“Dividing and conquering” and “caching” in molecular modeling

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Abstract

Molecular modeling is widely utilized in subjects including but not limited to physics, chemistry, biology, materials science and engineering. Impressive progress has been made in development of theories, algorithms and software packages. To divide and conquer, and to cache intermediate results have been long standing principles in development of algorithms. Not surprisingly, Most of important methodological advancements in more than half century of molecule modeling are various implementations of these two fundamental principles. To access interesting behavior of complex molecular systems in a wide range of spatial and temporal scales, the molecular modeling community has invested tremendous efforts on two lines of algorithm development. The first is coarse graining, which is to represent multiple basic particles in higher resolution modeling as a single larger and softer particle in lower resolution counterpart, with resulting force fields of partial transferability at the expense of some information loss. The second is enhanced sampling, which realizes “dividing and conquering” and/or “caching” in configurational space with focus either on reaction coordinates and collective variables as in Metadynamics and related algorithms, or on the transition matrix
and state discretization as in Markov state models. For this line of algorithms, spatial resolution is maintained but no transferability is available. With introduction of machine learning techniques, many new developments, particularly those based on deep learning, have been implemented to realize more efficient and accurate ways of “dividing and conquering” and “caching” along these two lines of algorithmic research. We recently developed the generalized solvation free energy theory, which suggests a third class of algorithm that facilitate molecular modeling through partially transferable in resolution “caching” of distributions for local clusters of molecular degrees of freedom. Connections and potential interactions among these three algorithmic directions are discussed. This brief review is on both the traditional development and the application of machine learning in molecular modeling from the perspective of “dividing and conquering” and “caching”, with the hope to stimulate development of more elegant, efficient and reliable formulations and algorithms in this regard.

**Impact of molecular modeling in scientific research**

Impact of molecular modeling in scientific research is clearly embodied by the number of publications. Results of a Web of Science (www.webofknowledge.com) search with various relevant key words is listed in Table 1. However, despite widespread applications, we remain far from accurately predicting and designing molecular systems in general. Further methodological development is highly desired to tap its full potential. Historically, molecular modeling has been approached from a physical or application point of view, and numerous excellent reviews are available in this regard.\textsuperscript{1–8} From an algorithmic perspective, “dividing and conquering” (DC) and “caching” intermediate results that need to be computed repetitively are two fundamental principles in development of many important algorithms (e.g. dynamic programming\textsuperscript{9}). In this review, I provide a brief discussion of important methodological development in molecular modeling as specific applications of these two principles. The content will be organized as the following. Part I describes fundamental challenges in
molecular modeling; Part II summarizes application of these two fundamental algorithmic principles in two lines of methodological research, coarse graining (CG)\textsuperscript{10–15} and enhanced sampling (ES);\textsuperscript{16–19} Part III covers how machine learning, particularly deep learning, facilitate DC and “caching” in CG and ES, part IV introduces a third strategy for partially transferable in resolution “caching” of local sampling based on the generalized solvation free energy theory; and part V discusses connections among these three lines of algorithmic development, their specific advantages and prospective explorations. Due to the large body of literature and limited space, we apologize to authors whose excellent work are not cited here.

Table 1: Number of publications from web of science search on Sep,8, 2020

| Key words                                                      | Number of publications |
|---------------------------------------------------------------|------------------------|
| Molecular dynamics simulation                                 | 241,748                |
| Monte Carlo simulation                                       | 189,550                |
| QM-MM (quantum mechanical - molecular mechanical) simulation  | 9907                   |
| Dissipative particle dynamics simulation                     | 3693                   |
| Langevin dynamics simulation                                 | 3893                   |
| Molecular modeling                                           | 2,072,091              |
| All of the above                                             | 2,243,182              |

Challenges in molecular modeling

Accurate description of molecular interactions

Molecular interactions may be accurately described with high level molecular orbital theories (e.g. CCSD) or sophisticated density functionals combined with large basis sets.\textsuperscript{20} These milestones were awarded Nobel prize in 1998. However, such quantum mechanically detailed computation is prohibitively expensive for any realistic complex molecular systems. Molecular interactions are traditionally represented by explicit functions and pairwise approximations as exemplified by typical physics based atomistic molecular mechanical (MM)
force fields (FF):\(^{21}\)

\[
U(\vec{R}) = \sum_{\text{bonds}} K_b(b - b_0)^2 + \sum_{\text{angles}} K_\theta(\theta - \theta_0)^2
\]

\[
+ \sum_{\text{dihedral}} K_\chi(1 + \cos(n\chi - \delta))
\]

