Subject Section

The application of Hadoop in Structural Bioinformatics

Jamie Alnasir 1,*, Hugh P. Shanahan 2

1 Institute of Cancer Research, 123 Old Brompton Road, London, SW7 3RP, United Kingdom and
2 Department of Computer Science, Royal Holloway, University of London, Egham, Surrey, TW20 0EX, United Kingdom.

* To whom correspondence should be addressed.

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The paper reviews the use of the Hadoop platform in Structural Bioinformatics applications. For Structural Bioinformatics, Hadoop provides a new framework to analyse large fractions of the Protein Data Bank that is key for high throughput studies of (for example) protein-ligand docking, clustering of protein-ligand complexes and structural alignment. Specifically we review in the literature a number of implementations using Hadoop of high-throughput analyses and their scalability. We find that these deployments for the most part use known executables called from MapReduce rather than rewriting the algorithms. The scalability exhibits a variable behaviour in comparison with other batch schedulers, particularly as direct comparisons on the same platform are generally not available. Direct comparisons of Hadoop with batch schedulers are absent in the literature but we note there is some evidence that MPI implementations scale better than Hadoop. A significant barrier to the use of the Hadoop ecosystem is the difficulty of the interface and configuration of a resource to use Hadoop. This will improve over time as interfaces to Hadoop e.g. Spark improve, usage of cloud platforms (e.g. Azure and AWS) increases and standardised approaches such as Workflow Languages (i.e. WDL, CWL, Nextflow) are taken up.

Keywords: Structural Bioinformatics, Hadoop, Cloud Computing.

1 Introduction

The Apache Hadoop project [1] is a software ecosystem i.e. a collection of interrelated, interacting projects forming a common technological platform [2] for analysing large data sets. Hadoop presents three potential advantages for the analysis of large Biological data sets. In the first instance, it is designed for the analysis of large semi-structured data sets; secondly it is designed to be fault tolerant (in essence by ensuring a very large amount of overall redundancy) which becomes almost inevitable for sufficiently large numbers of processors; finally the MapReduce formalism for describing the data sets allows for the easy construction of work flows.

Given the continued importance of computationally analysing protein structures in fields such as Membrane proteins [3], protein-protein interactions and their impact in Systems Biology [4, 5] and protein engineering [6] a platform for easily analysing semi-structured data sets such as those found in the Protein Data Bank [7] on a large would be an important step forward.

On the other hand, Hadoop also presents barriers to its adoption within the community for Bioinformatics and the analysis of structural data. In the first instance Hadoop runs as a series of Java libraries and hence there is a learning curve for any Bioinformatician or Structural Biologist who hasn’t used Java before, though we note that more recent additions to the Hadoop ecosystem such as Spark [8] have a wider range of languages (e.g. Python, Scala and R). Correspondingly, unlike data parallel languages such as High Performance Fortran [9], Hadoop cannot be easily retro-fitted into a stable code base even if the original code is written in Java though it is possible to use Hadoop to deploy instances of executables in the same way that batch schedulers do. Finally, implementing Hadoop (and therefore Spark) on a local cluster is not trivial and requires a significant level of expertise...
from the relevant systems administrator. As we note, this latter difficulty
is obviated on cloud platforms such as Azure and AWS [10].
This paper reviews the range of work that has been done on Hadoop in
Bioinformatics and in Structural Bioinformatics in particular. Specifically
this paper will determine how stable these platforms are in comparison
to other approaches such as batch schedulers and MPI (discussed in
sections 1.1.3 and 1.1.4 respectively). The rest of this paper is organised
as follows. In the first instance a brief overview of the Hadoop and Spark
systems as well as a description of batch schedulers and MPI. It then
describes the Hadoop formalism. A brief review of the application of
Hadoop in Bioinformatics is provided followed by an in-depth review
of the application of Hadoop in Structural Bioinformatics. Specifically we
examine the range of cases where Hadoop has been used, the scaling
behaviour of such systems in comparison with other platforms, where
available and its dependence on the internal configuration of the version
of Hadoop used. Finally this paper will draw some conclusions on the
scalability of Hadoop and its application in Structural Bioinformatics.

