

quickly the optimal setup for their calculations, and (if necessary) extend its functionality with an extremely small effort, unprecedented in the codes currently available. Also, the module is highly autonomous from the other NAMD source files, and can be easily adapted to other simulation programs as well. The set of features and their options will be introduced. Applications using the methods implemented so far (umbrella sampling, steered MD, adaptive biasing force and metadynamics) and make specific use of their combined advantages, will also be presented.

2094-Pos Board B64

Generating Pathways for Free Energy Calculations in Proteins Using Constraint-Based Conformational Sampling

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Constraint-based sampling [1] is a computational method for quickly exploring the allowed motions of a protein. Sampling of protein conformations is guided by a set of geometric constraints instead of a molecular mechanics force field. The geometric constraints preserve covalent bonding geometry, maintain favorable non-bonded contacts, and prevent steric overlap. We have tuned the constraints so that sampled conformations are low in energy according to a molecular mechanics force field (Amber). In this work, we apply the constraint-based sampling method in a targeted fashion to generate a pathway between two conformational end states in the protein dihydrofolate reductase (DHFR). The pathway we generate bridges the so-called “closed” and “occluded” states of DHFR, a transition that involves loop rearrangement near the binding site and relative rotations of subdomains. We then use this pathway as a starting point for free energy calculations. By performing molecular dynamics umbrella sampling [2] along the pathway, we obtain the free energy difference between the end states. Although the generated pathway is not necessarily the actual transition pathway, accurate calculation of the free energy difference between end states only requires that the pathway be low in free energy in the umbrella sampling method.

[1] Wells S, Menor S, Hespeneide B M, and Thorpe M F. Constrained geometric simulation of the diffusive motions in proteins. *Phys Bio* 2 S127-S136 (2005).

[2] Mamonova T and Kurnikova M. Structure and energetics of channel-forming protein-polysaccharide complexes inferred via computational statistical thermodynamics. *J Phys Chem* 110(49) 25091-25100 (2006).

2095-Pos Board B65

Conformational Transition Path Sampling For Proteins

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Defining a reaction path (or reaction coordinate) is an essential step to understand chemical reactions, conformational change, or ligand-binding processes in proteins, and it is also important to consider protein-protein associations, for example, in immunology. However, conventional molecular dynamics simulation methods often fail to find appropriate reaction paths even with very huge computing facilities. This is because such reactions occur much more slowly than the computationally feasible time, and the sampling efficiency of such reaction paths can be very low especially for large proteins. Recent advance in transition path sampling techniques helps us to circumvent this annoying situation, but the application of such methods to large proteins has been rarely done. Using such transition path sampling methods, we examine the conformational change of a protein, adenylate kinase, after ligand binding. In this work, we propose a novel coarse-grained model for the protein to describe the ligand-binding processes in a realistic way. The purpose of this study is to clarify the conformational transition pathways in the protein, that is, there are two moving domains in the protein, and we try to understand which domain moves first to make a transition from the open to closed structures at finite temperatures. To quantify the result, we calculate the free energy surface along such a reaction path. We compare a zero-temperature path (intrinsic reaction path) and finite-temperature paths, and discuss the difference in terms of conformational entropy and other quantities. We furthermore carry out the transition-path study of the protein using the corresponding all-atom model, and discuss the difference between the coarse-grained and all-atom models.

2096-Pos Board B66

Computing Transitions in Macromolecular Systems: Dynamic Importance Sampling

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Understanding and predicting conformational change in macromolecules is central to linking structure and function. Performing straight-forward all-atom molecular dynamics would, in principle, enable sampling of conforma-

tional changes. However, the time-scale for functionally important transitions, exceeds the usual molecular dynamics timescales by several orders of magnitude. For example, with large amounts of computer time all these transitions could be observed with good statistics and the results collected simply by waiting long enough. Thus to sample on longer time-scales requires the development of biased molecular dynamics methods, where the bias can be applied and corrected for at the end. In our approach, called ‘Dynamic Importance Sampling’ we generate a series of independent trajectories that are conditioned on starting and ending in defined conformations. Trajectories are generated using two different algorithms: one uses a soft-ratcheting scheme based on stochastic trajectories and the other uses information from the set of normal modes. The algorithms, which require no initial pathway, are capable of rapidly determining multiple pathways between known states. The associated probability scores, determined by correcting for the bias, allows us to rank order the most likely pathways. We will present examples from three-helix bundles and other systems for both analysis and possible experimental work.

2097-Pos Board B67

Improving The Computational Efficiency Of Non-Dynamical Approaches For Equilibrium Sampling Of All-Atom Protein Models

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We have been pursuing non-dynamical sampling methods which employ pre-calculation and storage of partial results, but limited energy calls during “production” sampling. Specifically, we have been using polymer-growth strategies to sample implicitly solvated all-atom polypeptides. Our original implementation was not efficient compared to standard Langevin dynamics (LD) simulations. We now describe a variety of technical advances - mostly in implementation, rather than in the algorithm - which have led to unprecedented efficiency compared to LD for several polypeptides. The efficiency comparison was performed using a novel statistical tool developed in our group.

2098-Pos Board B68

Evaluating The Effective Sample Size Of Equilibrium Molecular Simulations Using Automatically Approximated Physical States

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In order to assess “convergence” in molecular simulations and to quantify the efficiency of competing algorithms, we need a reasonable and universally applicable estimate of the “effective sample size,” N_{eff} . For equilibrium sampling, we suggest the most fundamental definition of N_{eff} to be that number governing the variance in populations of physical states measured from multiple independent simulations. We demonstrate a simple automated procedure for approximating physical states and show that the resulting estimates for N_{eff} agree well with intuitive transition counts. A wide variety of biomolecular systems are successfully analyzed. Our approach can be applied to systems with unknown physical states and to modern non-dynamical algorithms, such as those based on the “exchange” mechanism. The necessary software for estimating N_{eff} will be freely available on our website.

2099-Pos Board B69

The “Weighted Ensemble” Path Sampling Method Can Find Target States Blindly And Automatically

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The computational sampling of rare transition events is a well appreciated challenge in chemical and biomolecular systems. Previously, we employed the “weighted ensemble” (WE) approach to path sampling and found it efficient in a simple protein model (PNAS, 104:18043, 2007). However, one drawback of the original WE formulation, and of other path sampling methods, is the requirement for a previously known target state and/or approximate reaction coordinate. We show that an improved, fully “blind” WE method does not require choosing any coordinates in advance. We demonstrate the correctness of the new approach, and quantify its efficiency, using a previously studied united-residue model of calmodulin. In addition, we have performed WE simulations using the CHARMM package to study alanine dipeptide. We find multiple structurally distinct pathways, highlighting the strength of WE in sampling multiple barrier-separated pathways.

2100-Pos Board B70

Accelerated Subspace Iteration Method for Protein Normal Mode Analysis

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Normal mode analysis is commonly employed to elucidate the conformational dynamics of proteins and related biological function. In typical applications, only a small subset of the complete set of frequencies and normal modes of