Scalable Online Comparative Genomics of Mononucleosomes: A BigJob

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ABSTRACT

Our goal is to develop workflows for simulating arbitrary collections of mononucleosomes as an "on-demand" analysis tool for comparative genomics at atomic resolution. The limiting factor is resource availability. The aim of this paper is to document and share our experiences in providing a general-purpose, easy to use and extensible solution for such computations. At the core it involves supporting the execution of high-throughput workloads of high-performance biomolecular simulations on multiple XSEDE machines. Although conceptually simple, it is still a difficult practical problem to solve, especially in a flexible, robust, scalable manner. Specifically, we employ BigJob – an interoperable Pilot-Job. The bulk of this paper is about presenting our experience in executing a very large number of ensembles including the associated non-trivial data management problem. Our experience suggests that although a nascent and fledling technology, BigJob provides a flexible and scalable pilot-job to support workloads that were hitherto not easy, if not possible.

Categories and Subject Descriptors

H.4 [Information Systems Applications]: Miscellaneous; D.2.8 [Software Engineering]: Metrics—complexity measures, performance measures

General Terms

Experience, Technology

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Keywords

HPC, Distributed Computing, NAMD, MD, Large Scale, BigJob, SAGA, Python, Computational Workflow, XSEDE resources

INTRODUCTION 1.

The need to run many distinct instances of molecular dynamics (MD) simulations concurrently arise in many different scientific contexts: enhanced sampling and configurational space exploration to name just a couple. As a specific critical example of a scientific problem that critically depends upon the ability to execute and control large-scale ensembles of high-performance computing, we investigate the domain of "online genomics".

Nucleosomes represent the fundamental structural unit of eukaryotic genomes. As such, the nucleosome's structure and dynamics is relevant to all genomic processes. Our goal is to develop workflows for simulating arbitrary collections of mononucleosomes as an on-demand analysis tool for on-line comparative genomics at atomic resolution.

As an example of the aforementioned problem, we need to simulate 105 mononucleosome configurations for 20 nanoseconds in 1 nanosecond intervals, a total of 2100 tasks. As it typically takes 1 hour to simulate 1 nanosecond using 240 cores on Lonestar, the total compute time is about 504,000SU. By, running 10 simulations in tandem on 2400 cores the wall time is only 210 hours, or just under 10 days. This is a non-trivial undertaking and requires support for scalable, flexible and advanced execution modes. Since the 105 configurations are independent, a properly implemented solution allows us to complete the study in 1 day given sufficient resources. Here we demonstrate that on-line comparative genomics of mononucleosomes is feasible. The aim of this paper is to document and share our experiences in executing high-throughput workloads of high-performance biomolecular simulations on multiple XSEDE machines.

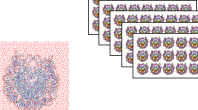
Whereas single high-performance simulations are commonplace and the norm, it still remains a challenge to execute many instances of a high-performance simulation concurrently. Contributing to the problem is the fact that most supercomputing centers, including XSEDE, have their environments tuned to support mostly single-job oriented work loads. On the one hand, although workflows and gateways have improved the mix of work loads supported, they do not present a natural "user interface" for multiple instances. In response to the deficiency, there are many tools and approaches to manage many multiple instances of highperformance simulations, however very few present a general purpose, uniform and flexible approach. Our approach builds upon important but incremental advances - both conceptual and implementation, of Pilot-Jobs (a conceptual model of which can be described by the P Model [4]). Specifically, we employ BigJob – an interoperable Pilot-Job. We use the Pilot-API to express the abstract workflow associated high-throughput workload.

An important difference between this paper and a previous related paper [6], investigating BigJob's use for nucleosomes, is that this work is focussed on understanding and improving simulations on a single machine (as opposed to collective distribution); additionally, the science problem being addressed in the current publication is distinct from Ref. [6].

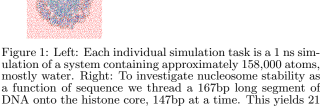
Our paper is logically organized into the following sections. After the introduction and the underlying scientific motivation (§1), we discuss the basics of pilot-jobs, we introduce BigJob and the interoperability layer (SAGA) and how they couple to provide an interoperable and flexible pilot-job. We then outline the abstract workflow of the scientific problem of interest, and examine how we use the Pilot-API to implement it. The bulk of this paper is about presenting our experience in executing a very large number of ensembles including the associated non-trivial data management problem. Our experience suggests that although a nascent and fledling technology, BigJob provides a flexible and scalable pilot-job to support workloads that were hitherto not easy, if not possible.

2. SCIENTIFIC PROBLEM

A genome is more than just a sequence of DNA. In eukaryotic organisms it exists as a biolecular complex of DNA and proteins called chromatin that resides inside the cell nucleus[5]. The length of DNA is typically many orders of magnitude larger than the diameter of the nucleus. The width of DNA is 2 nm so it can be readily folded to fit inside the nucleus. Herein lies the problem. Folding affects access to the instructions encoded in the DNA and therefore all biologic mechanisms that require these instructions. The proteins that affect DNA folding are called histones. Eight histones associated as two dimers and a tetramer ([H2A - $H2B[H3 - H4]_2[H2A - H2B]$), wrap 147 base pairs of DNA into 1.7 turns of a superhelix. The histone-DNA complex is known as the nucleosome. To a first approximation, folding of the DNA, or at least local access, is determined by the location and conformation of all nucleosomes on a chromosome. Genome wide maps of nucleosome positioning are now readily available, but the rules governing nucleosome positioning and the dynamics of nucleosomes are not well understood. X-ray crystallographic studies have provided a number of atomic resolution structures of the nucleosome [7], but because of experimental difficulties growing crystals for an arbitrary nucleosome, all structures to date contain essentially the same sequence of DNA and represent only one con-



(Water not shown for clarity.)



separate systems that must be simulated. The entire sim-

ulation ensemble includes 5 sets of 21 systems, as pictured.

