



Biomolecular simulations at the exascale: From drug design to organelles and beyond

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The rapid advancement in computational power available for research offers to bring not only quantitative improvements, but also qualitative changes in the field of biomolecular simulation. Here, we review the state of biomolecular dynamics simulations at the threshold to exascale resources becoming available. Both developments in parallel and distributed computing will be discussed, providing a perspective on the state of the art of both. A main focus will be on obtaining binding and conformational free energies, with an outlook to macromolecular complexes and (sub)cellular assemblies.

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Introduction

Molecular dynamics (MD) simulations have come of age [1]. Since their first application to biomolecular systems almost fifty years ago [2], both algorithmic developments as well as ever growing computational power have contributed to MD simulations now being an integral part of molecular biophysics. From deciphering protein function to computational drug design, from enzyme catalysis to the simulation of whole viruses or organelles, the field has matured substantially over

these years. In the example of classical atomistic MD simulations, the state of the art currently reaches milliseconds for small proteins on one side of the spectrum to nanoseconds for hundreds of millions of atom supramolecular complexes on the other side [3,4] (Figure 1).

Either by use of massively parallel simulation strategies or being able to access longer timescales, ever better statistics become available for quantitative simulation analysis. Ranging from alchemical methods in protein or drug design, to conformational landscapes sampled using collective coordinates employing a variety of methods, and to stitching simulations from massively parallel trajectories together via Markov state modeling and then assessing state populations from the statistics, methods to assess free energies have become commonplace.

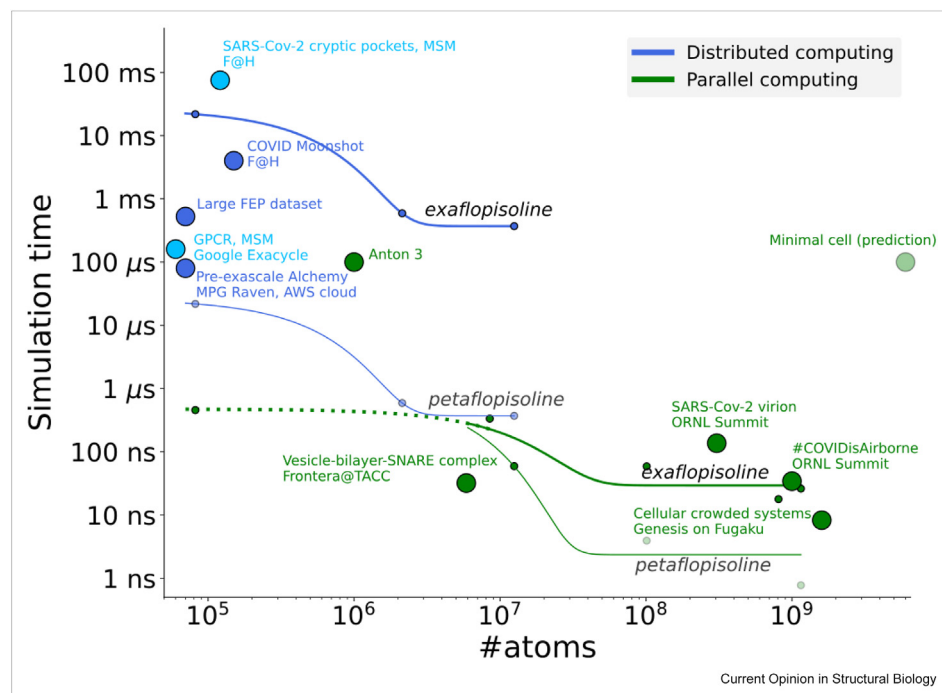
Currently at the dawn of exascale compute resources becoming available [5], opportunities are emerging across the field. In this review, we will concentrate on a subset of these developments with a focus on free energies, including alchemical free energy calculations of ligand–protein interactions, conformational free energy landscapes of individual biomolecules, as well as the sampling of supramolecular assemblies. We specifically focus on molecular dynamics simulations with molecular mechanics force fields, while we can recommend other reviews describing current state-of-the-art for large-scale biomolecular simulations with QM [6] and QM/MM [7].

Biomolecular simulations in drug design campaigns

In anticipation of the upcoming exascale resources, a number of studies reported on preparatory work to meet the challenge of efficiently utilizing such resources for alchemical free energy calculations. These studies on their own are still far from reaching the exascale level, rather they report on a concept of how protein–ligand binding free energy calculations can be distributed, deployed, and subsequently analyzed in an automated manner at a large scale.

