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Parallelization of Push-based System for Molecular Simulation Data Analysis with GPU

Iliiazbek Akhmedov, Yi-Cheng Tu †, Vladimir Grupcev, Joseph Fogarty, Sagar Pandit

Abstract—Modern simulation systems generate big amount of data, which consequently has to be analyzed in a timely fashion. Traditional database management systems follow principle of pulling the needed data, processing it, and then returning the results. This approach is then optimized by means of caching, storing in different structures, or doing some sacrifices on precision of the results to make it faster. When it comes to the point of doing various queries that require analysis of the whole data, this design has the following disadvantages: considerable overhead on random I/O while reading from the simulation output files and low throughput of the data that consequently results in long latency, and, if there was any indexing to optimize selections, overhead of storing those becomes too big, too.

There is a new approach to this problem presented in the previous paper – Push-based System for Molecular Simulation Data Analysis for processing network of queries proposed in the previous paper and its primary steps are: i) it uses traditional scan-based I/O framework to load the data from files to the main memory and then ii) the data is pushed through a network of queries which consequently filter the data and collect all the needed information which increases efficiency and data throughput. It has a considerable advantage in analysis of molecular simulation data, because it normally involves all the data sets to be processed by the queries.

In this paper, we propose improved version of Push-based System for Molecular Simulation Data Analysis. Its major difference with the previous design is usage of GPU for the actual processing part of the data flow. Using the same scan-based I/O framework the data is pushed through the network of queries which are processed by GPU, and due to the nature of science simulation data, this gives a big advantage for processing it faster and easier. In the old approach there were some custom data structures such as quad-tree for calculation of histograms to make the processing faster and those involved loss of data and some expectations from the data nature, too. In the new approach due to high performance of GPU processing and its nature, custom data structures were not even needed much though it didn't bear any loss in precision and performance.

Index Terms—Push-based system, molecular simulation, scientific databases, spatial distance histogram, GPU, parallel processing, CUDA.

I. Introduction

In various sciences simulation systems take big place and often times they may be the clue for results. One of such sciences, which is primarily related to this paper, is physics.

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In this case, from computer engineering point of view, simulations have the following flow:

- initial physical properties are given to simulation software as arguments which are interpreted and provided in a language defined by the simulation software
- then it runs for given amount of time that can generally last from milliseconds to months or even more
- finally, the file generated during the simulation is analyzed to extract useful information

The purpose of this paper is primarily focused on the third step, which basically involves the entire simulation data to be processed by the analysis software.

Big data processing is becoming one of the key issues with the amount of data being generated by modern systems. Eventually, this data is required to be processed on the fly, thus, analysis software should be able to handle massive data in a very short period of time. When working with huge volume of data, there are non-trivial issues arise. For example, it can take days and weeks to analyze big enough data sets, because the data cannot be simply loaded into memory, since it can get up to terabytes, thus, it has extra overhead because of widely used random access disk I/O framework to read from disk chunks by chunks. Besides it, analysis of the data can get even more complicated, since going through certain parts of the data once would not be enough, which leads to low throughput and efficiency of loaded data. One such example, the data analysis approach has polynomial complexity just for reading the data in order to come up with result, and since the data can't simply be saved in memory, it raises the overhead of disk I/O, too. Pull-based architectures in data processing engines are inefficient, since having a set of specific queries, in order to compute them all, it is needed to fetch data, filter it, and apply needed formulas. It is inefficient, since for every query the same chunk of data needs to be pulled into the memory at least the number of queries times or more in case of more complex queries.

As it has already been mentioned in the previous paper [1], one of the modern issues of analyzing massive data on the fly is social networks. . "In order for a system to be able to perform analytical examination of the data produced in such streaming media, the system should have the capability of fast data access. The reason, the millions of data records(tweets) produced every second. Moreover, these tweets may have different geographical origin, introducing different languages and forms and often times containing unsolicited messages, errors, malicious content, etc. Therefore, some low level data uniformity and cleaning on top of the data access and man-

agement issues should be considered and possibly incorporated in the process of analytical investigation in order to achieve relevant result. "[1]–[5]

The primary focus and problem in this paper is scientific data analysis. Particles simulation is one of the most popular methods of analyzing certain chemical reactions, physical processes, or other behavior of different materials. Molecular simulations (Molecular Dynamics) are applied in different fields and represent a method of analyzing physical movements of particles, atoms, and molecules in a fixed space with a given period of time, apparently with a possibility of giving initial state for each item that is involved in the process and can affect the system. This system is an N-body simulation. The number of atoms in simulations vary in hundred of thousands, particularly, we may observe two simulation systems of a collagen fiber structure and dipalmitoylphosphatidylcholine (DPPC) bilayer lipid system consisting of 890,000 and 402,400 atoms respectively on Figure 1. Simulation data represents number of records of physical properties such as mass, charge, velocity, coordinates, and forces for each item aggregated as frames, where each frame represents a snapshot of time, placed with a fixed time interval which may also vary depending on the simulation itself and simulation precision requirement. "Quantities measured during the simulations are analyzed to test the theoretical model [6], [7]. In short, the MS is proven and powerful tool for understanding the inner-workings of a biological system, by supplying a model description of the biophysical and biochemical processes that are being unfold at a nanoscopic scale." [1]