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formation of the nucleosome. The nucleosome is a dynamic entity[2] that can exist in various states of association[8]. A canonical octasome can be formed with $\mathcal{O}(4^{147}) \approx (10^{88})$ different sequences of DNA. An exhaustive study of nucleosome structure and dynamics is not possible. Our goal is to develop computational techniques that allow us to compare ensembles of nucleosomes for an arbitrary realization of the nucleosome. Having such capabilities available "ondemand" provides a powerful tool for comparative genomics. In our current studies we are investigating relationships between nucleosome conformation and dynamics and the material properties of DNA as a function of DNA sequence. To achieve this end we thread different sequences of DNA onto available x-ray structures of the nucleosome. Computationally, we are deliberately mis-positioning nucleosomes to determine the rules governing nucleosome positioning.

2.1 Nucleosome Modeling Study

In our previous study [6], we simulated 336 nucleosomes that were experimentally determined to be associated with the most highly occupied and least variable nucleosome positions in the yeast genome[3]. This simulation study contains 50 times more nucleosomes than in any other molecular dynamics study of the nucleosome, and collectively it represents 6.7 microseconds of dynamics, approximately 8 times more than any other nucleosome simulation study [1].

In the current study, we chose five sequences representing 167 base pair of DNA, that are known via experiment to be nucleosome free. For each of the five nucleosome free regions, we model 21 systems, see Figure 1. One system represents the center of the sequence. The other twenty represent 10 neighboring positions on each side of the center position, i.e. one full helical repeat of the DNA in either direction. Threading one helix repeat of the DNA around the histone core allows us to investigate the contribution from sequence specific defects in the DNA, e.g. DNA bends or highly flexible regions. As each helix repeat length of DNA is thread around the histone core such defects will be oriented toward, away and then back toward the histone core.

In total, there are 105 systems to model. Each is simulated for 20 ns using NAMD as the compute engine and Amber force field parameters. This is the typical simulation methodology for nucleosome simulations[1]. Each 20 ns trajectory is divided into twenty tasks where each task is

a 1 ns long simulation of a given position. We thus have 2,100 tasks consisting of 105 independent threads. Each of the 2,100 tasks requires 41MB of input data, runs for \approx 1 hour on 240 processors, and generates 3.6GB of data. In total this is nearly 7.6TB of data and over 540,000 SU of compute time.

2.2 Computational Requirements

Aside from DNA sequence, the systems simulated are nearly identical to our previous study[6]. Each system is represented by an all atom model containing $\approx 158,000$ atoms including: 13,046 atoms of protein, \approx 9600 DNA, 426 ions, and $\approx 135,084$ water atoms (differences due to differences in DNA sequence). The input data required to start any single task is $\approx 41 \text{MB}$: parm (28.1MB) molecular topology and parameter data, crd (5.5MB), vel (3.6MB), coor (3.6MB) files containing coordinate and velocity information, and xsc (177 B) periodic boundary condition data. Each task generates 3.6GB of output data including: dcd (1.8GB) coordinate file in highly compression binary format, dvd (1.8GB) velocity file in highly compressed binary format, out (167MB) text based log file, xst (68KB) dimensions for the period cell size as function time, restart files coor (3.7MB) and vel (311.4KB) containing coordinate and velocity information. Running 105 tasks concurrently would require 4.3 GB of input data, generate 7.56 TB of output data and could utilize 25,200 processors efficiently in a single 24hr run to yield 2.1 microseconds of molecular dynamics data.

3. SOFTWARE TOOLS AND COMPUTATIONAL INFRASTRUCTURE

XSEDE is inherently a complex infrastructure with heterogeneous resources. In order to harness the power of such a distributed environment, we utilize Pilot-Jobs. A Pilot-Job is a mechanism by which a proxy for the actual simulations is submitted on the resource to be utilized; this proxy agent in turn, is given responsibility to convey to the application the availability of resources and also influence which tasks are executed. The abstraction of a Pilot-Job generalizes the reoccurring concept of utilizing a placeholder job as a container for a set of compute tasks; instances of that placeholder job are commonly referred to as Pilot-Jobs or pilots.

In general, Pilot-Abstractions provide a suitable means to marshal heterogeneous sets of both compute and data resources and support the efficient utilization of different kinds of commercial as well science cloud resources. Pilot-Abstractions have been extensively used on both HPC and HTC infrastructures for a range of application scenarios as a resource management abstraction to, (i) improve the utilization of resources, (ii) to reduce wait times of a collection of tasks, (iii) to facilitate bulk or high-throughput simulations where multiple jobs need to be submitted which would otherwise saturate the queuing system, and (iv) as a basis to implement application specific execution, scheduling and policy decisions

The P* model [4], a model for Pilot-Abstractions, worked to clearly define the computation and data components of a distributed application as 'compute units' and 'data units' in the context of Pilot-Jobs and Pilot-Data. A compute unit describes a self-containing piece of work, e.g. a computational task that potentially operates on a set of input data,

while a data unit is a container for a logical group of data that is often accessed together or comprises a larger set of data; e.g. a data file or chunk.

3.1 BigJob: A Pilot-based Framework

BigJob is a pilot-job system implementation which provides a framework for running many types of distributed applications – including but not limited to very-large scale parallel simulations, many small high-throughput simulations, or ensemble-based workflows. Consistent with the P* model, BigJob (BJ) provides a unified run-time environment for Pilot-Jobs on heterogeneous infrastructures. For this purpose, BigJob provides a higher-level, unifying interface to heterogeneous and/or distributed data and compute resources. The framework is accessed via the Pilot-API, which provides two key abstractions: Pilot-Job and Pilot-Data.

Applications can specify their resource requirements using a Pilot description. In the compute case, the user typically specifies the application to run as well as the number of cores required by their application. Pilots are started via the Pilot-Compute Service. BigJob eliminates the need to interact with different kinds of compute resources, e. g. batch-style HPC/HTC resources as well as cloud resources, and provides a unified abstraction for allocating resources.

BigJob has seen its widest usage across the heterogeneous resources that XSEDE provides. Simple installation into user space on any resource that supports Python 2.5 or greater makes the uptake of BigJob easy for the end user. BigJob supports thousands of jobs and millions of SUs on XSEDE. It has been at the heart of two recent and successful ECSS projects.