In a proof of concept study, more than 1600 protein–ligand binding affinity differences were calculated by simultaneously utilizing 480 nodes on the MPG Garching supercomputer Raven [15]. Subsequently, these

Figure 1



Approximation of the simulation timescales for atomistic molecular dynamics simulations achievable on an exascale resource in 24 h of uninterrupted sampling. The upper blue line corresponds to the distributed computing scenario; the lower green line marks the true parallelization of a simulated system across an exascale machine. The isolines are based on the benchmarks in Refs. [8–10]: an exponential fit was used to extrapolate the results of the benchmarks to exa- and petaflop range (marked in small circles). Subsequently, another exponential fit was used to obtain the isolines through the extrapolated points. With the increase in system size, the achievable sampling time should eventually drop to zero, it is only due to the slow exponential decay the isolines appear to plateau for the larger system sizes. The dotted green line for smaller system sizes denotes the approximate range where parallelizing on peta- or exascale HPC machines is not efficient, i.e. using the full compute resource would rather degrade the performance. The simulation times were normalized to 4 fs integration time step. The large circles denote real simulation results without extrapolations. F@H SARS-Cov-2 cryptic pocket exploration [3] and the large FEP calculation [11] are marked assuming that all the calculations in the study could be performed in 24 h. Similarly, for COVID Moonshot [12] it was assumed that a single ‘Sprint’ would be performed in 24 h. Minimal cell system marked in the figure has not been simulated so far, but its size and aspired simulation time are described in Ref. [13]: we consider an atomistic representation of this system. References for the other marked studies: GPCR, MSM on Google Exacycle [14], Pre-exascale alchemy [15,10], Anton 3 [16], Vesicle-bilayer-SNARE complex [17], SARS-Cov-2 virion [9], #COVIDisAirborne [18], and Cellular crowded systems [8].

simulations were repeated on an AWS cloud infrastructure [10]. These studies, however, are only a preparatory step for approaching true exascale (Figure 1).

In one of the largest retrospective alchemical studies to date, Ross et al. [11] performed alchemical calculations for more than 1200 compounds (more than 2100 edges). This effort exceeds 500 μ s of total sampling time. While such sampling is readily achievable with the currently available computational resources, access to an exascale machine would allow to run all the simulations for this study in less than a day (Figure 1). Such a throughput would considerably speed up prospective protein-ligand binding affinity predictions in contemporary drug design campaigns [19,20].

In a community effort to find SARS-Cov-2 main protease (MPro) inhibitors [12], the established consortium gathered experts in structural biology as well as medicinal and computation chemistry. The designed compounds

were screened by means of alchemical relative binding free energy calculations prior to synthesis and evaluation in experimental assays. Simulations for the project were set up to make use of the Folding@home resources, which have recently reached an exascale mark [3]. The free energy calculations were performed in batches, termed ‘Sprints’. Assuming that one Sprint could be performed in one day (although this does not necessarily need to be the case) and considering a typical cumulative timescale noted in Ref. [12], we see that this alchemical approach could indeed come close to the exascale threshold for distributed computing (Figure 1).

The distributed computing character of many drug design campaigns renders it suitable to scale to large numbers of compounds by massively parallel deployment on exascale machines or distributed compute networks. Challenges therefore are mostly anticipated in the preparatory/management stages, efficient process orchestration as well as for data analysis, which should

ideally be integrated in the workflows rather than post-processed.

Sampling protein conformations with distributed computing

The kinetics of many biological phenomena, such as protein-ligand or protein–protein interactions, important to protein function and drug development can require millisecond to second sampling time.

However, even with exascale machines, it is not expected that simulation timescales beyond milliseconds will be immediately accessible for larger biomolecular complexes, as classical MD simulations will ultimately be limited by the need for small integration steps to avoid numerical error accumulation [21,22]. Therefore, distributed computing is not only applicable to large-scale alchemical free energy calculations, but also holds the promise to provide the computational resources and sampling strategies necessary to simulate the substantial conformational changes of proteins that occur on biologically relevant timescales. Using the first-ever exascale computing platform, created by interconnected personal computers from the Folding@home community of volunteers, Bowman and co-workers were able to scrutinize the opening dynamics of the SARS-CoV-2 spike protein complex and predict cryptic binding pockets near the active site and dimerization interface of MPro [3]. In fact, projects like Folding@home [23,24] and GPUGrid [25] have paved the way for massively parallel approaches to all-atom biomolecular simulations, by making use of graphics-processing unit acceleration, enhanced sampling methodologies and statistical methods such as Markov State Models [26] in MD-based high-performance computing [14].

