Supplemental material: Adaptive resolution simulation of an atomistic protein in MARTINI water

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In the following we present the remainder of results of our adaptive resolution simulations. The simulation protocol is given in the original paper1. Figs. 1 and 2 depict the normalized density profiles of water bundles and oxygen atoms, respectively, around the protein’s center of mass. The water density profiles are the same for all simulations. This is important because the stability of the protein structure depends critically on the density of water molecules around the protein.

FIG. 1. Solvent density around the center of mass of protein G for AdResS simulations with atomistic region radius sizes of 3.2nm (top), 3.4nm (middle) and 3.6nm (bottom) using the bundled water model 22 (The results for the model 12 are presented in the original paper1). The plots include error bars3. The results are compared to the fully atomistic bundled and SPC solvations. The vertical lines denote the boundaries between resolution domains, i.e., the atomistic (AT), hybrid (HY), and the coarse-grained (CG) domains.

FIG. 2. Oxygen density profiles around the center of mass of protein G. The presentation is the same as in Fig. 1 except that the results for the both models of bundled water are presented.

To achieve a flat density profile throughout the hybrid region, the adaptive resolution simulations (AdResS) require the use of thermodynamic (TD) force4,5. The latter compensates for the difference in the chemical potential at different levels of resolution. We calculate the TD force with an iterative procedure as described in the literature on a water system without the presence of protein. The TD force that acts on bundle’s center of mass in the hybrid region is shown for both models in Fig. 3.

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FIG. 3. The thermodynamics force that acts on molecule’s center of mass in the hybrid region.

FIG. 4. RMSD (top) and RMSF with error bars (bottom) of the backbone atoms with respect to the crystal structure as a function of time. We compare the results obtained from the fully atomistic simulation using SPC and bundled water (model 2) to AdResS simulations with three atomistic region sizes: spheres of radii 3.2, 3.4 and 3.6 nm, respectively. The corresponding results using the model 1 of the bundled water are presented in the original paper. The protein’s structural properties are verified with the root-mean-square deviation (RMSD) and the root-mean-square fluctuations (RMSF) of the backbone atoms with respect to the crystal structure (see Fig. 4). Using the model 2 of the bundled water the average RMSD values (in nm) are 0.23 ± 0.02, 0.17 ± 0.02, 0.18 ± 0.02, 0.17 ± 0.03 for the all-atom bundled and AdResS bundled (with the three atomistic region sizes) solvation, respectively. These values confirm that the protein structure is stable during our simulations.

We have also computed the protein’s radius of gyration ($R_g$) shown in Fig. 5. The results from the adaptive resolution simulations closely match the atomistic counterparts. Even for the adaptive resolution simulation with the smallest atomistic region the protein remains compact during the total simulation run. With atomistic domain size almost three times larger than $R_g$ we have ensured that the whole protein is always located within the atomistic region.

FIG. 5. The radius of gyration as a function of time. The presentation is the same as in Fig. 4 except that the results for the both models of bundled water are presented.

FIG. 6. Percentage of native contacts as a function of time. The presentation is the same as in Fig. 5.

Furthermore, we have determined the stability of native contacts, i.e., contacts between the side chains of two amino acids that are more than five residues apart in the protein but are spatially closer than 0.75 nm, by the MaxCluster tool. Figure 6 shows time evolution of the percentage of native contacts, i.e., the ratio between numbers of native contacts in the current and the crystal structure. The percentage remains con-
stant during the course of all simulations, further confirming the stable structure of the protein.

The solvent accessible surface area and the secondary structure were obtained using the DSSP program and are shown on Figs. 7 and 8.

The solvent accessible area results are in accordance with the relation between solvent accessible area $A$ and the protein’s molar mass $M_r$, i.e., $A = \alpha M_r^{0.73}$, where $\alpha = 0.063\text{nm}^2/\text{mol}$. For protein G the relation yields approximately 37 nm$^2$. We observe minor differences in the time evolution of secondary structure elements between different simulations. However, these differences appear to be uncorrelated with a specific simulation approach.

Finally, we have carried out additional simulations of protein enclosed in atomistic water droplet in vacuum. The results are shown in Figs. 9, 10 and 11. Simulations of pro-
Protein within water droplet in vacuum give similar results as the bulk AdResS and fully atomistic simulations for all considered structural properties of the protein. Such results indicate that protein’s effect on the water structure does not exceed the chosen sizes of atomistic regions and water droplets. The bulk is modeled in the AdResS simulations by CG water that is not able to form hydrogen bonds. Therefore, the stability of the protein does not seem to be affected strongly by the bulk hydrogen bond network. Note, however, that even in the smallest water droplet there are still several hydration shells around the protein.