\[
+ \sum_{\text{impropers}} K_{imp}(\phi - \phi_0)^2
\]

\[
+ \sum_{\text{nonbonded}} \left( \epsilon_{ij} \left[ \left( \frac{R_{minij}}{r_{ij}} \right)^{12} - \left( \frac{R_{minij}}{r_{ij}} \right)^6 \right] + \frac{q_i q_j}{\varepsilon r_{ij}} \right)
\]

or knowledge based potential functions:\(^{22}\) These simple functions, while being amenable to rapid computation and are physically sound grounded near local energy minima (e.g. harmonic behavior of bonding, bending near equilibrium bond lengths and bend angles), are problematic for anharmonic interactions, which are very common in many molecular systems.\(^{23}\) It is well understood that properly parameterized Lennard-Jones potentials are accurate only near the bottom of its potential well. Frustration are ubiquitous in biomolecular systems and are likely fundamental driving force for conformational transformations.\(^{24}\) One may imagine that a molecular system with all its comprising particles at their respective “happy” energy minima positions would likely be a stable “dead” molecule, which may be a good structural support but is likely not able to provide dynamic functional behavior.

Pairwise approximations are usually adopted for its computational convenience, both in terms of dramatically reduced computational cost and tremendously smaller (when compared with possible many body potentials) number of parameters that need to be fit in FF parameterization. It is widely acknowledged that construction of traditional FF (e.g. equation\(^1\)) is a laborious and error-prone process. Development of polarizable\(^{25,26}\) and more complex FF with larger parameter sets\(^{27}\) alleviate some shortcomings of earlier counterpart. Expansion based treatments were incorporated to address anharmonicity.\(^{28}\) However, to tackle limitation of explicit simple functional form and pairwise approximation for better description of molecular interactions remain challenges to be met for the molecular modeling community.
Additionally, even atomistic simulations are prohibitively expensive for large biomolecular complexes at long time scales (e.g. milliseconds and beyond).

**Inherent low efficiency in sampling nominal high dimensional space of molecular systems**

Complexity of molecular systems is rooted in their molecular interactions, which engenders complex and non-linear correlations among molecular degrees of freedom (DOFs). Consequently, effective number of DOFs are greatly reduced. Therefore, complex molecular systems are confined to manifolds of much lower dimensionality with near zero measure in corresponding nominal high dimensional space (NHDS), suggesting brute force random sampling is hopeless.

In stochastic trajectory generation by Monte Carlo (MC) simulations or candidate structural model proposal in protein structure prediction and refinement (or other similar scenarios), new configuration proposal are carried out in NHDS, a lot of effort is inevitably wasted due to sampling outside the actual manifold occupied by the target molecular system. Such wasting may be avoided if we understood all correlations. However, understanding all correlations implicates accurate description of global free energy landscape (FEL) and there is no need to investigate it further! Due to preference of lower energy configurations by typical importance sampling strategies (e.g. Metropolis MC), stochastic trajectories tend to be trapped in local minima of FEL, this is especially true for complex molecular (e.g. biomolecular) systems which have hierarchical rugged FEL with many local minima.

In trajectory generation by molecular dynamics (MD) simulations, configurational space is explored by laws of classical mechanics and no wasting due to random moves exists. However, molecular systems may well drift away from their true manifolds due to insufficient accuracy of FF. Similar to stochastic trajectory generation, it takes long simulations to map FEL since molecular system tend to staying at any local minimum, achieve equilibrium among many local minima is just as challenging as in the case of stochastic counterpart.
DC and “caching” in traditional molecular modeling

To cope with fundamental difficulties in molecular modeling, two distinct lines of methodological development (CG and ES) based on DC and “caching” strategies have been conducted and tremendous progress has been made in understanding of molecular systems. A brief summary is given below:

Coarse graining, a partially transferable “caching” strategy

Atomistic molecular dynamics simulations are the most well established coarse graining with a strong theoretical foundation, the Born-Oppenheimer approximation. Theoretically, MMFF are potential of mean force (PMF) obtained by averaging over all possible electronic DOFs for given atomic configurations. In practice, due to the fact that ab initio calculations are expensive and may have significant error when level of theory (and/or basis set) is not sufficient, reference data usually include results from both quantum mechanical (QM) calculations and well-established experimental data. A DC strategy is utilized by selecting atomic clusters of various size to facilitate generation of QM reference data. The essential information learned from reference data is then permanently and approximately “cached” in FF parameters through the parameterization process.