A huge number of short, completely independent, or loosely coupled trajectories of MD simulations run in conjunction with biasing schemes [22,27], can be used to efficiently explore simulation systems even with complex underlying free energy landscapes. In a recent example, simulations (on Folding@home) of the 299-amino acid long apolipoprotein E (ApoE) monomer made it possible to sample heterogeneous protein conformations on the millisecond time scale and enabled model building to compare to and corroborate experimental single molecule studies of the system [28]. Structural ensembles generated on massively parallel computer architectures equivalent to millisecond MD simulations made it feasible to gain atomistic insight into the transiently populated, partially ordered states of the N-terminal domain of p53 [29], the binding of ligands to the intrinsically disordered protein p27 [30] and catalytically relevant conformational dynamics of the protein methyltransferase SETD8 [31].

Over the past two decades Anton supercomputers [16], a family of special-purpose hardware spearheaded by D.E. Shaw Research and designed to run long-timescale MD simulations have pushed Moore's limit for time-per-MD step. This notable, alternative approach to generic CPU/GPU machines reportedly reaches 100 μ s per day, for a 1 million atom system in the latest generation, Anton 3 [16]. Although the direct FLOPs comparison between general and special-purpose machines is cumbersome, projecting the achievable sampling times in Figure 1 places Anton 3 well above the petaflop isoline. Recent accounts of MD simulations in the tens to hundreds of microsecond range achieved with Anton include the cancer drug imatinib binding to Abelson tyrosine kinase [32], the binding of small-molecule compound fasudil to monomeric α -synuclein [33] and the discovery of a mutant-selective inhibitor of the fibroblast growth factor receptor 2 [34].

Large systems requiring massive parallelization

The continuous progress toward exascale allows researchers to build and simulate molecular systems of increased complexity, in recent years particularly focusing on whole virus particles (virions), individual organelles [35], subcellular architectures, such as the nuclear pore complex [36] or an entire gene locus [37], or the bacterial cytoplasm [38,8]. Each of these systems presented an outstanding computational challenge, and, accordingly, provided unique insights, not attainable through simulations on a smaller scale. Indeed, the ever-increasing complexity and size of simulated biomolecular systems are important not only from the boundary-pushing perspective, but also promise discoveries of unknown phenomena that arise only at such scales and conditions, markedly different from typical *in vitro*-like conditions of routine simulations and experiments.

The interest in modeling complete envelopes of virus particles has recently been fueled by the COVID-19 pandemic. To this end, initial computational efforts from the Amaro group focusing on virus particles of various influenza variants, reaching ca. 160 million atoms simulated for up to 440 ns [39], paved the way to simulations of a full SARS-CoV-2 viral envelope (~305 million atoms, 84 ns) [9] and peaked with simulations of a SARS-CoV-2 virion in the respiratory aerosol environment, resulting in one of the largest molecular systems simulated to date, of 1 billion atoms (simulated for 2.42 ns, with the reported performance of 2.55 PFLOPS (34.21 ns/day), using NAMD on the 4096 ORNL Summit nodes) [18].

These first-of-its-kind simulations of an aerosolized virus open completely new research avenues, such as atomistic views of the virus infection. A similar performance of ca.

12 ns/day has been also reported for a similarly sized system (>1 billion of atoms) representing bacterial cytoplasm, using GENESIS software on the FUGAKU supercomputer (and up to 8.30 ns for 1.62 billion atoms) [8].

