure a correct activation of G and show that the stability of the 6 chain helices is tightly regulated because both its destabilization and over-stabilization are detrimental to the virus.

The number of residues involved in residue-residue interactions and a model of the interactions between each monomer denominated A-B, B-C, and C-A are shown in Figure 4. The model depicted in Figure 4 shows that the hydrogen bonds and other nonbonded interactions are responsible for the interactions among the monomers to build the trimer complex.

This trimer model shows the number of interactions across two interfaces as well as details of the individual residue-residue interactions across these interfaces (Figure 5).

The interactions analysis in the 3D structure was obtained by a PDBeSum server; for these hydrogen bonds and other nonbonded, disulfide bridges of CYS between two side chains or the formation of an amide bond (–CO–NH–) between side chains of Lys and a dicarboxylic aminoacid (Glu or Asp) were considered. The 3D structure presents electrostatic interactions between Glu 293 in chain A with Lys 297 in chain B, Glu 300 of chain A with Lys 313 of chain C, Lys 297 of chain A with Glu 300 of chain C, Lys 148 on chain B with Glu 291 on chain C, and Lys298 on 25 chain C with Glu 286 and 293 on chain B. These stabilize the structure (Figure 6).

Quality and reliability of the RABVG structure were checked using Z-score and Ramachandran plot. The stereochemical quality of the RABVG 3D structure was checked with a Ramachandran plot by analyzing the backbone dihedral angles residue by residue. The result showed that 89.9% of the residues were in the favorable region (Figure 7).

The overall model quality can be checked with ProsA Z-score that is used to check whether the input structure is within the range of scores typically found for native proteins of similar size [27]. The Z-score of the protein was –5.54. The model reliability was further checked by ERRAT [45] that analyzes the statistics of nonbonded interactions between different atom types and plots the value of the error function versus position of a 9-residue sliding window as calculated by
Salt bridges
Disulphide bonds
Hydrogen bonds
Nonbonded contacts

Figure 4: Schematic diagram showing the interactions between the subunits.

Figure 5: A model showing the number of interactions across the interfaces and the individual residue-residue interactions across the interfaces along with involved residues. Residue colors: positive (H, K, R) (blue); negative (D, E) (red); S, T, N, Q = neutral (green); A, V, I, M = aliphatic (gray); F, Y, W = aromatic (purple); P, G = Pro&Gly (orange); and C = cysteine (yellow). These include interactions between A and B interface, interaction between A and C interface, and interaction between B and C.
Figure 6: Glycoprotein trimer interactions (snapshot 80 ns): chain A (red), chain B (blue), and chain C (green); the interactions are shown between the chains, Glu 293 chain A with Lys 297 chain B, Glu 300 of chain A with Lys 313 of chain C, Lys 297 of chain A with Glu 300 of chain C, Lys 148 chain B with Glu 129 chain C, and Lys 298 chain C with Glu 286 and 293 B chain.

Figure 7: Ramachandran plot showing residues in the most favorable region and disallowed regions (RAMPAGE by Paul de Bakker and Simon Lovell available at http://mordred.bioc.cam.ac.uk/~rapper/rampage.php) [46].

Number of residues in favored region (~98.0% expected): 408 (89.9%)
Number of residues in allowed region (~2.0% expected): 30 (6.6%)
Number of residues in outlier region: 16 (3.5%)
The ERRAT model had good overall quality. (a) Graphic of ERRAT program for A monomer. (b) Graphic of ERRAT program for B monomer. (c) Graphic of ERRAT program for the C monomer. * On the error axis, two lines are drawn to indicate the confidence with which it is possible to reject regions that exceed that error value. ** Expressed as the percentage of the protein for which the calculated error value falls below the 95% rejection limit. Good high resolution structures generally produce values around 95% or higher. For lower resolutions (2.5 to 3 Å) the average overall quality factor is around 91% [45].

The ERRAT results showed 84.648 overall model quality (Figure 8). The Z-scores, Ramachandran plot, and ERRAT results confirmed that the quality of the RABVG trimer model is suitable for future theoretical studies.

Molecular dynamics (MD) simulations were carried on RABVG trimer at 310 K. From these theoretical studies we retrieved the RMSD values from Cα atoms. This suggests that the system reached structural stability and simulation integrity. The magnitude of the RMSD (7 Å) indicates that the RABVG trimer is stabilized at 12 ns and remains constant until the end of the simulation (Figure 9).