3.2 SAGA: Interoperability Layer

In order for BigJob to work on heterogeneous resources, it requires an interoperability layer which provides access to a variety of middleware. This is achieved through the use of The Simple API for Grid Applications (SAGA). SAGA defines a high-level access mechanism for distributed infrastructure components like job schedulers, file transfers, and resource provisioning services. Given the heterogeneity of distributed infrastructures, SAGA provides a much needed interoperability layer that lowers the complexity and improves the simplicity of using distributed infrastructure whilst enhancing the sustainability of distributed applications, services, and tools.

SAGA is an Open Grid Forum (OGF) recognized standard (GFD.90). It allows developers of distributed applications to construct higher-level functionality and abstractions, such as gateways, workflows, application management systems, and run-time environments. The key advantages to running with SAGA on XSEDE is that users do not need to worry about the individual batch queueing systems implemented on the various machines. Using the SAGA API and appropriate job adaptors, the different submission mechanisms for these queueing systems is handled on the SAGA backend, which is transparent to the user.

The SAGA API has been used to provide almost complete coverage over nearly all grid and distributed computing middleware/systems, including but not limited to Condor, Genesis, Globus, UNICORE, SGE, LSF/PBS/Torque, and Amazon EC2.

3.3 Deployment of BigJob

SAGA and BigJob are lightweight enough that they can be easily installed into the home directory of a user using the Python Package Index (PyPi). SAGA is packaged within BigJob, so users do not have to worry about installing two separate modules.

Since the main deployment is on XSEDE, we do not recommend altering the default PYTHONPATH of the machines. Instead, we encourage users to use a virtual environment. A virtual environment allows a user to create a local Python software repository in his or her home directory that behaves exactly like the global Python repository, except that it grants the user write access to it. In order to use the virtual environment, the Python version must be greater than 2.4. Since some XSEDE machines use Python 2.4 as the default python version, it may be required to load a python module file before installing BigJob.

After activating the virtual environment, the BigJob python package can be installed by typing:

easy_install bigjob

In addition to the BigJob package, the BigJob python dependencies, including the SAGA package, are also installed. The SAGA package includes the proper adaptors for a wide variety of middleware systems. This allows the user to submit jobs to any of the XSEDE batch queuing systems.

BigJob requires SSH password-less login to the machines and a redis server running either locally or on a remote server. The redis server is used for coordination of the pilotjob and its compute and data units. For the purposes of this project, we utilize a private redis server hosted on a virtual machine at Indiana University. In order to provide a more seamless uptake of BigJob by users, we will provide an open-access redis server available on XSEDE. This avoids the overhead of new users having to start a redisserver on an XSEDE machine's head node or on their local machines. This effort is currently underway with XSEDE ECSS staff to make the server only accessible to registered users of XSEDE.

After following the aforementioned steps, users will be able to write their own BigJob submission scripts using Python. These scripts can range from simple ensemble-based simulations to more complicated workflows based on the users' needs.

4. COMPUTATIONAL WORKFLOW

4.1 Overview

This particular study involves the molecular dynamics (MD) simulation of 105 nucleosomes representing 21 locations along 5 different chromosomes. The targeted simulation time is 20 nanoseconds for each system, preceded by a minimization and equilibration phase, but the simulation phase is broken into 20 sequential 1-nanosecond tasks, where the output (atomic coordinates and velocities, and extended cell dimensions) of one task becomes input to the next. So it's a chained sequence, with each task being dependent upon successful completion and output of the previous

The data (both inputs and outputs) for these systems are pre-organized in a simple hierarchy with a common base directory. The first tier represents the 5 chromosomes, and the

second tier represents the 21 locations along the DNA strand representing the start of the nucleosome. All 20 chained simulations for the same system (chromosome/location) share the same directory, using a file-naming convention that reflects the task sequence (dyn0 - dyn20), where files corresponding to the same dynamic step (task) share the same file prefix. Each file-type is represented by its own extension (.coor, .vel, .xsc, .dcd, .dvd, .xst). Some inputs, such as the forcefield parameters (sys.parm) and atom connectivity information (sys.pdb), are common to all tasks within a chained sequence and needn't be duplicated or sequenced. Yet other inputs, such as the configuration (.conf) files, are the same for all systems, but they differ from step to step. These configuration files are stored in a separate subdirectory off the base directory. The following is a brief schematic of the file organization.

	-dyn-conf-files	[NAMD configuration files common to all systems]
	dyn1.conf	[NAMD configuration file for first dynamic step]
	dyn2.conf	
	dyn20.conf	
	I-chr02	[1st of 5 chromosomes]
	0068270	[1st of 21 nucleosome locations on 1st chromosome]
	sys.crd	[Parameter files common to all 20 dynamic steps]
	sys.parm	
	min-eq.coor	[Input to 1st dynamic step]
	min-eq.vel	
	min-eq.xsc	
4	dyn1.coor	[Output of 1st dynamic step, input to next]
K	dyn1.vel	
	dyn1.xsc	Y
7	dyn1.out	[Output-only files]
	dyn1.err	
	dyn1.xst	
	Idyn1.dcd	
	dyn1.dvd	
	dyn20.out	[Output of last dynamic step]
	: I0068271	[2nd of 21 nucleosome locations]
	10008271	[ZNG OI ZI NGCIEOSOME IOCATIONS]
	I0068290	[Last of 21 nucleosome locations on 1st chromosome]
	:	[Baby of 11 Macroscome Todations on 180 chromosome]
۱	-chr04	[2nd of 5 chromosomes]
	-chr16	[Last of 5 chromosomes]

Directory structure showing organization of input and output files.

- Input to step X (1-20):
 - dynX.conf: same for all systems, one for each nanosecond MD step, kept in a separate directory
 - dynX-1.coor, dynX-1.vel, dynX-1.xsc: passed
 from previous step
 - sys.pdb, sys.crd, sys.parm: same for all simulations on the same system
- Output from step X:
 - dynX.dcd, dynX.cvd, dynX.xst: trajectory data, used for further analysis
 - dynX.coor, dynX.vel, dynX.xsc: passed to next
 step
 - dynX.out, dynX.err: stdout, stderr (primarily diagnostic information)

To help automate the workflow under BigJob (particularly for data staging with Pilot-Data), a naming convention had to be assumed for certain file-prefix keywords within the

dynX.conf files, since these files can not in general be inspected (unless they are local) during a Pilot-Job's run-time to extract the filenames and orchestrate the file staging. So the assumption is that input file prefix within a dynX.conf file would be set to 'dynX-1' and that the output file prefix would be set to 'dynX'. The only exception was for the first step (dyn1), where the input file prefix is assumed to be 'min-eq'.

