Performance of CHARMM36m with modified water model in simulating intrinsically disordered proteins: a case study

Laura I. Gil Pineda1,2, Laurie N. Milko1, Yi He1✉

1 Department of Chemistry & Chemical Biology, The University of New Mexico, Albuquerque, NM 87131, USA
2 Department of Chemical Sciences, Universidad Icesi, Cali 760031, Colombia

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Abstract Molecular dynamics simulations can be a powerful tool to complement experiments in the study of the structures and dynamics of intrinsically disordered proteins. Though the accuracy of the physics-based all-atom force fields has improved significantly in simulating structured proteins over the past twenty years, most of these force fields face a big challenge to simulate flexible proteins. Recently, CHARMM36m with modified TIP3P model was proposed as a possible solution to simulate intrinsically disordered proteins. Here, we tested the proposed solution using an extensively studied protein, namely NCBD, to explore the performance of CHARMM36m plus modified TIP3P water. Our results suggest that the modified TIP3P water model does enhance the sampling of conformational space compared to the standard TIP3P water model. However, the new CHARMM36m force field still leads to over-compact structures and over-stabilized helices.

Keywords Nuclear coactivator binding domain (NCBD), CHARMM36m, Protein simulations, Secondary structure preferences, Stabilization of proteins

INTRODUCTION

In traditional structural biology, a globular protein has a single stable tertiary structure. The discovery of intrinsically disordered proteins (IDPs), which do not have a unique stable structure under physiological conditions, is challenging the traditional structural biological paradigm (Click et al. 2010; Dunker et al. 2008; Dyson and Wright 2005; Tompa 2002; Wright and Dyson 1999). One unique property of IDP sequences is that the primary sequence of an IDP is enriched with polar and charged amino acids, along with decreased amounts of non-polar residues. Such decreases of non-polar residues have limited the capability of IDPs to form hydrophobic cores, which are the key contributors leading to stable structures in structured proteins (Dunker et al. 2001; Huang and MacKerell 2018). This distinct sequence composition enables IDP’s ability to switch between or sample different tertiary structural states. Their constant structural fluctuation allows a single IDP to perform a multitude of biological functions (Uversky et al. 2005), such as roles in cellular signaling (Smock and Gierasch 2009) and regulation (Fuxreiter et al. 2008; Babu et al. 2011). IDPs have also been associated with several pathological conditions, including cancer (Iakoucheva et al. 2002; Metallo 2010; Uversky et al. 2008) and neurodegenerative diseases (Uversky et al. 2014).

Experimental techniques, such as small-angle X-ray scattering (SAXS), nuclear magnetic resonance (NMR) and Forster resonance energy transfer (FRET) spectroscopy, are often being used to study IDPs (Eliezer 2009; Sapienza and Lee 2010; Yengo and Berger 2010). The major challenges are that the heterogeneous ensembles of IDPs and their rapid inter-conversion between conformations make it difficult to obtain detailed structural information solely from experiments. Over the past years, molecular dynamics (MD)
Simulate intrinsically disordered proteins using CHARMM36m with modified water model

RESULTS

Preference to compact structures

As mentioned in the Introduction section, a major problem when modeling IDPs through physics-based atomistic models is overly compact ensembles (Best et al. 2014; Henriques et al. 2015; Piana et al. 2014; Rauscher et al. 2015). Previous simulations of NCBD with different force fields faced this problem (Burger et al. 2012; Knott and Best 2012; Papaleo et al. 2018). The modified TIP3P water model intends to fix this problem by increasing the dispersion interactions between the protein and water (Huang et al. 2017). In the timescale studied, the modified water system sampled more "open" conformations when compared to the standard TIP3P water (Fig. 1).
However, when calculating the average \( R_g \) for each system, the difference is reasonably small (1.21 nm for modified vs 1.2 nm for standard). Details of the average value and the standard deviation of each trajectory are shown in Table 1. It also results much more compact when compared to that obtained by previous simulations (1.37–1.49 nm) (Knott and Best 2012; Papaleo et al. 2018) and of course lower to the values estimated experimentally under “native-like” conditions (1.52 nm) (Kjaergaard et al. 2010). This could be explained by the fact that previous works employed different techniques to improve their samplings, such as REMD and experimental restraints (Knott and Best 2012; Papaleo et al. 2018). Additionally, it is important to mention that only one Lennard–Jones well depth (\( \epsilon_H \)) value was tested and it was previously stated that no universal \( \epsilon_H \) applies to all IDP systems, so it might be necessary to decrease or increase this value to get the desired effect (Huang et al. 2017).