On the other hand, the residues we used in the root mean square fluctuations (RMSF) identify the regions responsible for the fluctuations during the MD simulations. The areas with higher fluctuations correspond to beta-loop-beta chains residues and turn regions. Those with lower fluctuations are regions of α-helices. During the MD simulation, no structural uncoiling was observed. At 310 K, the RMSF values go from 3 to 15 Å obtained from 12000 to 7500 frames; RMSF are similar for all chains except Ala87 to Thr100, Pro136 to Thr147, Phe173 to Asn201, and Ser422 to Val435 residues of chain C (Figure 9).

The regions with the most fluctuations have not been described as important areas to maintain the structural stability of the G protein trimer. This does not generate significant changes within the main trimer binding regions. In the residues 125 to 131 conformational change of chain C after 10 ns was detected and is stable from 40 ns to 80 ns when molecular dynamic end (Figure 10).

During molecular dynamic simulations, the structure of the G protein remained stable after 12 ns. It maintained the trimer bound and conserved key interactions to maintain the stability of the structure.
4. Conclusions

The molecular modeling of G protein was performed by a I-TASSER server and was evaluated via a Rachamandran plot and ERRAT program obtained 84.64% and 89.9% of the residues in the favorable regions and overall quality factor, respectively.

The interactions between residues, 274 to 293, are directly linked to the structure of the prefusion and postfusion of Glycoprotein. These interactions are important to maintain these structures.

This is important for structural stability of the G protein trimer. It might be a good target for antiviral compounds because such modifications would change the helical conformation and be detrimental to the virus.

The fluctuations that occurred during the molecular dynamics do not affect the stability of the structure of G protein trimer. Protein G structural stability was obtained by molecular dynamics analysis at 12 ns.

Competing Interests

The authors declare that they have no competing interests.

Acknowledgments

The authors would like to thank COFAA-IPN.

References

[1] G. M. Baer, “Rabies—an historical perspective,” Infectious Agents and Disease, vol. 3, no. 4, pp. 168–180, 1994.
[2] A. A. V. Albertini, R. W. H. Ruigrok, and D. Blondel, “Rabies virus transcription and replication,” Advances in Virus Research, vol. 79, pp. 1–22, 2011.
[3] A. A. V. Albertini, G. Schoehn, W. Weissenhorn, and R. W. H. Ruigrok, “Structural aspects of rabies virus replication,” Cellular and Molecular Life Sciences, vol. 65, no. 2, pp. 282–294, 2008.
[4] M. Rustici, L. Bracci, L. Lozzi et al., “A model of the rabies virus glycoprotein active site,” Biopolymers, vol. 33, no. 6, pp. 961–969, 1993.
[5] Y. Gaudin, R. W. H. Ruigrok, C. Tuffereau, M. Knossow, and A. Flamand, “Rabies virus glycoprotein is a trimer,” Virology, vol. 187, no. 2, pp. 627–632, 1992.
[6] Y. Gaudin, S. Moreira, J. Bénéjean, D. Blondel, A. Flamand, and C. Tuffereau, “Soluble ectodomain of rabies virus glycoprotein expressed in eukaryotic cells folds in a monomeric conformation that is antigenically distinct from the native state of the complete, membrane-anchored glycoprotein,” The Journal of General Virology, vol. 80, no. 7, pp. 1647–1656, 1999.
[7] L. Sissoëff, M. Mousli, P. England, and C. Tuffereau, “Stable trimerization of recombinant rabies virus glycoprotein
ectodomain is required for interaction with the p75NTR receptor.” *The Journal of General Virology*, vol. 86, no. 9, pp. 2543–2552, 2005.

[8] A. Anilionis, W. H. Wunner, and P. J. Curtis, “Structure of the glycoprotein gene in rabies virus,” *Nature*, vol. 294, no. 5838, pp. 275–278, 1981.

[9] Y. Gaudin, “Rabies virus-induced membrane fusion pathway,” *The Journal of Cell Biology*, vol. 150, no. 3, pp. 601–612, 2000.

[10] T. L. Lentz, T. G. Burrage, A. L. Smith, and G. H. Tignor, “The acetylcholine receptor as a cellular receptor for rabies virus,” *The Yale Journal of Biology and Medicine*, vol. 56, no. 4, pp. 315–322, 1983.

[11] T. L. Lentz, E. Hawrot, D. Donnelly-Roberts, and P. T. Wilson, “Synthetic peptides in the study of the interaction of rabies virus and the acetylcholine receptor,” *Advances in Biochemical Psychopharmacology*, vol. 44, pp. 57–71, 1988.