4.2 Planned Approach

The initial plan was to use the Python-based BigJob API (Pilot-Job, Pilot-Compute, and Pilot-Data) based on the Python-based SAGA framework (called *Bliss*) to develop a BigJob script that would orchestrate an ensemble of tasks as one or more batch pilot jobs on one or more XSEDE resources (like Lonestar or Kraken) using several internal sub-queues to distribute independent tasks in a round-robin fashion, while also incorporating a shared workflow mechanism to help manage the dependent tasks, deferring their submission and staging files from output Data-Units to input Data-Units as necessary and as depicted in Figure 2.

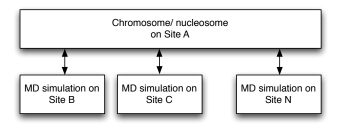


Figure 2: Orchestration of distributed resources using Pilot-Compute and Pilot-Data.

The workflow was to be defined in an external *Config* file, independent of the BigJob script, and read by the BigJob script. This workflow *Config* file would be generated semi-automatically based on the inspection of output files from prior BigJob runs.

The initial goal was to make use of the Pilot-Data API to stage data at run-time to and from a remote source, where the data originally resided and where further post-processing was to be done. However, an alternate approach depending on locally accessible files was also to be explored and evaluated against the remote staging approach.

4.3 Implementation

The actual implementation went mostly as planned, including the development of a BigJob script (namd_bigwork.py) in Python, using an INI style Config file to hold both BigJob configuration settings and the workflow definition, and a shell script (gen_workflow) was also written to generate the workflow section of the Config file. Each of these will be briefly described, starting with the format and generation of the workflow definition component.

Workflow defined using INI style Config file

Below is an example of a WORKFLOW section of the *Config* file. The WORKFLOW section contains subsections for the RE-MOTE_DATA_RESOURCE, DEFAULTS, and the TASKS. The TASKS section contains subsections for each task. Each task has a unique label, entries to help locate the various files, and a list of prerequisite tasks (DEPENDENCIES) identified by their la-

bels. Most entries have default values if not specified. The *Config* entries are parsed and converted into a *Dictionary* object within the Python script using the ConfigObj library (favored over the built-in ConfigParser library).

```
[WORKFLOW]

[[REMOTE_DATA_RESOURCE]]

PROTOCOL = "ssh"

USER = "jacks"

MACHINE = "lonestar"

DATACENTER = "tacc.utexas.edu"

BASEPATH = "vscratch/02059/jsolow"

CONF_DIR = "/scratch/02059/jsolow/dyn-conf-files"

PARM_DIR =

[[DEFAULTS]]

[[TASKS]]

[[[chr04-1357521-4]]]

DATAPATH = "chr04/1357521"

CONF = "dyn4"

PARM =

IN =

OUT =

DEPENDENCIES = "chr04-1357521-3",

[[[chr04-1357521-5]]]

DATAPATH = "chr04/1357521"

CONF = "dyn5"
```

Workflow section of a *Config* file, including configuration settings for the remote data source, showing two tasks, one depending on the other.

Script to generate workflow Config file - gen_workflow

To help automate the generation of the workflow definition, a shell script was written to traverse the directory tree containing all the input and output files looking for *signatures* that a simulation was completed and skipping it as a task. For this particular study, these *signatures* included:

- dynX.out file contains "WallClock" in one of the last two lines
- dynX.dcd and dynX.dvd files are greater than 1.8GB (proportional to the size of the dynX.coor file and the number of dynamics steps)

The script uses the directory path (chromosome/location) and the dynamic step (1-20) to generate the task label. It assumes that the task is dependent upon on the dynamic step that precedes it, X-1, except for the first step, which assumes $\min-eq.*$ as its input. The base paths for the data tree, the NAMD configuration files, and parameter files are set in the REMOTE_DATA_RESOURCE section.

The script takes as arguments the range of dynamics steps to include in its traversal, with the default of 1-20. Restarting an incomplete run is usually just a matter of rerunning the gen_workflow script, which automatically omits the completed tasks.

The output of this script can be appended to the main Config file or kept as a separate file and referenced by the FILE keyword in the WORKFLOW section. The latter is the cleaner and preferred method.

'NAMD BigWork' Python script - namd_bigwork.py

The primary deliverable of this exercise is a Python script (namd_bigwork.py) that uses the BigJob framework to launch a batch pilot job and help orchestrate the submission of an ensemble of NAMD MD simulations (a collection of independent and dependent tasks) to a single batch job instance

with a significant reservation of resources for maximum high-throughput performance. In this particular study, that reservation of resources is 2400 cores on Lonestar (@ TACC) for a 24-hour period, sub-divided internally into 10 sub-queues, to handle 200+ simulations.

The BigJob script sets up a Pilot-Job with both Pilot-Compute and Pilot-Data components, with the latter only used if the data resource is remote from the compute resource. If the data is local to the computer resource, no data staging is done, and the Compute-Unit is directed to use the input data directory as the working directory, which is also the destination directory for the output and where the next dynamic step expects its input data to be.

If the data is remote, then staging is needed to get the data to and from the Compute-Unit's temporary working directory. Three Data-Units are used to stage the data: an input Data-Unit, and output Data-Unit, and a chained Data-Unit. The chained Data-Unit is used to selectively redirect output data from one task to another dependent task as input. The Data-Unit-to-Data-Unit transfer is generally done by reference only and no data is actually moved. However, the input Data-Unit does need to copy data from the remote resource to the local input Data-Unit and then to the Compute-Unit's working directory, using SCP or SFTP for the former (determined by the SAGA-BigJob framework) and possibly symbolic links for the latter. The output data is shuttled from the Compute-Unit's working directory back to the output Data-Unit, where it remains until the completion of the Pilot-Job. It then becomes the responsibility of the user to rummage through the contents of the cryptic Data-Unit directories and move them to their final destination.