Scientist gives the properties to simulation software (for example, Gromacs), runs the simulations, and finally get the output file. The output file must then be analyzed to produce certain results which may help him come up with certain consensus on original theoretical model that resulted in the molecular simulation system [6]. Gromacs is simulation software tool that helps scientist to run the actual simulation. It is a molecular dynamics package primarily designed for biomolecular systems such as proteins and lipids. [9]. Besides the fact that it helps to generate the output files for the simulations, apparently it also helps to analyze the data itself, but the original problem is that it is not as optimized as it can be in order to analyze the data. Gromacs follows approach of pull-based design, which means that for any given query (e.g. total mass or total charge, which are very similar type of 1-body queries without sophisticated selection) it will pull data separately and generate addition overhead wasting disk read I/O in order to come up with the result just for a single query. As it has been proposed in the previous paper, in order to remedy such issues, the push-based design does exactly the opposite, where instead of loading the data on demand for each query, the queries are batched into a network, then the entire dataset is loaded chunk by chunk pushing it through the network which has its internal relationships and dependencies amonth the queries. In this case, since scientific simulation data is run once with specific physical properties, it is never modified, thus, on continuation, it will only append, which means that the processing can also be run on appended frames. [1] This type of approach has already been revised by other

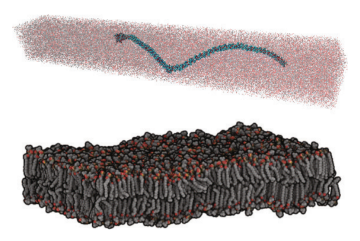


Fig. 1. Snapshots of two MS systems: a collagen fiber structure with 890,000 atoms (top) and a dipalmitoylphosphatidylcholine (DPPC) bi-layer lipid system with 402,400 atoms (bottom) [8]

systems [10]–[12] as of reusing loaded data through queries produced [13]–[15].

In this paper, we are incorporating parallelization into the processing part of the proposed design by means of CUDA programming language for GPU. Since storage of the simulation data is very expensive, it might come to the point of analyzing the data on the fly (meaning running the simulation and analyzing it at the same time in a streaming manner), which leads to a problem of optimizing the processing part of the design proposed in the previous paper, because time spent on generating data should be tried to conceal the time spent on the processing part by means of overlapping or simply running it in a quick manner.

A. Problem Statement

Simulation software systems, in general, follow the same methodology of running and storing simulation data. The simulation software system examples are: Gromacs [16], VMD [17], MDAnalysis [18], Wordom [19], MD-TRACKS [20], SimulaidOne [21], Charmm [22]. In the type of simulations brought up as examples above, the flow of the data is the following. Once the simulation is run, the output files are contained as trajectory files with descriptors (they contain information about space dimensions, number of atoms and frames, etc.) that can be easily transposed into simple flat files containing the physical properties atom by atom, frame by frame, which are consequently read and processed by the proposed push-based system. Since we have certain amount of queries needed to be run on given simulation data, generating high I/O traffic followed by design of pull-based system is not considerable. The approach proposed in the previous paper is very good in terms of performance in comparison with the original pull-based system [1]. The problem is still that some of the queries processed by those means are still improvable, especially taking into consideration the fact that in the used previous works for calculation of 2-body functions, which take the biggest time for processing [38], [39], we might have some error bounds, which might be unacceptable for certain simulation analysis where sacrifice on loss of

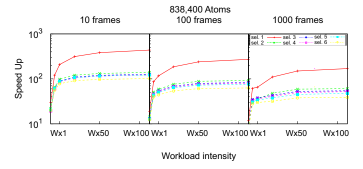


Fig. 2. Speed up over different levels of atom selection for 838,400 atoms [8]

data is unbearable. Besides it, although we might have huge performance boost on specific data structure as density map, we still have certain expectation from data nature (such as its uniformity), thus, it makes sense to have desire to simplify the processing and still having pure computation based on any values, considering that the queries should be available to be pre-programmed by user in a separate module of the code.

B. Our approach with improvement

Reading the data from files is a huge overhead, thus, speedup of push-based system in general is significant over pull-based system in our given case. It's been demonstrated in the previous paper with different amount of atoms, frames, and workloads. Since we have to load the data every time there is a different query versus push-based system loads a chunk of data once, it is quite noticeable how the framework contributes to efficiency of analysis of simulation data. You can observe some estimation samples in Figure 2. Even though the memory loading and pushing steps are the key features of the proposed design, nevertheless, having general knowledge about the network of queries, in this paper, we will try to focus on optimizing the actual execution of them.