Due to the separation of time scales for electronic and atomic motion, elimination of electronic DOFs is straightforward but comes with the price of incapability in describing chemical reactions. To harvest benefits of both quantum and atomistic simulations, a well-established DC strategy is to treat a small region participating interested chemical reaction at QM detail and the surrounding with atomistic MD simulation. This series of pioneering work was awarded Nobel prize in 2013.

The united atom model (UAM) is the next step in coarse graining, where hydrogen atoms are merged into bonded heavy atoms. This is quite intuitive since hydrogens have much smaller mass on the one hand, and are difficult to see by experimental detection tech-
nologies utilizing electron diffraction (e.g. X-ray crystallography) on the other hand. For both polymeric and biomolecular systems, UAM remains to be expensive for many interested spatial and temporal scales. Therefore, further coarse graining in various forms have been constructed. As a matter of fact, CG is usually used to denote modeling with particles that representing multiple atoms in contrast to atomistic simulations, and the same convention will be adopted in the remaining part of this review. Both “Top-down” (that based on reproducing experimental data) and “bottom-up” (that based on reproducing certain properties of atomistic simulations) approaches are utilized. For polymeric materials, beads are either utilized to represent monomers or defined on consideration of persistent length, and dissipative particle dynamics (DPD) were proposed to deal with complexities arise from much larger particles. For biomolecular systems, a wide variety of coarse grained models have been developed. Another important subjects of CG methodology development is materials science. Earlier definition of CG particles are rather ad hoc. More formulations with improved statistical mechanical rigor appeared later on, with radial distribution function based inversion, entropy divergence and force matching algorithm being outstanding examples of systematic development. Present CG is essentially to realize the following mapping as disclosed by equation (4) in ref.:

\[ \exp[-\beta V_{CG}(R_{CG})] \equiv \int dR \delta(M_{R}(r) - R_{CG}) \exp[-\beta V(r)] \]  

with \( r \) and \( R \) being coordinates in higher resolution and CG coordinates, \( M_{R}(r) \) being the map operator from \( r \) to \( R \), \( V \) and \( V_{CG} \) being potential energy of higher resolution and CG representation respectively. Due to lack of time scale separation for essentially all CG mapping, strict realization of this equation/mapping is not rigorously possible. A naive treatment of CG particles as basic units (with no internal degrees of freedom) would result in wrong thermodynamics. Due to corresponding significant loss of information, it is not possible to develop a definition of CG and corresponding FF parameterization for comprehensive repro-
duction of all atom description of corresponding molecular systems. Different coarse graining have distinct advantages and disadvantages, so choosing proper CG strategy is highly dependent upon specific goal in mind. CG particles are usually isotropic larger and softer particles with pairwise interactions, or simple convex anisotropic object (e.g. soft spheroids) that may be treated analytically.\textsuperscript{15,48,49} Such simplifications provide both convenience of computation and certain deficiency for capturing physics of the target molecular systems. CG may be carried out iteratively to address increasingly larger spatial scales by “caching” lower resolution CG distributions with ultra CG (UCG) FF.\textsuperscript{13,50–52} Pairwise approximation and explicit simple function form remain to be limitations of interaction description for traditional CG FF. When compared with atomistic FF, pairwise approximation deteriorate further due to lack of time scale separation (Fig. 1).
Figure 1: Schematic illustration of time scale separation issue in CG. A) and B) show two situations with $C_\alpha$ distances between two amino acids GLU and ALA being $R$, but with GLU have different conformations. If $C_\alpha$ atoms were defined as CG site, then these two relative conformation with distinct interactions would be treated as the same. In A) and B), CG site distance in both A) and B) are $R$, but many other pairs of atoms have distinct distances as exemplified by $r_1$ and $r_2$. Such treatment would only be true if for any small amount of displacement of $C_\alpha$, side chains accomplished many rotations and thus may be accurately represented by averaging (i.e. with good time scale separation). This is apparently not the case not only for the specific definition of $C_\alpha$ being CG site.