The introduction of workflow (task) management to a BigJob script is the primary contribution from this effort [NAMD + BigJob + Workflow = 'NAMD BigWork']. The task workflow is defined the in the Config file (namd_bigwork.conf by default). The WORKFLOW section of the Config file is parsed and converted to a hierarchy of Python Dictionary structures for easy processing. The main construct is an array of task Dictionary elements that contain all the information about a particular task, including it's current status, which goes from WAITING to PENDING (if it has unsatisfied dependencies) to New to Unknown to Running to Done to COMPLETED or FAILED as it progresses along. The script continually loops through the tasks looking for changes in Compute-Unit and Data-Unit (if data staging) statuses and releasing PENDING tasks when their dependencies are satisfied. Any task that is in a WAITING (not PENDING) state is set up for submission to the Pilot-Job's agent. This set up involves the creation of the input, output, and chained Data-Units (if data staging) and a Compute-Unit. If data staging is being done, any requisite chained Data-Units are transferred to the input Data-Unit. Once all the Data-Units are processed, the Compute-Unit is submitted.

The Pilot-Job is generally a batch job submitted to the system's batch queuing system (like PBS, SGE, or SLURM), and it can sit in the system queue for an indefinite period of time. Meanwhile, the Pilot-Job agent queues up Compute-Units for execution in its internal queuing system as sub-jobs. Each sub-job will have its own sub-allocation of cores from the total allocation given to the Pilot-Job. Once the batch job (Pilot-Job) starts running, the backlog of sub-jobs will begin to get spawned as cores become available. In this

particular study, there were 2400 cores distributed among 10 sub-jobs, 240 cores each. Once a sub-job is completed, those cores become available for another sub-job held in the Pilot-Job's internal queue. When there are no remaining tasks (sub-jobs) to submit, the Pilot-Compute, Pilot-Data, and Pilot-Job agents are all gracefully shutdown and the batch job terminated.

'NAMD BigWork' log and status files

To help monitor, perform diagnostics, and do some post analytics (performance measures) on all the tasks, both a running log and a snapshot status file are generated during the run. The names of the log and status files are prefixed by the *Config* file name, followed by the a date and PID of the script, and suffixed with .log and .status, respectively.

That status file is written out with every state change showing details of each task organized by state, as seen in the snippet below. It contains the rather cryptic Compute-Unit and Data-Unit directory names to help map them to specific tasks. It also contains the current run time for each task since it entered the Running) state.

The running (accumulative) log file contains every task status change and the details of each submission, followed by a summary (tally by state) with time-stamps, as seen in the snippet below. The overall level of detail for logging to stdout is set in a BigJob configuration file (bigjob.conf) or by an environment variable (BIGJOB_VERBOSE), but the written log file generally only contains INFO level output, although it can be overriden up to the overall logging level.

Filtering the log file (with grep, for example) can generate individual task summaries, such as the example below for task chr16-0503364-13, which has no dependencies, or another example for task chr16-0503364-14, which depends on the completion (and the output) of the previous task. Filtering for the state summaries gives a trace of the task throughput, which can be plotted against the time-stamps for a visual summary like that in Figure 3.

'NAMD BigWork' Config file

The BigJob script is generally free of specifics about the compute and data resources, details of the NAMD executable, or of any workflow details. All that is externalized to the 'NAMD BigWork' Config file (namd_bigwork.conf by default). The Config file is broken into sections - PILOT, NAMD, and WORKFLOW. The WORKFLOW section was described earlier, and its content can be further externalized using the FILE keyword to reference a separate Config file for just the workflow details. A snippet of an example Config file is given below.

4.4 Results

Typical run

A typical run for this study was a 24-hour 2400-core job broken into 10 240-core sub-job queues (within the Pilot-Job), which handled 210 MD simulations, representing 2 1-nanosecond simulations for each of 105 systems (21 locations along 5 chromosomes). Each simulation was 500,000 MD timesteps and took about 1 hour on 240 cores (on Lonestar). 2400 cores represents almost 10% of Lonestar's 28,800 cores.

Figure 3 shows a typical run profile for an ensemble of sub-jobs progressing through their states within the Pilot-Job. Note that BigJob tends to use a LIFO-like queuing

```
Task status as of 2013-03-05 05:48:36.093657
0 Tasks still WAITING to be processed
75 Tasks PENDING dependent tasks to be completed
chr02-0068270-14 | Waiting for ['chr02-0068270-13']
chr02-0068271-14 | Waiting for ['chr02-0068271-13']
:
28 Tasks submitted to queue (New) and waiting to run
chr02-0068271-13 | Compute Unit: sj-fd97230c-8572-1162-8f2a-f04da2004b3c
chr02-0068271-13 | Compute Unit: sj-fd97230c-8572-1162-8f2a-f04da2004b3c
:
67 Tasks submitted but changed to Unknown status
chr02-0068272-14 | Compute Unit: sj-fd97230c-8572-1162-8f2a-f04da2004b3c
chr02-0068272-14 | Compute Unit: sj-856fac12-8572-1162-8f2a-f04da2004b3c
:
10 Tasks currently Running
chr04-1357530-13 | Run time: 303 | Compute Unit: sj-52c5da08-8573-1162-8f2a-f04da2004b3c
chr04-1357533-13 | Run time: 324 | Compute Unit: sj-540a0084-8573-1162-8f2a-f04da2004b3c
:
0 Tasks Done and waiting to complete data export
30 Tasks OMPLETED
chr02-0068272-13 | Run time: 3297 | Compute Unit: sj-076340f0-8573-1162-8f2a-f04da2004b3c
chr02-0068274-13 | Run time: 3296 | Compute Unit: sj-0b6d947a-8573-1162-8f2a-f04da2004b3c
chr02-0068274-13 | Run time: 3296 | Compute Unit: sj-0b6d947a-8573-1162-8f2a-f04da2004b3c
```