Some of the queries may be very slow due to their nature on a sequential type of computation, especially, 2-body functions. Since the nature of the data that comes with simulation is basically physical properties of atoms, there is a lot of computation that involves independent primitive mathematical operations, which makes it a perfect problem for parallelism.

Although the original idea of push-based system for molecular simulation data analysis was focused on optimizing throughput and usage of loaded data, there were some extra approaches specifically for 2-body functions. For example, Density Map for Spacial Distance Histogram was used in order to avoid additional memory allocation and latency reduction with proven error bounds. SDH is a quite intensive and computational problem, especially with increasing amount of atoms.

We believe that having this nature of computational problems, the proposed GPU improved version of push-based system will significantly change in terms of performance by incorporating parallelism with CUDA.

C. Contribution and roadmap of the paper

We have designed and implemented a push-based system that can be used as a tool for analysis of MS systems. We have tested the system on real MS datasets. The experimental results evidently show the superiority of the proposed system over one of the most widely used MS analysis system across many scientific fields, i.e., GROMACS. The efficiency improvement ranges anywhere from 3 to 100 times improvement (of the overall running time) per set of selected queries run on a different selection of the MS data. If we take in account the number of such queries that the scientist run every day, we believe our system would be of great benefit to MS community. In addition to the higher efficiency that our system achieves, we also believe that our idea will initiate a new breed of database related, push-based systems that could be used to analyze the MS systems in a more efficient way. Having a push-based like system, we were also able to improve the computation of the more complex quantities. Namely, the computation of some such complex quantities involves computation of much less complex functions (sub-parts), and many such sub-parts are being shared among some of the complex quantities. So, by computing these more basic, sub-quantities in advance, we were able to achieve however minuscule time gain when computing the more complex queries.

The major technical contributions presented here are:

- Design the network (tree like) of the most commonly used queries in MS (physics);
- Build the system: design and build the modules, representing the quantities to be computed in an efficient manner, following the already built network;
- Develop a scientific simulation database benchmark that can be used for evaluating similar systems and products.

The remainder of this paper is arranged as follows: in Section II we give a survey of the systems used in the field of MS systems analysis. We continue the paper with Section III in which we show the design of the query network build from the most widely used quantities in MS system analysis. In Section IV we describe our push-based system for MS data analysis. Then, in Section V we present the benchmark designed to test our system as well as the results attained through comprehensive experiments run on real MS generated data. At the end, we conclude this paper with Section VI in which we give an overview of our possible future endeavors in the field of MS data analysis.

II. RELATED WORK

The idea of data streaming has been broadly used in many fields. The main usage however is aimed at processing live data generated online. There is an ocean of references for data stream management, but we believe the presentation in [23] encapsulates the majority of the ideas, problems and solutions. In the past decade, however, the database community started to follow the data stream idea to process stored data. Processes can take advantage of the streaming data at any time the data is being pushed through the system. This gives rise to push-based type design for data management systems. The idea of such push-based design was previously considered in projects such

as DataPath [10], Volcano [11] and QPipe [12] among others. These works show ideas in which the data-driven dataflow is compared to the demand-driven dataflow showing the need for the later. They also talk about maximizing the data and work sharing among queries at runtime. Essentially, we incorporate such ideas in our design of the push-based system.

On the other side, the scientific community has steadily progressed from processing massive data files towards employing database systems for the storage, acquisition, and analysis of large-scale scientific data [24], [25]. The widely used and popular relational database systems are conventionally designed and optimized to better manage the data produced by the business type applications. But such conventional database systems (DBMS) are not well equipped to deal with the type and quantity of scientific data, such as the data produced by the molecular simulations. In the recent past, the DBMS community has made some attempts into the design and construction of database systems optimized for handling scientific data. Such examples include the BDBMS project [26] that deals with annotation and provenance of the sequence data in biosciences, and the PeriScope project [27] is designed to efficiently handle declarative queries against bio sequences. On top of the aforementioned examples, there are also ideas for new DBMS frameworks aimed at the management of scientific data [28]–[30]. One of those systems is the SciDB [30], [31] and it is closest to the idea presented in this paper. SciDB is data management and analytics system that is primarily used is in application domains involving very big scale array data. This system, like the one presented in this paper is designed around a multi-dimensional array data-model and it uses arrays to store the data. SciDB stores petabytes of data on a number of machines and runs its queries on those machines. It is made for high performance, high-availability, fault tolerance, and scalability. However, to the best of our knowledge, it too follows the pull based design where its queries demand the data they need. As seen earlier in this paper, this type of design can impose I/O overhead and decrease the data throughput when doing the analysis. Aside the mentioned issue, the design and build of such DBMS optimized for scientific data management come with a additional challenges. Such challenges as well as their probable resolution are outlined in [32]. Recently, there have been some efforts aimed at designing and building MS data managements systems on top of relational databases. Such efforts are presented through projects like BioSimGrid [33] and SimDB [34] that were developed especially for molecular simulations. However, to the best of our knowledge, such systems still lack the efficiency needed for MS data management as well as efficient query processing strategies.