### Table 1

| System   | Average | Standard deviation |
|----------|---------|--------------------|
| Modified |         |                    |
| MD1      | 12.200  | 1.047              |
| MD2      | 12.090  | 0.951              |
| Standard |         |                    |
| MD1      | 11.971  | 0.669              |
| MD2      | 11.947  | 0.640              |

**Over-stabilization of helical structures**

Another problem that is common when modeling IDPs is the over-stabilization and preference bias to secondary structures, in particular, helices. To evaluate this effect in CHARMM36m, the secondary structure (specifically the helicity) was calculated for each system, using the define secondary structure of proteins (DSSP) algorithm (Kabsch and Sander 1983). Both systems (Fig. 2) presented three regions of high helical propensity that correspond in sequence location to the helixes presented in unbound NCBD (Kjaergaard et al. 2010). However, previous works on NCBD rarely report helicity above 0.8 (or 80%). This indicates possibly over-stabilization of the helices, which was also observed for NCBD with previous force fields (Naganathan and Orozco 2011; Zhang et al. 2012) even at high temperatures. Additionally, some of the previous studies (Knott and Best 2012; Zhang et al. 2012) found a bimodal behavior in the region corresponding to Helix 2 (residues 23–35), which was not observed here.

Contact maps (Fig. 3) were also constructed. Two residues were considered to be in contact if the distance between two heavy atoms of these residues, which must be more than four residues apart in the sequence, was less than 0.55 nm. Based on this definition, the intra-helical contacts were identified \((i, i + 4)\) presenting a high probability of contact (diagonal), which correlates with the high helicity found using DSSP. Perpendicular to these are the inter-helical contacts, which were also similar for both systems, with minor differences in the probability. The other tertiary contacts appear to be distinctive or at least with a different probability.
between the two systems. Specifically, there were some contacts between the C-terminal and the first helix, as well as with the N-terminal. This could be an indication of a compact structure which is in agreement with the low radius of gyration obtained. These contacts were presented with a slightly higher probability in the standard water system than in the modified ones. Finally, both systems seem to have similar results, presenting the most difference in the C-terminal region of Helix 3. This is because of the low percentage of interactions between the C-terminal of the protein and the rest of NCBD as shown in Fig. 3.

Modified TIP3P samples larger conformational space

Next, the conformational space sampled by each system was studied. First, protein structures of both systems were clustered with a cutoff of 0.25 nm. This clustering was done by using the core (no N or C-terminal) for the least-squares fit and RMSD calculations. By excluding the terminal movements (which can be significant in terms of RMSD values), conformational changes in helical packing can be identified. With this cutoff, the simulations with the modified TIP3P water generated about twice more clusters than the ones using the standard TIP3P water. Additionally, the modified system, in general, had clusters that represented smaller amounts of conformations (low percentage). These could be indications that the modified system sampled a more heterogeneous free energy space. The top five clusters (Fig. 4), which on average represent ~65% of the structures, consisted of folded structures (high residual structure) and some appear to be more compact than the initial structure. All the top structures present three highly structured helixes, as expected from DSSP analysis. However, there are some changes in the packing of the helixes, which can be seen clearly in Clusters 2 and 4 of the modified TIP3P water systems and to a less extent in Clusters 4 and 5 of the standard TIP3P systems.
Fig. 4 Representative structures for the top five clusters, fitted to the initial structure, for each system. Below each structure is the percentage it represents and its radius of gyration.