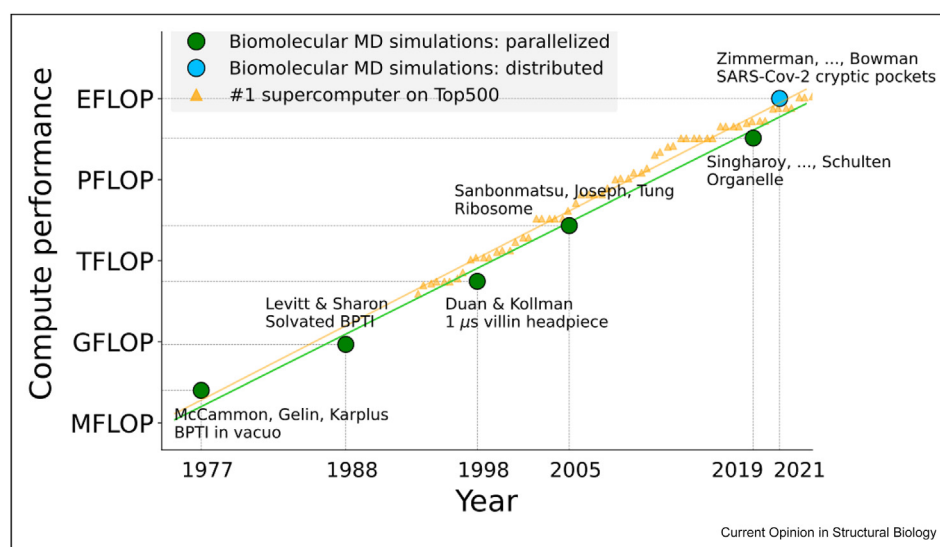
A natural next step in these modeling efforts is the construction and subsequent simulation of a whole-cell model. The availability of high resolution data of many cellular complexes, together with the integrative modeling tools, particularly AI-based, allowed researchers to build the first near-atomistic model of a whole minimal cell [13], Syn3A, owing to its extensive experimental characterization. The final system consisted of 550 million coarse-grained (MARTINI) particles (i.e. more than 6 billion atoms), involving a full chromosome, cytosol, and a lipid envelope. Putting aside the impressive feat of assembling such a massive system, it turned out the MD code of choice, GROMACS, was not ready (early 2023) to handle this system, thus the actual MD steps of this whole cell model have not been reported. The mentioned problems included handling multiple large molecules (e.g. the genome) spread over many domains in the domain decomposition approach, typically used to efficiently parallelize MD simulations. A related issue appeared in the simulations of the SARS-CoV-2 virion in the respiratory aerosol: a reduced particle density in an aerosol results in large ‘empty’ spaces, which is difficult to handle by standard domain decomposition. Finally, the amount of data generated by such simulations will be a challenge on its own.

These examples underlie the necessity of concomitant development of algorithms and novel software solutions (e.g. AI-driven), to be able to fully exploit the exascale (and beyond) resources for MD simulations. It has been nevertheless estimated that dedicated computational resources of the current type would enable reaching whole-cell simulations of 10–100 μ s (taking into account the typical MARTINI timestep of 20–30 fs). The fully atomistic simulations of such a system, however, would require further advancements of molecular dynamics engines in terms of scaling. For example, the widespread Particle Mesh Ewald (PME) method used to handle long-range electrostatic interactions might become too inefficient in all-atom simulations of cell-like systems. More flexible approaches, such as Fast Multipole Method (FMM) [40] offer a beneficial alternative to PME.

Outlook

The widespread use of MD simulations and their applications to various scientific questions naturally led to very diverse approaches of executing them on distinct computational architectures in recent years. In this review, we provide an overview over recent state-of-the-art studies that approach the exascale computing range (Figure 1). Apart from millisecond timescales becoming more and more reachable, another emerging trend is moving away from single molecule systems towards biomolecular complexes, protein or ligand series, to organelles and (sub)cellular assemblies.

Figure 2



Historic overview of the hallmark biomolecular simulations. The green symbols mark largest systems that were amenable for molecular dynamics simulations in the respective year. The cyan symbol reaches the exascale threshold by employing distributed computing Folding@home platform to simulate multiple smaller systems. The orange symbols correspond to the performance of the most powerful supercomputer from the Top500 list [42]. References for the marked studies: BPTI in vacuo [2], Solvated BPTI [43], 1 μ s villin headpiece [44], Ribosome [45], Organelle [35], and SARS-Cov-2 cryptic pockets [3].

Crossing the frontier to exascale computing power poses novel technical challenges concerning the administration and execution of the MD simulations (e.g. workflow management). These may eventually require building simulation architectures from “algorithms to transistors” [16] to make full use of capabilities provided by the (physical) exascale infrastructure, once its widespread availability is achieved. Preliminary studies have also already highlighted that the biosimulation community will likely have to evolve the ways by which the generated data is handled for analysis and shared (e.g. Exabyte problem) [41].

From a historical perspective, the compute resources used by biomolecular simulations follow closely the trend of the most powerful supercomputers (Figure 2). The extent, in terms of FLOPs, used by the hallmark applications of molecular dynamics grows exponentially, doubling every 15 months. At this rate, a 1000-fold increase in FLOP usage comes in about every 12 years. If the trend remains, the zettaflop barrier could be achieved in the next decade.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

No data was used for the research described in the article.

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