Another simple and powerful type of CG model for biomolecular systems is Go model\textsuperscript{53,54} and elastic network models (ENM)\textsuperscript{55,56} or gaussian network models (GNM)\textsuperscript{57} with native structure being defined as the equilibrium state, and with quadratic/harmonic interactions between all residues within given cutoff. Only a few parameters (e.g. cutoff distance, spring constant) are needed. Such models “caching” the experimental structures and are proved to be useful in understanding major conformational transitions and slow dynamics of many biomolecular systems.\textsuperscript{58,59}
Enhanced sampling, a nontransferable in resolution DC and/or “caching” strategy

Umbrella sampling (US)\textsuperscript{60} is the first combination of DC and “caching” strategy for better sampling of molecular system along a given reaction coordinate (RC) (or order parameter) \(s\). DC strategy is first applied by dividing \(s\) into windows, information for each window is then partially “cached” by a bias potential. Later on, adaptive US (AUS)\textsuperscript{61,62} and weighted histogram analysis method (WHAM)\textsuperscript{63} was developed to improve both efficiency and accuracy. MABR\textsuperscript{64,65} was developed to achieve error bound analysis which is lack in WHAM. Further development including adaptive bias force (ABF)\textsuperscript{66} and metadynamics.\textsuperscript{67} Details of these methodologies were well explained by excellent reviews.\textsuperscript{68–71} The common trick to all of these algorithms (and their variants) is to “cache” visited configurational space with bias potentials/force and thus dramatically accelerate sampling of interested rare events. Denote CV as \(s(r)\) (\(r\) being physical coordinates of atoms/particles in the target molecular system), equilibrium distribution and free energy on the CV may be expressed as:\textsuperscript{18}

\[
p_0(s) = \int dr \delta[s - s(r)] p_0(r) = \langle \delta[s - s(r)] \rangle \\
p_0(r) = \frac{e^{-\beta U(r)}}{\int dr e^{-\beta U(r)}} \\
F(s) = -\frac{1}{\beta} \log[p_0(s)] \\
F(s) = -\frac{1}{\beta} \log[p(s)] - V(s)
\]

with \(p(s)\) being the sampled distribution in simulation with corresponding bias potential \(V(s)\) for “caching” of visited configurational space.

The starting point of these “caching” algorithms is selection of reaction coordinates (RC) or collective variables (CV), which is a very challenging task for complex molecular systems in most cases. Traditionally, principle component analysis (PCA)\textsuperscript{72} is the most widely utilized and a robust way for disclosing DOFs associated with the largest variations. To
deal with ubiquitous nonlinear correlations, kernels are often used albeit with the difficulty of choosing proper kernels.\textsuperscript{73} Additional methodologies, include multidimensional scaling (MDS),\textsuperscript{74} isomap,\textsuperscript{75} locally linear embedding (LLE),\textsuperscript{76} diffusion map\textsuperscript{77,78} and sketch map\textsuperscript{79} have been developed to map out manifold for high dimensional data. However, each has its own limitations. For example, LLE\textsuperscript{76} is sensitive to noise and therefore has difficulty with molecular simulation trajectories which are quite noisy; Isomap\textsuperscript{75} requires relatively homogeneously sampled manifold to be accurate. Both LLE and Isomap do not provide explicit mapping between molecular coordinates and CVs; Diffusion and sketch maps are likely to be more suitable to analyze molecular simulation trajectories. Nonetheless, their successful application for large and complex molecular systems remain to be tested. When we are interested in finding paths for transitions among known metastable states, transition path sampling (TPS)\textsuperscript{80,81} methodology maybe utilized to establish CV.

Apparently, RC and/or CV based ES is a different path for facilitate simulation of complex molecular systems on longer time scales from coarse graining. One apparent plus side is that these algorithms are “in resolution” as no systematic discarding of molecular DOFs occur. With specification of RC and/or CVs, computational resource is directed toward the presumably most interesting dynamics of target molecular system, and RC and/or CV maybe repetitively refined/updated to obtain mechanistic understanding of interested molecular processes. However, the down side is that “cached” information on local configurational space is not transferable to other similar molecular systems. While rigorous transferability may not be easily established for any CG FF, practical utility of CG FF for molecular systems with similar composition and thermodynamic conditions have been quite common and useful.\textsuperscript{15} Therefore, CG FF may be deemed as partially transferable.

An important recent development of DC strategy for enhanced sampling is Markov state models (MSM),\textsuperscript{82–84} one great advantage of which is that no RC or CV is needed. Instead, it extracts long-time dynamics from independent short trajectories distributed in configurational space. Many important biomolecular functional processes have been characterized
with this great technique.\textsuperscript{85,86} The most fundamental assumption is that all states for a target molecular system form an ergodic Markov chain:

\[
\pi(t + \tau) = \pi(t)P
\]  

with \(\pi(t)\) and \(\pi(t + \tau)\) being a vector of probabilities for all states at time \(t\) and \(t + \tau\) respectively. \(P\) is the transition matrix with its element \(P_{ij}\) being probability of the molecular system being found in state \(j\) after an implied lag time \((\tau)\) from the previous state \(i\). Apparently as \(t\) goes to infinity for an equilibrium molecular system, a stationary distribution \(\pi\) will arise as defined below:

\[
\pi = \pi P
\]