'NAMD BigWork' status file - a snapshot in time showing details of each task organized by state

```
WAITING: 1 | PENDING: 75 | New: 28 | Unknown: 66 | Running: 10 | Done: 0 | COMPLETED: 30 | FAILED: 0 | Time: 2013-03-05 05:48:10.288700
Total tasks remaining to complete: 180 of 210 at 2013-03-05 05:48:10.288912
Preparing Task chr02-0068285-14 for submission
Compute Unit for Task chr02-0068285-14 submitted to Compute Data Service
Compute Unit status for Task chr02-0068285-14 changed to New at 2013-03-05 05:48:10.290884
WAITING: 0 | PENDING: 75 | New: 29 | Unknown 66 | Running: 10 | Done: 0 | COMPLETED: 30 | FAILED: 0 | Time: 2013-03-05 05:48:18.155282
Total tasks remaining to complete: 180 of 210 at 2013-03-05 05:48:18.155496
Compute Unit status for Task chr02-0068285-14 changed to Unknown at 2013-03-05 05:48:36.093026
WAITING: 0 | PENDING: 75 | New: 28 | Unknown: 67 | Running: 10 | Done: 0 | COMPLETED: 30 | FAILED: 0 | Time: 2013-03-05 05:48:43.793960
Total tasks remaining to complete: 180 of 210 at 2013-03-05 05:48:43.794202
```

'NAMD BigWork' log file - running log of task status changes and sub-job submission details, with a task state summary after each status change, including time-stamps.

```
Preparing Task chr16-0503364-13 for submission
Compute Unit for Task chr16-0503364-13 submitted to Compute Data Service
Compute Unit status for Task chr16-0503364-13 changed to New at 2013-03-04 12:28:22.006905
Prerequisite Task (chr16-0503364-13) found for Task chr16-0503364-14
Compute Unit status for Task chr16-0503364-13 changed to Unknown at 2013-03-05 02:59:48.492535
Compute Unit status for Task chr16-0503364-13 changed to Running at 2013-03-05 03:54:21.593379
Compute Unit status for Task chr16-0503364-13 changed to Done at 2013-03-05 04:49:42.614543
Compute Unit completed for Task chr16-0503364-13
```

Extraction from log file showing history of task chr16-0503364-13, which has no dependencies.

```
Prerequisite Task (chri6-0503364-13) found for Task chri6-0503364-14

Compute Unit status for Task chri6-0503364-14 changed to PENDING at 2013-03-04 12:28:22.013608

All prerequisite tasks completed for Task chri6-0503364-14

Compute Unit status for Task chri6-0503364-14 changed to WAITING (from PENDING) at 2013-03-05 04:50:53.570544

Preparing Task chri6-0503364-14 for submission

Compute Unit for Task chri6-0503364-14 submisted to Compute Data Service

Compute Unit status for Task chri6-0503364-14 changed to New at 2013-03-05 04:51:01.186614

Compute Unit status for Task chri6-0503364-14 changed to Unknown at 2013-03-05 04:51:27.003795
```

Extraction from log file showing history of task chr16-0503364-14, which depends on the completion of chr16-0503364-14.

```
WAITING: 0 | PENDING: 78 | New: 29 | Unknown: 63 | Running: 10 | Done: 2 | COMPLETED: 28 | FAILED: 0 | Time: 2013-03-05 05:46:58.886167
WAITING: 0 | PENDING: 77 | New: 28 | Unknown: 64 | Running: 10 | Done: 2 | COMPLETED: 28 | FAILED: 0 | Time: 2013-03-05 05:47:06.513090
WAITING: 0 | PENDING: 77 | New: 29 | Unknown: 64 | Running: 10 | Done: 2 | COMPLETED: 28 | FAILED: 0 | Time: 2013-03-05 05:47:16.520071
WAITING: 0 | PENDING: 77 | New: 29 | Unknown: 64 | Running: 10 | Done: 1 | COMPLETED: 29 | FAILED: 0 | Time: 2013-03-05 05:47:12.2.220978
WAITING: 1 | PENDING: 76 | New: 29 | Unknown: 65 | Running: 10 | Done: 1 | COMPLETED: 29 | FAILED: 0 | Time: 2013-03-05 05:47:30.113406
WAITING: 0 | PENDING: 76 | New: 29 | Unknown: 65 | Running: 10 | Done: 1 | COMPLETED: 29 | FAILED: 0 | Time: 2013-03-05 05:47:30.113406
WAITING: 0 | PENDING: 76 | New: 28 | Unknown: 66 | Running: 10 | Done: 1 | COMPLETED: 29 | FAILED: 0 | Time: 2013-03-05 05:48:02.363673
WAITING: 0 | PENDING: 75 | New: 28 | Unknown: 66 | Running: 10 | Done: 0 | COMPLETED: 30 | FAILED: 0 | Time: 2013-03-05 05:48:10.288700
WAITING: 0 | PENDING: 75 | New: 28 | Unknown: 66 | Running: 10 | Done: 0 | COMPLETED: 30 | FAILED: 0 | Time: 2013-03-05 05:48:10.288700
WAITING: 0 | PENDING: 75 | New: 28 | Unknown: 66 | Running: 10 | Done: 0 | COMPLETED: 30 | FAILED: 0 | Time: 2013-03-05 05:48:10.288700
WAITING: 0 | PENDING: 75 | New: 28 | Unknown: 67 | Running: 10 | Done: 0 | COMPLETED: 30 | FAILED: 0 | Time: 2013-03-05 05:48:10.288700
```

Extraction from log file showing task state summaries, including time-stamps, that can be used for time-series plots like the one in Figure 3.

```
[PILOT]

COURDINATION_URL = "redis://localhost:6379"

COMPUTE_USER = "jacks"

COMPUTE_DATACENTER = "tacc.utexas.edu"

COMPUTE_DATACENTER = "tacc.utexas.edu"

COMPUTE_SENVICE_PROTOCOL = "sge+ssh"

COMPUTE_ACCOUNT = "TG-MCSIOO111"

COMPUTE_QUEUE = "development"

DATA_USER = "jacks"

DATA_MACHINE = "lonestar"

DATA_SERVICE_PROTOCOL = "ssh"

DATA_SIZE = 100

SUBJOBS = 2

PROCESSORS_PER_JOB = 48

PROCESSORS_PER_DOB = 12

[NAMD]

ROOT_DIR = "/home1/00288/tg455591/NAMD_2.9"

BIM_DIR = "NAMD_2.9_Linux_x86_64-MVAPICH-Intel-Lonestar"

EXEC = "namd2"

[VORRFLOW]

FILE = "namd_workflow_jsoloe.conf"
```

'NAMD BigWork' Config file showing the PILOT and NAMD sections and an external FILE reference for the WORKFLOW section.

mechanism, as the dependent tasks get interlaced early in the queue even though they were submitted much later (after their prerequisite tasks were completed).