Generally, the data produced in the process of molecular simulation is being stored in large, plain files with no structure whatsoever. Queries, which are implemented in a stand alone programs within simulation/analysis systems, are executed onto such files producing the quantities that scientist use to analyze the molecular system. Such simulation and/or analysis packages include: Gromacs, VMD, MDAnalysis, Wordom, MD-TRACKS, SimulaidOne, Charmm among the others. But to the best of our knowledge, all of these systems work on a similar basis: they take a user defined query and execute

it onto the MS simulated data. In order for the query to be executed, the data has to be loaded into the main memory. Then the result is either produced onto the display or written to a file. When the next user query comes, the system again loads significant part of the dataset into the main memory and executes the query. We believe that there is a room for improvement of such systems, given the fact that many of the user defined queries executed during system's analysis are fairly static. In other words, there is a number of queries that a user would always want to execute on a given simulation data. Furthermore, the selections of MS data onto which such queries might be executed, are also fairly constant (i.e., oftentimes the user selects the same group of atoms (e.g., all hydrogen atoms) to calculate given quantity, like center of mass for instance). So, by precoding many such queries and running them automatically once the system has loaded the data into memory, we believe we can save a lot of time that otherwise would have been spent in loading the same data into main memory anytime a query is executed. On top of the automated query execution, our system can take user's query as input as well. With this, we believe our system is an improvement over the MS analysis systems that are used

III. NETWORK OF QUERIES

MS Queries. In order to study some important statistical features of an MS system, scientist need to "extract" various statistical quantities out of the data produced by the simulation. To achieve this, queries are executed against the data. Most of the gueries used in the analysis of MS systems are analytical in nature. Essentially, these analytical queries are mathematical functions that translate a selection of atoms (atoms' measurements) to a scalar, vector, a matrix, or a data cube [34]. Once the simulation is done, the analysis carried out will depend on the structure being studied as well as the features of the system that need exploring. In other words, not all system's quantities need to be computed every time the system is being analyzed. Some of the more popular queries, including density (atom counts), first-order statistics (mean), secondorder statistics (variance), and histograms among others, can be seen in Table I. The queries shown in this table are the ones that we have also incorporated in our system. Just to clarify some of the notation in Table I: we assume that the MS system comprises of n particles and r_i , m_i , c_i and q_i denote coordinates (vector form), mass, charge, and number of electrons of a particle i, respectively.

There are two types of queries/functions among the ones used to analyze an MS system. The first type are one-body functions. Such functions usually are algebraic functions [24] and only involve quantities(attributes) from a single atom at any given time in the process of computation. Each atom (atom's attributes) is being processed a constant number of times, thus the total running time of such functions/queries is O(n). This type of functions is very suitable for the idea of push-based system, or an online system in which the data is being read once and acted upon. In other words, in a single run of the incoming data, all such queries will produce useful final

Function Name	Equation/Description
Moment of Inertia	$I = \sum_{i=1}^{n} m_i r_i$
Moment of Inertia on z axis	$I_z = \sum_{i=1}^{n} m_i r_{zi}$ $M = \sum_{i=1}^{n} m_i$
Sum of masses	$M = \sum_{i=1}^{n} m_i$
Center of mass	$CoM = \frac{I}{M}$
Radius of Gyration	$RG = \sqrt{\frac{I_z}{M}}$
Dipole Moment	$D = \sum_{i=1}^{n} q_i r_i$
Dipole Histogram	$D_z = \sum_{i=1}^n \frac{D}{z}$
Electron Density	$D = \sum_{i=1}^{n} q_i r_i$ $D_z = \sum_{i=1}^{n} \frac{D}{z}$ $ED = \frac{\sum_{i=1}^{n} (e_i - q_i)}{dz \cdot x \cdot y}$
Heat Capacity	$HC = \frac{3000 \cdot \sqrt{T \cdot boltz}}{2 \cdot \sqrt{T} - n \cdot df \cdot VarT}$
Isothermal Compressibility	$I = \frac{VarV}{V_{avg} \cdot boltz \cdot T \cdot PresFac}$
Mean Square Displacement	$msd = \langle (r_{t+\Delta_t} - r_t)^2 \rangle$
Diffusion Constant	$D_t = \frac{6 \cdot msd(t)}{t}$
Velocity Autocorrelation	$V_{acor} = \langle (V_{t+\Delta_t} \cdot V_t) \rangle$
Force Autocorrelation	$F_{acor} = \langle (F_{t+\Delta_t} \cdot F_t) \rangle$
Density Function	Histogram of atom counts
SDH	Histogram of all distances
RDF	$rdf(r) = \frac{SDH(r)}{4 \cdot \pi \cdot r^2 \cdot \sigma_r \cdot \rho}$

TABLE I: Popular analytical queries in MS

results. Except the SDH and the RDF (i.e., the last two in the table), all other queries shown in Table I fall into this category of one-body functions. Most of these functions are defined on a single frame of the MS data. Only the autocorrelation functions are defined on two distinct frames.