1.1 Distributed computing architectures

1.1.1 Hadoop and MapReduce

Apache Hadoop is a software framework for distributed processing and
storage which is typically installed on a Linux computer cluster to facilitate
large scale distributed data analysis, though it can be run on a standalone
single computer node usually for the purposes of prototyping. Hadoop
clusters may be built using commodity hardware, for instance off the shelf
equipment such as used in computer farms, and key features are fault-
tolerance and data-locality. In the former case scaling up a cluster to add
more machines and disks increases the probability of a failure occurring
and hence systems must have a built-in redundancy to compensate for it. In
the latter case data-locality provides the ability of the framework to execute
code on the same node or at least the same rack of the cluster as where
the input data resides. This reduces the amount of network traffic during
processing thereby avoiding network bottlenecks [11]. The fault-tolerance
and data-locality of Hadoop are made possible by its distributed file system
(HDFS) [12] and by Hadoop’s resource scheduler YARN (Yet Another Resource
Negotiator) [13] which is responsible for cluster management, in particular resource allocation and job scheduling.
The distributed data on HDFS is accessed programmatically using the
MapReduce formalism, originally implemented in Java. In this formalism
the distributed data accessed from HDFS is a set of tuples i.e. pairs of keys
and values \( \langle k_i, v_i \rangle \), \( 1 \leq i \leq N \) where \( N \) are the total number of data
entries. For example, the entries in the Protein Data Bank (PDB) would
be a set of pairs where \( k_i \) would be a specific PDB id and \( v_i \) could be a single
string with the corresponding PDB data in XML format. All operations are
based on these tuples, either creating new tuples (i.e. through a Map step)
or summarising the data stored in the tuples (i.e. through a Reduce step).
Extending the above example, via a Map step, a specific executable (e.g. a
docking program run with a specific small molecule) could be run on
each PDB entry to create a log file for each PDB entry. In the MapReduce
formalism this means creating a new set if tuples \( < k_i, l_i > \) where \( k_i \) is
again the PDB entry and \( l_i \) is a single string with the log file. A Reduce
step could then be applied on this second set of tuples to create a single
tuple which carries some specific set of summary data (e.g. how many
structures had a docking score greater than some threshold in the previous
Map step).

The rise of the use of Hadoop has mirrored the increasing use of cloud
platforms. MapReduce is offered as a Platform as a Service (PaaS) by all
of the major cloud-service providers (Amazon AWS, Google Cloud and
Microsoft Azure) [14].

1.1.2 Apache Spark

Spark is a cluster computing framework that can be used standalone or can
utilize Hadoop’s distributed file system (HDFS) and a resource scheduler
(often Apache YARN), providing an application programming interface
(API) for distributed computation. Spark is designed to overcome some
of the constraints of Apache Hadoop offering significant performance
improvements but keeping the fault-tolerance and scalability features of
Hadoop by utilising HDFS [15].

Although Spark supports MapReduce programs, the main constraint
Spark overcomes is Hadoop’s acyclic data flow model by allowing the re-
usability of intermediate data in the form of a data structure that is central
to Spark, the Resilient Distributed Dataset (RDD). The RDD serves as an
abstraction for distributed memory that allows in-memory computations
on large clusters in a fault-tolerant manner, and because an RDD is
partitioned across multiple compute nodes, can be rebuilt if for some reason
a partition is lost [16]. The RDD is an immutable (read-only) data structure
that can encapsulate objects from a number of different programming
languages and typically on Spark this is Python, Java, or Scala and in
contrast Hadoop development, notwithstanding that Hadoop MapReduce
programs can be written in any language supporting the UNIX POSIX
standard I/O streams, is predominantly Java based. This is significant from
a development perspective because Java requires programmers to possess
more specialist object-oriented programming (OOP) knowledge than for
example Python, and Java programs tend to have more dependencies on
runtime libraries.

An RDD is usually created in two ways, by referencing an external
data source, for instance a dataset or file on HDFS, or by parallelising
an existing Spark data structure. Parallelising a data structure, for instance an
array in Spark, allows it to be operated on in a parallel fashion whereby
Spark handles the caching of the RDD in memory across nodes of the
cluster. As a result of this, Spark MapReduce operations gain significant
performance enhancements over their Hadoop counterparts. This can be
achieved because Spark creates an execution plan, in the form of a directed
acyclic graph (DAG), of Spark and MapReduce operations to be performed
on RDDs. The execution plan models dependencies and allows Spark
to optimise execution of a job’s components in a way that is not constrained to
linearity of execution like MapReduce on Hadoop. Spark categorises some
functions/procedures in Spark programs as action events, for instance a
reduce step, and others as transformations, for instance a map step. This
allows Spark to process the execution plan DAG using a method known
as lazy evaluation, that is only action events cause data to be loaded into
memory, whereas transformations in the execution plan are only executed
when an action with dependency on that transformation is executed [17].
This method improves on cluster utilisation over Hadoop because it allows
cluster resources to be acquired and released on an ad-hoc basis throughout
a complex job. This avoids the rather less desirable but typical scenario in
Hadoop in which resources (for instance executor processes - which are
finite and dictated by cluster configuration) are reserved at the outset then
released back to the cluster (hence making them available to other jobs)
only when the running job is finished.