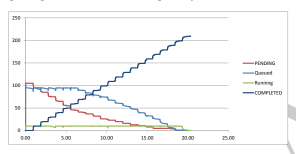


Figure 3: Progression of 210 sub-jobs (number in each state) over time (hrs)

Figure 4 shows the progression of the last seven BigJob runs over a ten day period. The typical BigJob completed all 210 tasks assigned to it. Exceptions are the first and last two BigJobs. The first BigJob was only assigned 185 (the other 25 having been completed in an earlier run). The next to last BigJob completed only 209 sub-jobs. The failed sub-job was then automatically added to the last BigJob run. Careful inspection of the running simulations indicates that during the last hour of each BigJob less than 10 sub-jobs might be running. For example, the number of running jobs for the next to last BigJob do not abruptly drop to zero at the end of day 69. It turns out that that sub-job failed to run due to an mpiexec problem and was never detected by the Pilot-Job, hence left to run out the clock - essentially waiting for a "dead" sub-job to complete.

Summary

The total study took about 11 runs to cover the full 20 nanoseconds of simulation for each system plus an initial minimization-equilibration step, for a total of 2310 simulations. At 240 cores for 1 hour each, that's about 550,000 SUs, or 55,000 SUs per run (per day).

There were a few runs that did not complete for various reasons (see *Issues* below) and had to be restarted using a regenerated workflow *Config* file.

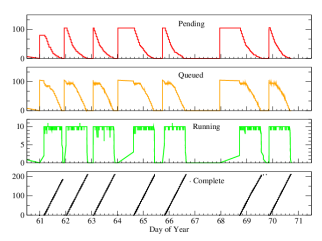


Figure 4: Progression of the last seven BigJob runs over a ten day period.

Most simulations ran in just under an hour, but there were some anomalies (outliers). Figure 5 shows the statistical distribution (histogram) of run-times (wall times) for all 2100 NAMD simulations. The shortest run-time was 54 minutes with 2027 tasks completing in under 1 hour. A total of eight simulations required longer than 2 hours to successfully complete. The longest single task (for a 1-nanosecond simulation) was 8 hours long. The mean run-time for all simulations was 57 minutes. If the simulations that required longer than 2 hours to run are excluded, the variation in run-times is less than 4 minutes or less than 6% of the expected run time. These results indicate that BigJob could proactively kill any sub-job taking longer than 1.5 hrs and resubmit it with the assumption being that the sub-job has encountered a hardware or system-level failure. When sub-jobs with anomalous run-times are manually rerun they complete within the expected time frame.

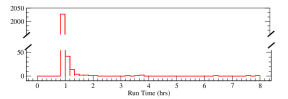


Figure 5: Histogram of run-times for the 2100 NAMD simulation tasks.

5. DISCUSSION: EXPERIENCE AND CON-CLUSION

5.1 Issues

This experiment with BigJob and the integration of a workflow was not without its issues. The following highlight some of those issues.

One of the first challenges was to identify the right NAMD executable to use within the BigJob framework. NAMD is usually run via the charmrun frontend, which handles all the MPI details. Furthermore, the executable is usually built with the Charm++ pre-processor/compiler/library, which interacts with the charmrun frontend. So it is not compat-

ible with mpirun, which is used by BigJob. Fortunately, the NAMD developers provide a builder for an MVAPICH version of the NAMD executable, which appears to work appropriately under the BigJob environment.

The next big surprise was to discover that data staging only works one way, from the remote data resource to compute resource, and not the other way. Getting data back to the remote data source is expected to be done external to BigJob. One can export to the local system (where the BigJob script is running), but unless that is the original data source, it is of limited value. The workflow in the study requires a good deal of data shuffling between dependent tasks, and the directory structure is non-trivial, so an alternate scheme needed to be explored, at least temporarily. The temporary solution was to copy all the data to the compute resource (to scratch space on Lonestar in this case) and run everything local. As output was generated, it was rsync-ed back to the original source (in this case, a workstation at LA Tech) on a scheduled basis using cron. With the data and compute resources both being local, there is no need for data staging, and the working directory can be set to where the data resides - and run in situ. The volume of data requires that scratch storage be used for the data, so long-term integrity of the data of the course of the study was a potential issue. It was important to keep all the data and its directory structure intact to help automate generation of the workflow Config file.

In an effort to work within the constraints of BigJob regarding output data staging (back to its remote origin), another issue arose. It appears that SCP has issues with very large files. The trajectory files (.dcd and .dvd) for this study are around 1.8GB each, and it was discovered (after many SUs were consumed and scratch files purged) that they were getting truncated to about 500MB (without any warning or error messages) while they were being copied from the working directory to the Data-Unit store (or even when being exported locally). So, any further data staging was abandoned until this issue could be resolved. However, it was demonstrated that selective Data-Unit-to-Data-Unit transfers (via a "chained" Data-Unit) can dramatically reduce the amount of data transfer (copying) that needs to take place for chained tasks. Sync-ing the data back to the original remote source is still a troubling issue, and the cryptic directory naming and flat directory structure used by BigJob don't help.

Going to an all-local solution avoids many issues, but it also presents a few subtle issues of its own. For example, if the NAMD *Config* file is not in the working directory (which they are not in this study), one must force NAMD to ignore its location as the default location for the input files. This was done by adding a "CWD getenv(\$PWD)" entry to the NAMD *Config* file to force it to look in the current working directory.

Not only is exporting output data back to a remote resource the user's responsibility, but so is cleaning up afterward. The Compute-Unit's working directories and the Data-Unit directories can tend to grow rather quickly and need to be cleaned up by the user. One can sweep this issue under the rug by using scratch or temporary space, but it would be better citizenship to clean up manually if it can't be done automatically or programmatically within the BigJob script. Ideally, BigJob would support both remote exporting and the purging of working and staging directories.