The advantage of not needing RC/CV does not come for free but with accompanying difficulties. Firstly, one has to distribute start point of trajectories to statistically important and different part of configurational space, then select proper (usually hierarchical, with each level of hierarchy corresponds to a specific lag time) partition of configurational space into discrete states. This is the key step of DC strategy in MSM. No formal rule is available and experience is important. In many cases some try and error is necessary. Secondly, within each discrete state at a given level of hierarchy, equilibration is assumed to be achieved instantly and this assumption causes systematic discretization error, which fortunately may be controlled with proper partition and sufficiently long lag time.\textsuperscript{87} Apparently, metastable states obtained from MSM analysis is molecular system specific and thus not transferable.

Another important class of enhanced sampling is to facilitate sampling with non-Boltzmann distributions and restore property at targeted thermodynamic condition through proper reweight.\textsuperscript{88} Most outstanding examples are Tsallis statistics,\textsuperscript{89,90} parallel tempering,\textsuperscript{91} replica exchange molecular dynamics,\textsuperscript{92} Landau-Wang algorithm\textsuperscript{93} and integrated tempering sampling (ITS).\textsuperscript{94,95} These algorithms are not direct applications of DC and “caching” strategies and are not discussed here.
Machine learning improves “caching” in molecular modeling

Toward ab initio accuracy of molecular simulation potentials through machine learning

Fixed functional form and pairwise approximation of non-bonded interactions are two major factors limiting the accuracy of molecular interaction descriptions in both atomistic and CG FF. Neural network (NN) has capability of approximate arbitrary functions and therefore has the potential to address these two issues. Not surprisingly, significant progress has been made in this regard as summarized by recent excellent reviews.\textsuperscript{96–102} Cutoff and attention to local interactions remains the DC strategy for development of machine learning potentials. The major improvement over traditional FF is better “caching” that overcomes pairwise approximation and fixed functional form limitations. NN FF naturally tackle both issues as explicit functions are not necessary since NNs are universal approximators that can fit any function. The significance of many-body potentials\textsuperscript{103} and extent of pairwise contributions were analyzed.\textsuperscript{104,105} There are also efforts to search for proper simple functional forms, which are expected to be more accurate than usual functional forms in traditional FF on the one hand, and alleviate overfitting/generalization and computational cost of complex NN FF on the other hand,\textsuperscript{106,107} especially when training dataset is small. While most machine learning FF are trained to predict energy,\textsuperscript{97,101} gradient-domain machine learning (GDML) approach\textsuperscript{108} directly learns from forces and realizes great savings of data generation.

Just as in the case of traditional FF, transferability and accuracy is always a tradeoff. More transferability implicates less attention is paid to “cache” detailed differences among different molecular systems, hence less accuracy. Exploration in this regard, however, remains not as much as necessary.\textsuperscript{109–111} Unlike manual fitting of traditional FF, systematic investigation of tradeoff strategy is feasible for machine learning fitting,\textsuperscript{112} and yet to be done for many
interesting molecular systems. With expediency of NN training, development a hierarchy of NN FF with increasing transferability/accuracy and decreasing accuracy/transferability is likely to become a pleasing reality in the near future.

Rapid further development of machine learning potentials, particularly NN potentials, are expected. However, significant challenges for NN potentials remain on better generalization capability, description/treatment of long range interactions,\textsuperscript{113} wide range of transferability, faster computation and proper characterization of their error bounds. Should significant progress be made on these issues, it is promising that through these potentials we may have routine molecular simulations with both classical efficiency and \textit{ab initio} accuracy in the near future.

**Machine learning and coarse graining**

As in the case of constructing atomic level potentials, machine learning has been applied to address two outstanding pending issues in coarse graining, which are definition of CG sites/particles and parameterization of corresponding interactions between/among these sites/particles.