Apache Spark supports Python (through the PySpark API) allowing
rapid development with easily installable modules, and offers substantial
performance benefits over Hadoop. It is noteworthy that other popular
languages in the field of bioinformatics, such as R, also have interfaces to
Spark - for example SparkR [18].

1.1.3 Batch schedulers

A batch-scheduler (also referred to as a job-scheduler or workload
management software), is a central software component of a compute
cluster that manages the workload of computational jobs on a cluster and
allocates cluster resources to them [19]. We will refer to the technique as
batch-scheduling and the central software component as the job-scheduler. Generally a computational job on a batch-scheduled system is a normal user program that runs on a single compute node, but can also be a more specialist distributed program that comprise of components written to run on multiple nodes which communicate by passing messages, for example using message passing interface (MPI) (discussed in the following section).

Computational jobs are submitted in a batch to the job scheduler which adds them to a queue. The job scheduler decides on how to prioritise competing jobs, what resources to allocate to the jobs. The jobs are then submitted by the job scheduler to compute nodes of the cluster using a scheduling algorithm [20].

Batch-scheduled cluster systems couple job flow control with the ability to reserve (and limit) the allocation of cluster resources such as, for example, CPU cores, physical RAM, virtual memory and execution time. However batch-scheduled clusters offer only “course granularity” control of concurrency at the job-level (unlike MPI systems) and does not render the same level of fault-tolerance and data-locality through lack of a distributed file system such as HDFS that Hadoop provides (discussed above) unless an external distributed file system such as, for example, GPFS or Lustre is employed. Batch-scheduled cluster systems use the job scheduler to deploy “whole” executable programs to compute nodes of the cluster which may or may not run in a parallel fashion - for instance a single program when submitted will run on only one node, while multiple submitted jobs may run either on a single node or be distributed across multiple nodes depending on the load on each compute node and the scheduling rules set.

1.1.4 MPI
High performance computing (HPC) systems are typically reliant on a high degree of inter-process communication. The Message Passing Interface (MPI) is a standard for the development of parallel programming software and libraries for parallel computing architectures that standardises syntax [21]. MPI facilitates concurrent programming by specifically dictating the standard syntax to be used for the messages passed between communicating processes. MPI supports a wide variety of architectures such as multiple computers with distributed memory, shared memory, multiple processors and heterogeneous combinations of these.

MPI offers an extremely fine granularity of control over the processes involved in the execution of parallel, concurrent programs running across networked computers in clusters. Although MPI is extensively used in high performance computing, it can be used on clusters of standard machines or workstations. MPI requires the programmer to explicitly handle parallel functionality at a lower level than for instance Hadoop which automates parallelism of users programs through the MapReduce formalism. Whilst MapReduce has parallels to MPI programming, especially in relation to MPI functions scatter and reduce, it offers automatic parallelism, as well as data-locality and fault-tolerance (discussed previously) [32].

2 Applications of Hadoop in Bioinformatics
The emergence of platforms that utilise Hadoop infrastructure that we have discussed, namely Hadoop and Spark, have not been overlooked by researchers in different areas of bioinformatics. A number of projects within the Apache Hadoop ecosystem find useful application in bioinformatics [1]. These include the data-warehousing framework Hive [23] which has an SQL type query language, the high level data-flow language Pig [24] which compiles scripts into sequences of MapReduce steps for execution on Hadoop, the machine-learning and clustering facilities offered by Mahout [25], and HBase a distributed scalable database [26]. All of these projects utilise Hadoop’s cluster infrastructure and distributed file system and therefore gain from the scalability and fault-tolerance inherent in their design, as discussed earlier.

2.1 MapReduce
In terms of software applications MapReduce has been employed for a variety of problems in processing biological and sequencing datasets. Some notable projects in the area of sequence alignment are, CloudBurst [27] and CloudAligner [28], which are both based on the RMAP alignment algorithm [29], and CloudBlat [30] which is based on the BLAST algorithm [31]. It is noteworthy that MapReduce can be especially suited for, for example the construction of a de Bruijn graph for de novo genome assembly. For example, Contriall is able to build adjacency lists for all the k-mers in the genomic sequence reads and then uses distributed aggregation functions such as reduce to compress simple chains of length N in \( O(\log(N)) \) rounds using a randomized parallel list ranking algorithm [32].