Although most of BigJob's work is offloaded to a batch Pilot-Job, the launch script remains a key component in orchestrating the complex workflow, and often that script is launched on the head (login) node of the compute resource. If care isn't taken to throttle some of its activities it becomes prone to be canceled by the system's administrator. This project experienced that first hand and put nearly 50,000 SUs at risk by having the batch Pilot-Job sit idle until the maximum walltime was exceeded. So care must be taken to throttle tight loops (using strategically placed sleep() calls) and minimize CPU usage if the script is being launched on a limited or controlled resource like a login node.

On the flip-side, if a the batch Pilot-Job should somehow be aborted or terminated (e.g., after a series of mpiexec problems, which happened more than once during this study), the launch script can be left running unaware. In one instance a single sub-job got hung with some mpiexec problem and the Pilot-Job was unaware, causing the Pilot-Job to run with all cores idle until the walltime was exceeded. It would be helpful if the $\mathtt{get_state}()$ for the Pilot-Job could detect such events or allow user-configured criteria to terminate the batch job (e.g., terminate if X% of the cores are idle for more than Y minutes).

BigJob uses an Advert service (usually a Redis server) to communicate and transfer information between various distributed components of the BigJob framework. The status of the Advert service is usually checked upon initiation of the BigJob environment; however, if the Avert service shuts down later, the Pilot-Job batch job can abort and leave the BigJob launch script running unaware of the problem. Again, it would be helpful to be able to detect such events and shut everything down gracefully.

Much of the system-dependent details of working in a distributed heterogeneous environment are handled by using SAGA and the BigJob framework as middleware, and much of that is done using somewhat generic adapters; however, various subtle and system specific details often remain that need to be managed or configured by the user. This can defeat much of the transparency intended by SAGA and BigJob. For example, a user shouldn't have to know what particular batch queuing system a compute resource uses (Torque/PBS, SGE, SLUM,...) or whether to use SSH or GSISSH or fork. System-specific configuration files should be able to handle such nuances. He/she shouldn't have to completely overhaul his/her strategy based on whether the data is local to the compute resource. Data staging ought to be smart enough to figure that out. A global file system model/view or abstraction might be helpful - where everything looks local.

The MPI environment can vary between systems, especially when they're adapted to take advantage of certain architectures or interconnect fabric. The use of aprun on Kraken and ibrun on TACC machines in place of mpirun are good examples. The use of charmrun for NAMD is an even more extreme example. These are tough issues to deal with even for a veteran MPI user, let alone for someone trying to be shielded from such details and nuances. SAGA/BigJob needs to develop some generic wrapper mechanism and/or configuration file to mask this level of detail.

One final issue, which is more of a workflow management issue than a BigJob issue, is the potentially inefficient use of cores at the end of a run, when sub-queues aren't being kept full, leaving more and more cores idle as it approaches the end. Either the workflow manager (the BigJob script) needs to allocate more cores to tasks (sub-jobs) near the end of a run, or the Pilot-Job needs to be configurable to not accept any more sub-jobs if it can't utilize a certain portion of its available resources. This, of course, is very workflow dependent and probably shouldn't rely on BigJob for anything but hints about resource utilization.

5.2 Future

This was just the first of a series (hopefully) of experiments with BigJob and distributing large ensembles of chained NAMD simulations across a heterogeneous mix of compute and data resources like those on XSEDE. The following are a few things suggested for future work.

The INI-style Config file format is rather limited in its ability to express scientific workflow. Workflow management theories often use directed acyclic graphs (DAG) to express workflow, where nodes are tasks and edges represent dependencies and/or data flow. There are now markup languages to express DAGs, such as DAX (DAG in XML), used by Pegasus [] [http://pegasus.isi.edu/projects/pegasus] and other workflow management systems. There are even graphical programs like Triana [] [http://www.trianacode.org] that generate DAX. Adopting a language/format like DAX would help broaden the scope of workflow management and allow some degree of plug-and-play with other workflow management tools.

The original goal was to demonstrate a more distributed example of using the BigJob framework, with multiple Pilot-Jobs running on different compute resources, data being staged from yet another resource (and back), and all controlled by a single workflow management front-end. Although each aspect was demonstrated independently, with a few caveats, the chore of tying it all together remains.

One thing that was not fully explored was the interlacing of data staging with compute processing to avoid blocking while large data files are being moved around. This will require a better understanding of the states and their transitions within the Pilot-Data framework to help eliminate the blocking wait() calls.

The workflow (task) management needs to better address the "end game" by re-allocating cores (reducing the number of sub-queues and redistributing allotted cores) near the end of a run to avoid idling resources as the number of tasks (sub-jobs) drops below the number of available sub-queues. Not all tasks scale well, however, so this is a non-trivial matter. More is not always better, and being busy isn't the same as being productive - even for computers.

Many of the issues faced in this exercise were due to lack of monitor and control points among the various components involved. Exposing more of the underlying Pilot-API could help trap and respond to some of these events inside a BigJob script. However, exposing more of the Pilot-API (and the some of the other APIs, like to the Advert service) to the command language interface (CLI) could provide a separate external layer of monitoring and control that would help dynamically manage sub-jobs and tweak the workflow as needed. In that regard, building a GUI client for the CLI could be a valuable tool, especially if it's portable across platforms and robust to connection loss and restoration - a REST-based interface, perhaps, that runs in a web browser. These are obviously outside the scope of the end-user of BigJob, but some food for thought for the BigJob develop-

ment team.

5.3 Conclusions

To date less than ten molecular dynamics studies of the nucleosome have been reported. All except a control study performed by this group used nominally the same sequence of DNA. The longest all-atom trajectory reported for a nucleosome is 500ns. By using BigJob to marshall resources distributed across XSEDE we have achieved a sampling of nucleosome sequence variability and dynamics that is over two orders of magnitude higher than any previous effort. We note that the "experiments" reported here were actually production runs for our nucleosome study that successfully completed. Successful completion alone supports what is known biologically: that any sequence of DNA can be wrapped around a histone core to form a nucleosome, albeit with varying affinities. Achieving the level of throughput reported here is a necessary first step in dissecting DNA sequence specific properties of nucleosomes in atomic detail which, because of the fundamental role of nucleosomes, impacts our understanding of virtually all genetic processes.

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