[12] C. Tuffereau, J. Bénéjean, D. Blondel, B. Kieffer, and A. Flamand, “Low-affinity nerve-growth factor receptor (P75NTR) can serve as a receptor for rabies virus,” *The EMBO Journal*, vol. 17, no. 24, pp. 7250–7259, 1998.

[13] M. Lafon, “Rabies virus receptors,” *Journal of NeuroVirology*, vol. 11, no. 1, pp. 82–87, 2005.

[14] K. Hotta, Y. Motoi, A. Okutani et al., “Role of GPI-anchored NCAM-120 in rabies virus infection,” *Microbes and Infection / Institut Pasteur*, vol. 9, no. 2, pp. 167–174, 2007.

[15] C. Tuffereau, K. Schmidt, C. Langevin, F. Lafay, G. Dechant, and M. Koltzenburg, “The rabies virus glycoprotein receptor p75NTR is not essential for rabies virus infection,” *Journal of Virology*, vol. 81, no. 4, pp. 13622–13630, 2007.

[16] Y. Gaudin, R. W. H. Ruigrok, M. Knossow, and A. Flamand, “Low-pH conformational changes of rabies virus glycoprotein and their role in membrane fusion,” *Journal of Virology*, vol. 67, no. 3, pp. 1365–1372, 1993.

[17] A. Maillard, M. Domanski, P. Brunet, A. Chauffotte, E. Guittet, and Y. Gaudin, “Spectroscopic characterization of two peptides derived from the stem of rabies virus glycoprotein,” *Virus Research*, vol. 93, no. 2, pp. 151–158, 2003.

[18] A. P. Maillard and Y. Gaudin, “Rabies virus glycoprotein can fold in two alternative, antigenically distinct conformations depending on membrane-anchor type,” *The Journal of General Virology*, vol. 83, no. 6, pp. 1465–1476, 2002.

[19] J. Yang, R. Yan, A. Roy, D. Xu, J. Poisson, and Y. Zhang, “The I-TASSER suite: protein structure and function prediction,” *Nature Methods*, vol. 12, no. 1, pp. 7–8, 2015.

[20] M. R. Wilkins, E. Gasteiger, A. Bairoch et al., “Protein identification and analysis tools in the ExPASy server,” *Methods in Molecular Biology*, vol. 112, pp. 531–552, 1999.

[21] C. Cole, J. D. Barber, and G. J. Barton, “The Ipred 3 secondary structure prediction server,” *Nucleic Acids Research*, vol. 36, pp. W197–W201, 2008.

[22] F. Ferré and P. Clote, “DiANNA: a web server for disulfide connectivity prediction,” *Nucleic Acids Research*, vol. 33, no. 2, pp. W230–W232, 2005.

[23] A. Krogh, B. Larsson, G. von Heijne, and E. L. L. Sonnhammer, “Predicting transmembrane protein topology with a hidden Markov model: application to complete genomes,” *Journal of Molecular Biology*, vol. 305, no. 3, pp. 567–580, 2001.

[24] T. Nugent and D. T. Jones, “Transmembrane protein topology prediction using support vector machines,” *BMC Bioinformatics*, vol. 10, article i59, 2009.

[25] A. Roy, A. Kucukural, and Y. Zhang, “I-TASSER: a unified platform for automated protein structure and function prediction,” *Nature Protocols*, vol. 5, no. 4, pp. 725–738, 2010.

[26] R. A. Laskowski, J. A. C. Rullmann, M. W. MacArthur, R. Kaptein, and J. M. Thornton, “AQUA and PROCHECK: programs for checking the quality of protein structures solved by NMR,” *Journal of Biomolecular NMR*, vol. 8, no. 4, pp. 477–486, 1996.

[27] M. Wiederstein and M. J. Sippl, “ProSA-web: interactive web service for the recognition of errors in three-dimensional structures of proteins,” *Nucleic Acids Research*, vol. 35, no. 2, pp. W407–W410, 2007.

[28] D. Kozakov, D. R. Hall, D. Beglov et al., “Achieving reliability and high accuracy in automated protein docking: ClusPro, PIPER, SDU, and stability analysis in CAPRI rounds 13-19,” *Proteins*, vol. 78, no. 15, pp. 3124–3130, 2010.

[29] S. R. Comeau, D. Kozakov, R. Brenché, Y. Shen, D. Beglov, and S. Vajda, “ClusPro: performance in CAPRI rounds 6-11 and the new server,” *Proteins: Structure, Function and Genetics*, vol. 69, no. 4, pp. 781–785, 2007.