The second type of functions are multi-body functions and are holistic in nature. The computation of such functions involve more than one atom's attributes and cannot produce final result in a single run of the MS data (i.e., if traditional methods are used for their computation). Such queries include the Radial Distribution Function (RDF) [6], [35], [36] as well as some quantities associated with chemical shifts [37]. Generally, such functions are computed through histograms. For instance, the RDF is obtained from a histogram of all pairwise atom distances (this is the Spatial Distance Histogram or SDH). The traditional, straightforward (often the bruteforce) way of computing these holistic functions is a very time consuming process. On top of that, these methods cannot produce the final result in a single run of the MS data, making such functions unsuitable for our idea of a push-based system. However, in our previous work, we have designed a data structure together with an algorithm that opens up the possibility for such queries to be executed in a push-based type environment. Further details on this are given in Section IV.

Fig. 3 represents an idea to show how such query processing system can be improved. The idea behind it is that some of the queries share same sub-routines. Having all the queries made as separate modules, this sub-routines can be computed once the data is being pushed through the system and then be used anytime a more complex query needs it. Having in mind the amount of data in a single frame that the queries (sub-routines as well) need to go through, and the fact that in a single MS there are thousands of frames, we believe this can be immense improvement in terms of total running time.

IV. BUILDING THE SYSTEM

A. MS Data retrieval and in memory organization

A typical MS system generates and stores the data in a number of trajectory files, usually including multiple frames (snapshots of the simulated system taken at certain time intervals). Such MS generated data oftentimes goes through a simple lossless compression and, depending on the simulation software it may be stored in a binary format. Such trajectory file format is one of most often used MS file format (e.g., GROMACS, PDB). But such format is unrecognizable to our system. So our system has to do three things before it starts executing the queries: 1) Read the MS data from a trajectory file, 2) Translate the MS data to a form recognizable to our system, and 3) Load the data to memory.

Data read - transform. As mentioned above, in order for our system to be able to read the MS generated data, the data needs to be transformed. The reason for this is following: the MS data is stored in multiple files and possibly in different formats as well. One such file holds the global data (identifying the system and the simulation). Another file holds each frame's data. The frame data contain general information about the frame, but the main part is a sequential list of each atom's info, including atom's mass, position, charge, number of electrons, velocities, forces, etc. Another file (topology file) holds the molecule/residue info, essentially identifying what atom belongs to which molecule.

So, in order to extract the data from these files, we have created a sort of "extractor/transformer" of the attributes needed for the execution of the queries in our system. This transformer, essentially, is a separate piece of code that does three things: 1) it reads the MS generated data that the MS system stores in one of the often used MS file format (e.g., GROMACS); 2) it translates the data into a format that our system can read (taking only the information our system needs); and 3) it stores the data in a file that has basic structure to it. So, in the end, the data transformer produces a data file that our system takes as input. This code serves as a connection between an MS system (e.g., GROMACS) and our system. With this, our system can essentially be used as an add-on to GROMACS or other simulation systems and help improve the efficiency of the data analysis.

Data organization in main memory. Once the data is in

¹This paragraph talks only about the data organization used by one-body queries. For two-body queries (e.g., SDH), the data organization is discussed later on.

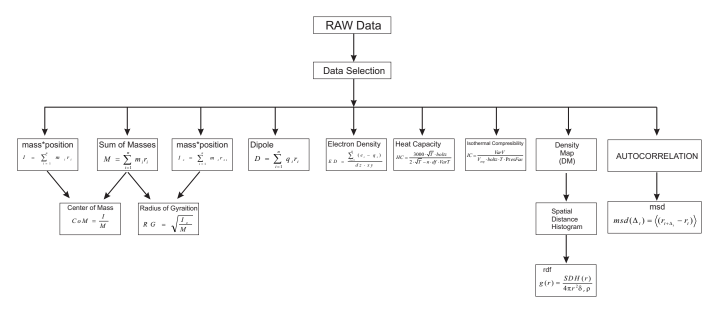


Fig. 3. MS Modules Structure

a format our system can read, the data is being loaded into the main memory one frame at a time. The organization of the in-memory particle's data is in the form of a simple, two dimensional array where a single row represents an atom in the system with all its attributes (e.g., coordinates, mass, charge, residue info, etc.). We also keep (in a one dimensional array) crucial system's information for each frame, like temperature, energy, pressure, etc. We have used such structures because they are very suitable for simulating a push-based type of system: a simple sequential read of the array gives that online type of data stream. So, as the system reads the array, it pushes the data onto the query-modules. As mentioned earlier, the one-body (algebraic) queries will produce a final result at the end of the first sequential read. However, the two-body (holistic) functions, like SDH and RDF, cannot do this in a single read of the data.² Therefore, each frame's data array can be continuously read (in a loop manner) as many times as a query needs it.

B. Query modules

As mentioned earlier, there are two types of functions/queries used for analysis of MS systems: algebraic or one-body, and holistic or two-body queries (these in general can be multi-body, but in this paper we only deal with a two-body functions).