Traditional CG FF, suffers from both pairwise approximation and accuracy ceiling of simple fixed functional forms which are easy to fit. By using more complex (but fixed functional form) potentials with a machine learning fitting process, Chan \textit{et. al.}\textsuperscript{114} developed ML-BOP CG water model with great success. Deep neural network (DNN) was utilized to facilitate parameterization of CG potentials when given radial distribution functions (RDF) from atomistic simulations.\textsuperscript{115} CGnet demonstrated great success with simple model systems (alanine dipeptide).\textsuperscript{116} DeePCG model was developed to overcome pair approximation and fixed functional form and demonstrated with water.\textsuperscript{117} Using oxygen site to represent water is rather intuitive. However, for more complex biomolecules such as proteins, possibility for selection of CG site explodes. To improve over intuitive or manual try and error definition of CG sites, a number of studies have been carried out\textsuperscript{118–121} and the methods established
provide better and faster options for choosing CG sites. However, no consensus strategy is available up to date and more investigations are desired. The fundamental difficulty is that there is no sufficient time scale separation between explicit CG DOFs and discarded implicit DOFs, regardless of specific selection scheme being utilized. Intuitively, one would expect CG FF parameters to be strongly dependent upon definition of CG sites/particles. In this regard, auto-encoders were utilized to construct a generative framework that accomplishes CG representation and parameterization in a unified way.\textsuperscript{122} The spirit of generative adversarial networks was utilized to facilitate CG construction and parameterization, particularly with virtual site representation.\textsuperscript{123} It was found that description of off-target property by CG exhibit strong correlation with CG resolution, to which on-target property being much less sensitive.\textsuperscript{124} Such observation suggests that adjust CG for specific target properties might be a better strategy than searching for a single best CG representation. Despite potentially more severe impact of pairwise approximation for CG FF than in atomistic FF, quantitative analysis in this regard remain to be done.

**Machine learning in searching for RC/CVs and construction of MSM**

To overcome difficulties of earlier nonlinear CV construction algorithms\textsuperscript{75–77,79} and reduce reliance on human experience, auto-encoders, which is well-established for trainable (non-linear) dimensionality reduction, are utilized in a few studies.\textsuperscript{125–128} Chen and Ferguson\textsuperscript{126} first utilized autoencoders to learn nonlinear CVs that are explicit and differentiable functions of molecular coordinates, thus enabling direct further utility in molecular simulations for more effective exploration of configurational space. Further improvement\textsuperscript{125} was achieved through circular network nodes and hierarchical network architectures to rank-order CVs. Wehmeyer and Noé\textsuperscript{127} developed time-lagged auto-encoder to search for low dimensional embeddings that capture slow dynamics. Ribeiro \textit{et. al.}\textsuperscript{128} proposed the reweighted autoencoded variational Bayes to iteratively refine RC and demonstrated in computation of the binding free energy profile for a hydrophobic ligand-substrate system.
Building a MSM for any specific molecular system requires tremendous experience and many steps in process are error prone. To overcome these pitfalls, VAMPnet that based on variational approach for Markov process was developed to realize the complete mapping steps from molecular trajectories to Markov states.\(^{129}\)

**Partially transferable and in resolution local distribution**

**“caching” with generalized solvation free energy theory**

Both CG and ES methodologies facilitate molecular simulation by effectively reducing local sampling. In CG, it is realized through “caching” (integration) of faster DOFs with proper CG FF, and thus has the inevitable cost of losing resolution (information), accompanied by the desired attribute of (partial) transferability to various extent. ES dramatically reduces lingering time of molecular systems in local minima through “caching” visited local configurational space, which is usually defined by relevant DC strategies, with biasing potentials. When compared with CG, there is no resolution loss. However, “cached” pattern of local configurational space is molecular process specific and thus not transferable at all. In molecular modeling community, these two lines of methodologies are developed quite independently. Nevertheless, one might want to ask why not have both of them in one method, that is to reduce repetitive local sampling without loss of resolution and with “cached” results partially transferable. We explored a first step toward this direction through a neural network implementation of the generalized solvation free energy theory (GSFE).\(^{130}\)

In GSFE theory, each comprising unit in a complex molecular system is solvated by its neighboring units. Therefore, each unit is both a solute itself and a comprising solvent unit of other units. Let \((x_i, y_i) = R_i\) denote a region \(i\) defined by a solute \(x_i\) and its solvent \(y_i\), a molecular system of \(n\) units has \(n\) overlapping regions with corresponding free energy and joint distribution:
Free energy minimization of the molecular system in thermodynamic equilibrium may be treated as maximization of joint probability (eq. 9). The first product term on the right hand side of the equation 10 is the local approximation of the joint distribution, and the fraction term is the contribution from global correlation. While direct calculation of the global correlation is daunting, due to significant overlapping of local regions, self consistence among all \( n \) local regions give a practical approximation for the global contribution term.