Fig. 5 Free energy landscapes for each system, using as order parameters: radius of gyration ($R_g$) and fraction of native contacts ($Q$). Notice that both have multiple minimums but the modified system samples a wider space.
Having studied representative structures from each system, free energy landscapes based on these trajectories were generated to examine the sampling capabilities of molecular dynamics simulations using different water models. Given that the modifications in the water model were implemented to try to replicate chain dimensions and the ability to model an intrinsically disordered protein is being tested, the order parameters chosen were the radius of gyration ($R_g$) and the fraction of native contacts ($Q$). As can be seen in Fig. 5, the modified TIP3P water system samples a larger conformational space. The standard water system is mostly limited by its inability to sample higher radius of gyration, causing repeated sampling of the same space, many times with $R_g$ values lower than the initial structure (~1.45 nm). While the modified system also has its minimums at low $R_g$ values, its sampling is more evenly spread with no significant energy barriers. Additionally, it is also clear that the modified TIP3P system samples more conformations with $Q < 0.5$, which although may not imply the sampling of unfolded structures, does indicate the sampling of structurally different conformations compared to the initial/NMR structure.

**DISCUSSION**

The purpose of this study is to evaluate the proposed solution of using the latest CHARMM36m with a modified TIP3P water model to simulate IDPs as well as determine whether the proposed modification in the protein–water interactions is enough to improve chain dimensions in MD simulations. The modification of the water model allows NCBD to sample more open conformations (and larger conformational space), based on the larger value of the calculated average radius of gyration. It is clear that, for the tested $\varepsilon$ value and the timescale reached, such modification is not enough to replicate the chain dimensions obtained experimentally. Higher $\varepsilon$ values may need to be tested, although the physical validation of this is uncertain. Additionally, it may be necessary to perform REMD simulations, as limitations set by the starting structure and general problems with convergence may contribute to these results. As far as the CHARMM36m force field, with or without the water model modification, there seems to be some weakness in the force field. To the extent of this study, there is a potential overemphasis on secondary structure, over-stabilization of the protein in general, and the possible underestimation of protein–water interactions.

**METHOD**

**All-atom simulation details**

NCBD (59 residues) was simulated using two water models: the standard TIP3P and the modified TIP3P water as described by Huang et al. (2017). The only difference between these two water models is that the parameter describing $\varepsilon_H$ between the water hydrogen atoms is changed from $-0.046$ kcal/mol in the standard CHARMM TIP3P water model to $-0.1$ kcal/mol in the modified CHARMM TIP3P water model (Huang et al. 2017). The structure for NCBD was obtained from the ligand-free state solution NMR structure (PDB ID: 2KKJ) (Kjaergaard et al. 2010) as shown in Fig. 6. All simulations were carried out using the CHARMM36m force field with explicit solvents and the Groningen Machine for Chemical Simulations (GROMACS) package (version 2018.3) (Abraham et al. 2015; Berendsen et al. 1995; Pall et al. 2015). The protein was placed in a cubic box with the corresponding water model and counter ions (Cl–) to neutralize the whole system at 304 K. Long-range electrostatics is calculated using the particle-mesh Ewald (PME) algorithm (Darden et al. 1993; Essmann et al. 1995). Periodic boundary conditions were applied in all directions. Each system of protein, water, and counter ions was prepared using CHARMM-GUI (Jo et al. 2008; Lee et al. 2016), which generates a series of GROMACS inputs for subsequent MD simulations.
To generate equilibrated starting structures for the MD simulations, steepest-descent minimization was carried out, followed by a 1-ns MD equilibrium simulation with a time step of 1 fs, to heat the whole system from 1 K to the desired temperature. All bonds with hydrogen atoms are converted to constraints with the algorithm LINear Constraint Solver (LINCS) (Hess et al. 1997), using the default parameters of the GROMACS package. The equilibrated structures obtained from the above steps were used for subsequent production runs. A Nose–HOOVER temperature thermostat (Nose 1984; Hoover 1985) was used to maintain the temperature. The time step was 2 fs, and snapshots were taken every 100 ps. For both the standard TIP3P and the modified TIP3P water model systems, a cubic water box size of 10 nm was employed and run for a total of 20 μs, including two 10-μs long MD trajectories.

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Compliance with Ethical Standards

Conflict of interest Laura I. Gil Pineda, Laurie N. Milko, and Yi He declare that they have no conflicts of interest.

Human and animal rights and informed consent This article does not contain any studies with human or animal subjects performed by any of the authors.

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**RESEARCH ARTICLE**

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