There are also tools implemented in MapReduce for the analysis of assembled sequencing data, for instance Crossbow [33] is designed for SNP (Single Nucleotide Polymorphism) detection. It uses the Bowtie [34] and the SNP caller SOAPpop [35]. Differential expression (using RNA-Seq) can be measured using the Myma software pipeline [36] - pipelines are data-flows comprising of sequential steps in which bioinformatics software are applied to the data [37].

Additionally, a number of programming libraries that facilitate the manipulation and processing of sequencing data file formats such as SAM Sequence Alignment Map and BAM (Binary Alignment Map) have arisen such as the Java based libraries Genome Analysis Toolkit (GATK) [38] developed by the Broad Institute and Hadoop-BAM [39] as well as the Scala based SparkSeq [40] (discussed below). The GATK provides functions for data management in the form of data access patterns, namely the low level implementation is separated from higher level functions, and also provides functions for analysis calculations. The Broad Institute have also developed a Workflow Definition Language (WDL) for use in data analysis pipelines (discussed in the next section). It is a high-level language that is designed to be human readable and writable. It allows researchers to describe analysis tasks, daisy-chain tasks into workflows, and utilise advanced features such as parallelization [41]. WDL was developed out of the necessity for standardisation amongst a number of different pipeline solutions, thereby providing a universal standard. In order to execute analysis pipelines written in WDL, an execution engine is necessary. Cromwell is such an engine, also designed by the developers of WDL, to run on many platforms (Locally, HPC or Google - support for other platforms such as Microsoft Azure and AWS is forthcoming) and can scale effectively to workflow needs [42]. Furthermore, other notable workflow languages that aim to facilitate reproducible workflows are coming to the fore and have started to gain traction, such as CWL (Common Workflow Language) [43] and Nextflow [44]. CWL, targeted at Bioinformatics, medical imaging, Chemistry, Physics and Astronomy is supported by Cromwell, Galaxy and Taverna. Nextflow, which is based on the Groovy language, allows for abstraction of the underlying platform by virtue of an execution engine which can run and parallelise workflow jobs on platforms such as LSF, SLURM and AWS [44].

The provision of pipeline development specifically for the Hadoop platform is also available. For instance, SparkSeq is a library for building genomic analysis pipelines using Scala on Apache Spark. Whilst Scala is supported on the Spark platform it lacks the same user base in bioinformatics as it enjoys amongst the data analytics and machine learning communities. Given the vast amounts of sequencing data being produced [45, 46], the purpose of these tools is to exploit the scalability which the Hadoop and Spark platforms offer, and this offsets any difficulty in
2.2 Spark

A number of applications have been developed specifically for the Apache Spark platform. Guo et al. have carried out an extensive review of Bioinformatics applications that use Spark [47]. Nothaft et al. [48] have developed an example genomics pipeline in Spark on HDFS. They have demonstrated performance gains on the platform running on Amazon EC2 in comparison to the equivalent pipeline run using existing tools. Their pipeline comprises alignment of reads, pre-processing of reads for QC, variant calling and filtration of positive variant calls. ADAM pipeline steps were evaluated against other well-known tools used for these steps such as GATK, SAMTools and Sambamba. They achieved superlinear speedup when increasing from 8x to 16x nodes, and near linear speedup when scaling from 32x to 128x nodes. This corresponded to a 63% reduction in execution cost on Amazon EC2.

VariantSpark, developed by [49], is a machine learning analysis framework for genomic data implemented using Apache Spark. It operates on large feature vectors and large datasets for genetic classification and can perform real-time analysis. Currently VariantSpark utilises k-means clustering, the next version, however, will implement a random forest algorithm.

The Genome Analysis Toolkit (GATK), used extensively in Variant discovery, provides some of its tools implemented in Spark - previous versions provided Java libraries for Hadoop. Whilst the Spark implemented GATK tools can be run on a single Spark node, performance gains are made when the Spark platform is a cluster of compute nodes which are horizontally scaled up.

While there are a variety of different Genomics applications that use Spark, there are at present very few applications in Structural Bioinformatics. MTTF [50] is a new compact binary data format for PDB data and a single Hadoop sequence file of the data set is available for analysis by Hadoop and Spark. Other papers, such as [51] and [52] point to future plans to transition to Spark, but at present there are no other examples of Spark being applied in Structural Bioinformatics.