The local term may be further expanded:

\[
P(R_i) = P(x_i, y_i) = P(x_i | y_i) P(y_i)
\]  

Both terms may be learned from either experimental or reliable computational datasets. The first term in eq.11 is the likelihood term when \( x_i \) is the given, it quantifies the extent of match between the solute \( x_i \) and its solvent \( y_i \). The second term is an effective local prior term, it quantifies the stability of the solvent environment \( y_i \). Computation of the prior term is more difficult than the likelihood term, but certainly learnable when sufficient data is available. A local maximum likelihood approximation of GSFE is to simply ignore the local prior term.

Through local approximations, free energy of a molecular system is expressed as a function of local likelihood and local priors, both of which may be learned from proper dataset, which are local distributions of corresponding molecular systems. It is important to note here what to be “cached” is a fit for local likelihood or prior, not a fit for the energy or force as in regular machine learning of both atomistic and CG potentials.

A particular implementation of the local maximum likelihood approximation of GSFE
for protein structure refinement with residues defined as comprising unit was conducted. In this scheme, GSFE is integrated with autodifferentiation and coordinate transformation to construct a computational graph for free energy optimization. With fully trainable local FEL (LFEL) derived from backbone and $C_\beta$ atoms, we achieved superb efficiency and competitive accuracy when compared state of the art all atom refinement methodology. Refinement of typical protein structure (within 300 amino acids) takes less than 30 seconds on a single CPU core, in contrast to a few hours by typical efficient sampling based algorithms (e.g. FastRelax) and thousands of hours for MD based refinement. In the latest CASP14 refinement contest (predictioncenter.org/casp14/index.cgi), our method ranked the 12th among 40 participants. We expect addition of complete heavy atom information and improved prior terms to further improve this method in the future. GSFE theory, and the LFEL line of thought, is certainly extendable to modeling of general soft matter.

More on connection among CG, ES and GSFE theory

All of these algorithms have a common goal of reproducing a joint distribution of a given molecular system at some target resolution, albeit from distinct perspectives. The fundamental underpinning is the fact that molecular correlations among its various DOFs limit a molecular system to a manifold of significantly lower dimension. Additionally, complex molecular systems (e.g. biomolecular systems) have hierarchical FEL. The necessary number of DOFs and corresponding dimensionality of manifold decreases accordingly as our interests move to higher hierarchies. ES,CG and GSFE are thus distinct strategies to “cache” correlations from a configurational space (Fig. 2 or physical space perspective Fig. 3).
Figure 2: Schematic illustration of essential features for enhanced sampling by Metadynamics and MSM. A) The “S” shape grey line represent the unknown manifold in the configurational space (represented by the square) of a molecular system. B) Small circles connected by blue arrows represent computed (guessed) RC/CVs for the molecular system, which is utilized to conduct Metadynamics simulations. C) The FEL of the molecular system along the computed/selected RC/CV in B). D) Characterization of the FEL by bias potentials (gaussians represented by blue bell shaped lines) accumulated in the course of Metadynamics simulations. E) Distribution of the molecular system to the whole configurational space at the start of a MSM simulation, small circles represent initial start points for short MSM trajectories. F) Sampling results of short MSM trajectories fall mainly near the manifold, distinct “states” are represented by different colors. G) Establishment of transition matrix by transition counts between “states” obtained from short trajectories.

When viewed from the focal point, Both CG and GSFE are motivated to “cache” relevant information on the full configurational distribution for local clusters of molecular DOFs. Such local clusters of molecular DOFs may be found in many similar molecular systems (e.g. protein molecular systems) and consequently have limited and approximate transferability. In CG, strongly correlated local clusters of molecular DOFs are represented as a single particle, correlations/interactions of CG particles within selected cutoff distances are represented by CG FF and longer range correlations/interactions are either neglected or incorporated through more coarser CG models. In GSFE, all complex many body correlations within selected cutoff (i.e. specific solvent of each solute) is directly “cached” by two terms in
equation 11, local likelihoods and local priors. In contrast, both RC/CV based ES and MSM are designed to first describe local parts of the configurational space formed by all molecular DOFs of the target molecular system. Information for such local configurational space is "cached" either as bias potentials or transition counts, which are further processed to map FEL and dynamics of interested molecular processes. All points (or local parts) of the full configurational space are molecular system specific, thus results are not transferable at all.