[30] S. R. Comeau, D. W. Gatchell, S. Vajda, and C. J. Camacho, “ClusPro: a fully automated algorithm for protein-protein docking,” *Nucleic Acids Research*, vol. 32, pp. W96–W99, 2004.

[31] S. R. Comeau, D. W. Gatchell, S. Vajda, and C. J. Camacho, “ClusPro: an automated docking and discrimination method for the prediction of protein complexes,” *Bioinformatics*, vol. 20, no. 1, pp. 45–50, 2004.

[32] R. A. Laskowski, “PDBeSum new things,” *Nucleic Acids Research*, vol. 37, no. 1, pp. D355–D359, 2009.

[33] J. C. Phillips, R. Braun, W. Wang et al., “Scalable molecular dynamics with NAMD,” *Journal of Computational Chemistry*, vol. 26, no. 16, pp. 1781–1802, 2005.

[34] A. D. MacKerell Jr., D. Bashford, M. Bellott et al., “All-atom empirical potential for molecular modeling and dynamics studies of proteins,” *The Journal of Physical Chemistry B*, vol. 102, no. 18, pp. 3586–3616, 1998.

[35] N. M. Glykos, “Software news and updates. Carma: a molecular dynamics analysis program,” *Journal of Computational Chemistry*, vol. 27, no. 14, pp. 1765–1768, 2006.

[36] J. Hsin, A. Arkhipov, Y. Yin, J. E. Stone, and K. Schulten, “Using VMD: an introductory tutorial,” *Current Protocols in Bioinformatics*, vol. 24, unit 5.7, pp. 5.7.1–5.7.48, 2008.

[37] P. Smialowski, A. J. Martin-Galiano, A. Mikolajka, T. Girschick, T. A. Holak, and D. Frishman, “Protein solubility: sequence based prediction and experimental verification,” *Bioinformatics*, vol. 23, no. 19, pp. 2536–2542, 2007.

[38] M. Saxena, S. S. Bhunia, and A. K. Saxena, “Docking studies of novel pyrazinopyridoindoles class of antihistamines with the homology modelled H(1)-receptor,” *SAR and QSAR in Environmental Research*, vol. 23, no. 3-4, pp. 311–325, 2012.

[39] I. R. Ramachandran, W. Song, N. Lapteva et al., “The phosphatase SRC homology region 2 domain-containing phosphatase-1 is an intrinsic central regulator of dendritic cell function,” *The Journal of Immunology*, vol. 186, no. 7, pp. 3934–3945, 2011.

[40] A. Ferlin, H. Raux, E. Baquero, J. Lepaut, and Y. Gaudin, “Characterization of pH-sensitive molecular switches that trigger the structural transition of vesicular stomatitis virus glycoprotein from the postfusion state toward the prefusion state,” *Journal of Virology*, vol. 88, no. 22, pp. 13396–13409, 2014.
[41] A. A. Albertini, C. Mérigoux, S. Libersou et al., “Characterization of monomeric intermediates during VSV glycoprotein structural transition,” PLoS Pathogens, vol. 8, no. 2, Article ID e1002556, 2012.

[42] Y. Gaudin, “Reversibility in fusion protein conformational changes. The intriguing case of rhabdovirus-induced membrane fusion,” Sub-Cellular Biochemistry, vol. 34, pp. 379–408, 2000.

[43] X. Sun, S. Belouzard, and G. R. Whittaker, “Molecular architecture of the bipartite fusion loops of vesicular stomatitis virus glycoprotein G, a class III viral fusion protein,” The Journal of Biological Chemistry, vol. 283, no. 10, pp. 6418–6427, 2008.

[44] S. Roche, S. Bressanelli, F. A. Rey, and Y. Gaudin, “Crystal structure of the low-pH form of the vesicular stomatitis virus glycoprotein G,” Science, vol. 313, no. 5784, pp. 187–191, 2006.

[45] C. Colovos and T. O. Yeates, “Verification of protein structures: patterns of nonbonded atomic interactions,” Protein Science, vol. 2, no. 9, pp. 1511–1519, 1993.

[46] S. C. Lovell, I. W. Davis, W. B. Arendall III et al., “Structure validation by Cα geometry: φ/ψ and Cβ deviation,” Proteins: Structure, Function & Genetics, vol. 50, no. 3, pp. 437–450, 2002.