1) One-body queries: Most of the query modules in Table I (except the SDH and RDF) are not that involved, only containing computations of fairly simple, one-body functions. These queries were coded as separate modules in our system. Each of these modules take few attributes as input (e.g., atom selection, frames selection(for the autocorrelation functions), number of atoms, etc.). The system pushes the data as it

becomes available onto these modules. The queries are being executed on the selection and are put in a "ready" mode, awaiting the next frame's data. First, the more basic queries, like total mass, are being computed. The results of such queries are temporary stored (in main memory) and are available for use anytime a more complex query needs them.

2) Two-body queries: In general, queries involving two-body functions are a bit more complex and cannot provide the final result in a single data read if a straightforward method is used for their computation. However, in the proposed system we have incorporated a data structure and an algorithm for the SDH (also RDF) from our previous work that is suitable for push-based type of system. In this subsection we give a brief description of the data structure (DM) and the algorithm (DM-SDH) designed in [38], [39] and implemented in the system proposed in this paper. For more detailed information, please refer to our previously published work on this topic [38], [39]

The data structure. The simulation data space is represented by a conceptual data structure we named Density Map (DM). The density map splits the simulation space into a grid of equal size regions (or cells). The cells are cubes in 3D and squares in 2D³ Resolution of a density map is the reciprocal of the cell size in that density map. In order to generate higher resolution density map, we split each cell of the current resolution's grid into four smaller cells of equal size. This design allows us to use a region quad-tree [40] to organize density maps of the same data but with different resolutions. So, essentially, a node in the quad-tree represents a single cell from the DM. Therefore, a density map of a certain resolution basically is the set of all nodes of one level of the tree. Each of the tree nodes records the cell's location in the density map (coordinates of corner points) as well as the number of particles in each cell. We name the afore describe tree the Density-Map tree (DMtree).

²However, we have previously deigned and created a data structure and an algorithm that can take the advantage of a single data read and produce final results for SDH computation [38], [39]. We have incorporate this into our system presented in this paper.

³In this paper, we focus only on the 2D data to elaborate and illustrate the proposed ideas.

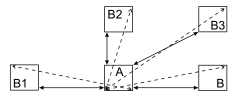


Fig. 4. Computing minimum (i.e., length of solid lines) and maximum distance (i.e., length of dashed lines) range between two cells

The algorithm. The essential part of the DM-SDH algorithm is a procedure named RESOLVETWOCELLS. The input to this procedure are two cells from the density map (e.g., A and B in Fig. 4). It computes, in constant time, the minimum and maximum distance between the two cells. A pair of cells is resolvable if both the min and max distance between them fall into the same SDH bucket i. If that is the case, the distance count of that bucket is being increase by $n_A n_B$ (n_A and n_B are the number of particles in cell A and B, respectively). Otherwise, the cells are non-resolvable and we either:

- (1) Go to the next density map with higher resolution and resolve all children of A with those of B, or
- (2) If leaf-level has been reached: compute every distance between particles of A and B and update the histogram accordingly.

In order to generate the complete SDH, the RESOLVETWO-CELLS procedure is executed for all pairs of cells for a given density map DM_k and the algorithm would recursively call the procedure (action (1) above) until leaf-level has been reached (action (2) above).

On top of the aforementioned DM-SDH algorithm, we have also incorporated two approximate SDH algorithms (ADM-SDH), introduced and described in [39], and [41]. These approximate algorithms are substantially faster than the the brute-force algorithm and also than the DM-SDH algorithm as they take advantage of some heuristic. For more details on the ADM-SDH algorithms please see the aforementioned work.

C. Working of the system

In this subsection we give a brief overview of how the system works at runtime. Please note that the first, preliminary part is only executed once, i.e., the data transformation from MS data files to a file that our system can read.

Here are the steps taken through out the analysis:

- (1) Execute the data transformer
 - (a) Read the MS data from trajectory files
 - (b) Extract the info needed for our system
 - (c) Save the read data to a file recognizable to the system
- (2) Load the data into main memory (one frame at a time)
 - (a) Load data into a double array (for one-body queries)
 - (b) Load data into the quad-tree structure (for two-body queries:SDH, RDF)
- (3) Push the data to all queries
- (4) A query, if available, acts upon the pushed data (first executing the lower level, sub-queries)
- (5) Store intermediate results (results of sub-queries)

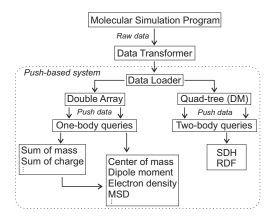


Fig. 5. Push-based system flow

- (6) Repeat steps 3-5 if needed.
- (7) Output results
- (8) Go to step 2 and load the next frame (if needed).

Fig. 5 depicts the flow of the system.