Computational process (or educated guess) for establishment of RC/CVs is essentially "caching" results from sampling/guessing of local parts of the configurational space formed by all DOFs within the target molecular system, and extract implicit/explicit mapping function from molecular DOFs to RC/CVs, full description of the later is hoped to disclose our interested molecular processes (e.g. biomolecular conformational transitions, substrate binding/release in catalysis). Trajectories of these molecular processes in configurational space usually correspond to part of the manifold which are made explicit by correctly constructed CVs. Involved molecular DOFs for RC/CVs are not necessarily spatially adjacent on the one hand, and may be different for different molecular processes of the same molecular system. Apparently, CVs are molecular process specific and not transferable, even among different molecular process of the same molecular system. Nonetheless, the methodology for searching CVs may be applied to many different molecular processes/systems.

In contrast, traditional CG focus on local correlations in real/physical space. The first step of CG is to partition atoms/particles of high resolution representation into highly correlated local clusters that will be represented by corresponding single CG particles, and moderately correlated regions define cutoff for CG particles; The second step is to select a site (usually one of the comprising high resolution particles) to represent the corresponding highly correlated cluster; The third step is to select functional forms to describe molecular interactions among newly defined CG particles, and parameters are optimized by selected loss functions (e.g. differences of average force in force matching) in the whole configurational space of molecular systems and hopefully to be transferable to some extent. One may
imagine that both best clustering and optimal representation sites of clusters may vary with
different functional forms used to describe CG particle interactions and in different part of
configurational space. Neural network based CG potentials do not have limitation of fixed
functional form and pairwise approximations. However, the need to partition molecular
systems into transferable clusters and to specify representation site/particle remain.

For all different forms of CG, the fundamental essence is to “cache” many body potential
of mean force (PMF) in CG FF. In contrast, GSFE is to first using a DC strategy to divide
molecular systems into local regions, then directly “cache” many body PMF (or local FEL)
of such local regions. The advantage of CG is a simpler resulting physical model, but is
inflexible due to fixed clustering and representation on the one hand, and loses resolution on
the other hand. Properly implemented GSFE while has a selected spatial region of specific
solvent for each comprising unit of a molecular system, the constitutes of such regions are
fully dynamic and no loss of resolution is involved. Hence all difficulty and uncertainty as-
associated with partition and site selection, which apparently limit transferability, disappears.
Correspondingly, the extent of transferability of a GSFE model is in principle at least no
worse than CG. Differences of CG and GSFE is schematically illustrated in Fig. 3.
Figure 3: Schematic illustration of difference between CG and GSFE using protein as an example. A) Target molecular systems in physical space. Due to the goal of construct partially transferable models and/or force fields, usually many different but similar molecular systems are considered. B) Selection of local atom/particle clusters to be represented as one particle in CG model. C) Selection of CG sites. D) Comparison between atomistic (or higher resolution) simulation results and CG (lower resolution) results. E) Adjust of CG FF parameter according to comparison from D). F) Definition of solvent region for each solute unit. G) Feature extraction for each solute. H) “Caching” of LFEL with neural network by training with prepared data sets.

These strategies may be combined to facilitate molecular modeling. For example, one might first utilize deep learning based near quantum accuracy many body FF to perform atomistic simulations for protein molecular systems, and then extracting local distributions properly with GSFE, which may potentially be utilized to analyze protein molecular systems with near-quantum accuracy and at regular amino-acid based CG speed! Similarly, one may extract and “cache” large body of information from residue level CG simulations with GSFE, which may be utilized to achieve ultra CG (UCG) efficiency with residue resolution. Application of CV and MSM based ES algorithm for CG models is straight forward. Combination of GSFE with CV or MSM based ES is more subtle and yet to be investigated.
Conclusions and prospect

The application of “dividing and conquering” and “caching” principle in development of molecular modeling algorithms is briefed. Historically, coarse graining and enhanced sampling have been two independent lines of methodological development. While they share the common goal of reducing local sampling, the formulations are completely different with distinct (dis)advantages. Coarse graining obtains partial transferability FF but loses resolution, enhanced sampling retains resolution but results are not transferable. The GSFE theory suggests a third strategy to directly approximate global joint distribution by superposition of LFEL, which may be learned from available dataset of either experimental or computational origin. Through integration of coordinate transformation, autodifferentiation and neural network implementation of GSFE, our recent work of protein structure refinement demonstrated that simultaneous realization of transferable in-resolution “caching” of local sampling is feasible. I hope this perspective stimulates further development of “dividing and conquering” strategies for complex molecular systems through more elegant, efficient and accurate ways of “caching” potentially repetitive computations in molecular modeling. With diverse molecular systems (e.g. nanomaterials, biomolecular systems), specialization of methodology is essential to take advantage of distinct constraints.

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