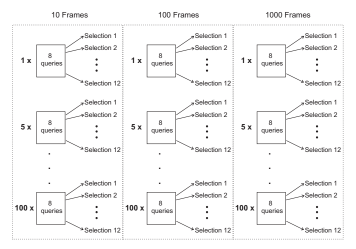
Step 2, loading the data into main memory is different for SDH query compared to the one for the one-body queries. The reason for that, as mentioned earlier, is the different data structure used to store the data in memory. While we use double array to store the data for the one-body queries, we use quad-tree like data structure to store the data needed to compute the SDH. The loading to the double array is straightforward. However, to load the quad-tree structure, we need to use some of the info from the data itself. Namely, the coordinates of the atoms are used to determine in which tree node an atom belongs. That way we build the so called density map (DM), i.e., the different regions with a certain number of atoms in them (including all the atom's attributes). So basically, to solve the SDH problem in a push-based manner, we convert the problem into populating a data structure in push-based manner. This data structure will then be used as an input to our DM-SDH algorithm that, although not completely in "on-the-fly" way, is a great improvement over the naive methods used in much of today's MS analysis systems (i.e., GROMACS, PDB, CHARMM, etc.).

V. EXPERIMENTAL RESULTS

The system was implemented in C++ programming language and tested on real molecular simulation data sets. The experiments were carried out on an Apple MacPro machine with 8GB of physical memory and two Quad-Core Intel Xeon 3GHz processors. The MacPro was running OS X Mavericks 10.9.3 operating system. We have compared the results obtained by our system to those obtained by running the analysis through GROMACS (v. 4.5.7). Both systems were analyzing the same data sets.

Data sets. In our experiments, six data sets from different simulations were used. All simulations were done on a POPC ⁴ lipid bilayer, but were all set to produce data of different

⁴POPC is a chemical compound composed of a diacylglycerol and phospholipid. Its full name is 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine and it is one of the most important lipids in bio-physical molecular simulation.



Such workload set ups are being executed on 6 different size data sets : 50K, 200K, 800K, 2.5M, 4.5M, and 8.8M atoms

Fig. 6. Workload setup

sizes (i.e., different number of particles in the simulation). Namely, we have tested the system on simulations with $52,400;\ 209,600;\ 838,400;\ 2.5M;\ 4.4M;$ and 8.8M atoms. Also, since the simulations were run separately, they produced six different MS systems with distinct characteristics (distinct structure, atom's positioning, etc.). From all of the generated data sets we have randomly selected sets of 10,100, and 1000 consecutive frames for the purpose of our experiments. This gave us 18 different datasets on which we tested our system.

Query work load. Two types of query workload were used: 1) one involving one-body queries only 2) one including two-body queries (SDH and RDF) as well. The reason for this is that GROMACS, the system we used to compare our system to, only has a naive method of solving the RDF (SDH) problem (like almost all MS analysis systems). In our system we have incorporated SDH (RDF) algorithms that are far more superior to the naive method, and comparing the systems like that would not have been fair (we believe).

One-body queries only. The following set of one-body queries were included in the test workload: mean square displacement (msd), radius of gyration, dipole moment, center of mass, velocity autocorrelation, electron density, mass density, and charge density. This set of queries were pointed to us, by a group in the physics field with extensive MS background, as one of the most commonly used in the field of collagen bilayer MS system analysis. A workload group contains all 8 queries executed on one of the 12 selections, making 12 groups. Such groups are executed on six different size data sets, with 10, 100 and 1000 frames. This workload is then repeated 5 more times, by executing each of the queries in the groups 5, 10, 25, 50, and 100 times, essentially just magnifying the workload intensity. In total, we have 12x6x3x6 = 1,296different workload setups to test the system on. Fig. 6 shows the organization of the workload setup.

A. Benchmark

Through extensive collaboration with a research group from the Physics department at USF, we have come up with a benchmark that can be used for testing the efficiency of an analysis system for molecular simulations. The benchmark consist of three essential parts: 1. Simulation data produced by an MS, 2. Queries that are to be executed onto that data in order to produce some information of interest, and 3.Benchmark parameters that control the size of the benchmark.

- 1) Benchmark Data: The data used in the benchmark was real molecular simulation data, produced through the GROMACS MS system. The initial, pre-simulation data file consisted of 200 POPC and 12000 solvent molecules, or 12200 molecules in total. This type of system was used because it is sufficiently diverse, containing enough distinct POPC and solvent molecules (e.g., each POPC molecule includes approximately 52 different atoms) and yet simple enough to be easily transformed into another system of different size. By using the *qenconf* function in GROMACS, we produced pre-simulation files of different sizes (essentially by changing the system's size (box)). Six different sized pre-simulation files were created. A molecular simulation was then run on these 6 files, each producing an MS system of certain size (volume/number of particles). All of the simulations were set up to produce 1000 frames (snapshots in time of the systems), each frame containing the same number of particles as the base one. The produced files contained: 52, 400; 209, 600; 838, 400; 2.5M; 4.4M; and 8.8M atoms per frame. So, for example, the file with 52,000 atoms holds 52,000,000 records in total (1000 frames, each containing 52,000 records). As mentioned earlier, this simulation data comes mostly in binary formats and in trajectory files having a lot of unneeded overhead. Therefore, it was transformed to a data arrays files containing only crucial information of the particles and the system. The size of the files ranged from 135MB for 52,000 atoms to 24GB for 8.8 million atoms (this is for data with 100 frames).
- 2) Benchmark Queries: The queries selected to be included in this benchmark were derived through a thorough observation of the way an MS system is being analyzed. They were found to be the base of the analysis of many MS systems. In other words, no mater how small or big the analysis was, these queries were included in that analysis. As mentioned earlier, they are of two types: one-body (and algebraic) and two-boy (and holistic). Table I shows these queries.
- 3) Benchmark Parameters: There are several parameters that can be used to control the overall size of the system. We divide the parameters into two groups:

Data size parameters:

- Select different sized dataset
- Number of frames
- Data selection (within the selected dataset) onto which the queries are being executed

Workload size parameters:

- Number of queries to be executed
- Number of times each query is executed

By changing these parameters, we can produce a versatile testing benchmark for MS analysis systems.

B. Results

We have run extensive experiments over all the different setups of workload mentioned previously in this section. However, in this paper we present only the workload setups of 4 data size sets: 838,400, 2.5M, 4.2M, and 8.8M atomsbecause we believe they convey enough information about the efficiency of our system compared to that of the Gromacs system. The running times of our push-based system were compared to those of the Gromacs system. The first set of figures, namely Figures ??-??, represent the speedup that our system obtains over the Gromacs system with various atoms selection levels. We define the selection levels based on the number of comparisons we have to make in order to extract the needed group(selection) of atoms. For example, if we want to do analysis on all molecules containing oxygen, or hydrogen, or carbon we would go over each molecule and compare its components to the selection list. The bigger the selection list, the higher the select level in our system. For better visualization, we note three different selection levels: high (at least 10 comparisons made), medium (between 1 and 10 comparisons made), and low select level (with one or less comparisons made). As seen in the figures, for high selection level, the speedup is smaller compared to that achieved in low select levels. The reason for this, we believe is in that the amount of time our system spends extracting the atoms group increases with the level of selection. Even though our system still shows considerable speedup over Gromacs in high level selections, we do believe there is room for improvement in our system and that is our immediate future work we are planning on doing. These figures also show the relation of the speedup to the workload intensity, i.e., the higher the workload intensity the higher the speedup.

The connection between the workload intensity and the speedup is better represented in the next set of figures, Figures ??-??. They show the speedup our system achieves over the Gromacs system on a varying workload intensity. Each of those figures show the speedup with different dataset sizes (e.g., 838,000, 2,567,600 atoms, etc.), including 10, 100, and 1000 data frames. The speedup is calculated simply as a ratio between the running time of our system on a certain set of workload and that of the Gromacs system on the same workload. These figures show that the speedup over varying workload intensity achieved by our system ranges anywhere from about 10 to 1000 times, depending on the size of the dataset, number of frames and the selection of the atoms.

Figure ??, shows the speedup our system achieves over all workload intensity (average workload intensity) with varying dataset sizes. It is clear that, again our system has better performance than the Gromacs system. The speedup presented in this set of figures ranges anywhere from about 15 to 650 times

The last figure, Figure ??, shows the speedup our system achieves over all workload intensity and all select levels with varying dataset sizes. This figure, in a way, summarizes the previous two sets of figures, bringing together the workload and the different selections through the average. It is clear that, again our system has better performance than the Gromacs system. The speedup ranges anywhere from 50 to 250.

All four sets of figures show that such push-based design has clear advantages over the pull-based type of design incorporated in the Gromacs system.

VI. CONCLUSIONS AND FUTURE WORK

The objective of our work is to design and implement improved data analysis system that can be used in the field of molecular simulation system's analysis. In this paper, we introduce the idea for such system. We build our system on a push-based type design, where data from data arrays is being pushed onto available queries in the system. These queries are being executed on the pushed data and produce intermediate / final result that would be used as part of the data analysis. We are able to achieve an improvement over existing, pull-based type designs because of the I/O overhead such designs introduce when dealing with large volumes of scientific data. Also, our queries are able to be executed on the same stream of data, making it suitable solution for streaming circumstances. We designed a benchmark that can be used to test data analysis systems. This benchmark comprises of three parts: 1)benchmark data, 2)benchmark queries, and 3) benchmark parameters. We use this benchmark to compare our system to one of the most frequently used MS analysis systems, Gromacs. The efficiency and speedup achieved by our system is supported by extensive experiments and their results. The results show that our push-based design achieves up to about 1000 times speedup in comparison to a pull-based design, i.e., Gromacs.

One direction of our future work will be to further improve our push-based design. Through the extensive experiments we have learned that our design can be improved when the atom selection clause involves many conditions. This improvement may be in the direction of improving the algorithmic part, but it can also be in the direction of improving the data presentation/organization we have used in the system.

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