3 Applications in Structural Bioinformatics

The Protein Data Bank (PDB) is an archive of data describing the 3D shapes of proteins, nucleic acids, and molecular-complex assemblies derived from x-ray crystallographic, Nuclear Magnetic Resonance spectroscopy (NMR) and electron microscopy techniques [53, 5]. It also serves as a portal for structural genomics [54].

There are also a number of applications for Structural Bioinformatics implemented using MapReduce on Hadoop, specifically to carry out high-throughput analyses of such data sets which will be discussed. Schematically these are shown in figure 1. Whilst the focus of this section will be on systems developed for the Hadoop platform, for purposes of comparison, it will also refer to similar systems implemented on other platforms. We have provided a table summarising all the applications discussed below in the Supplementary Information.

3.1 Molecular docking

Molecular docking typically involves simulating the electrostatic interactions between a ligand (often a potential drug molecule) and a target protein [58, 59]. It is used to score ligands on their affinity to the target, usually for the purposes of drug development - a process that is complex, time-consuming and expensive [60, 61].
They also tested the fault-tolerance of Hadoop in running their protein-ligand pair docking system by simulating task failures in 50% of the map steps by removing node(s) from service. They observed that the docking jobs still completed. As discussed previously, fault-tolerant distributed computation is a feature of Hadoop based applications, and this resilience in the execution of tasks is important, because the likelihood of a node failing increases with the scaling-up of a cluster. Fault-tolerance is also desirable in web-based services such as Cloud-PLBS, which serve to automate computational jobs and present the results to the user, without requiring third-party intervention to rectify failed jobs. However, it should be noted that no reference to source code for their system is provided in their paper, and the Cloud-PLBS service at http://bioinfo.cs.pu.edu.tw/cloud-PLBS/index.html is no longer available.

3.1.2 Clustering of protein-ligand complexes

One of the challenges in the field of molecular docking studies arises from the requirement to search the conformational space of protein-ligand complexes generated across docking experiments. This is necessary to select the most likely conformations of protein-ligand complexes, and, therefore, the putative ligands (potential drug molecules) which partake in these interactions. In such experiments large numbers of protein-ligand complexes are generated, docked, and scored [68], and it is, therefore, necessary to select a subset of putative ligands based on significant protein-ligand interactions.

Estrada et al. observe that selecting the native conformation, based on the assumption that the lowest energetically scored conformation (as computed by an energy function) represents the native binding of the ligand and protein, is not reliable, even in larger sets of conformations. This is often due to non-native ligand-protein complexes generating falsely low energy scores. They point out that, whilst hierarchical clustering techniques are a logical way of addressing this problem - as the lowest scoring, most densely populated clusters overlap with native conformation - most clustering algorithms are computationally expensive, and scale poorly with large datasets. They implemented a system using MapReduce on Hadoop to address this issue. They used two datasets, of size 5 TB (3,872 million ligand conformations) and 1 TB (768 million ligand conformations), generated from the Docking at Home volunteer grid computing project (Docking@Home) [69], which utilised CHARMM [70].

The examples discussed in the section 3.1.1 did not fully implement their solutions in MapReduce. This would have involved implementing (or re-implementing) algorithms using MapReduce, but instead exploited Hadoop’s map step to encapsulate and execute external applications. The method discussed in [68] however, is fully implemented in MapReduce. A map step is used which geometrically reduces the conformational space. This is stored in an Octree data structure [71], together with a unique identifier (an Octkey) used for traversal. This is achieved by projecting the x, y, z components of the conformations onto a 2D plane, and calculating their gradients (for x, y, and z) which are then encoded into a single point in the Octree. A reduce step is used to aggregate conformations in the Octree. Further MapReduce operations are then used to traverse the Octree using the Octkey identifier.

In order to compare the accuracy of their Hadoop-based Octree method (for selecting native conformations from the ensemble of complexes) against other approaches, namely Hierarchical clustering and Minimum Energy selection, they docked 100,000 protein-ligand complexes each for HIV, Trypsin, and P3Salpha. They obtained 80%, 75% and 25% accuracy for Hadoop based Octree, Hierarchical Clustering and Minimum Energy methods, respectively.

They also examined the accuracy of selecting native conformations from the cross-docking data in the Docking@Home datasets, whereby each conformation of the ligand in the set of complexes is docked with each conformation of protein. In doing so, they compared their Octree method with the Energy Minimum approach, and observed 43.8% and 5.8% accuracy, respectively.

In testing the scalability in processing the 5 TB dataset which, as discussed contains 3,872 million ligand conformations, they used a Hadoop cluster where each node possesses 32 cores (4x Octacore AMD Opteron 2.4 GHz), and up to a maximum of 32 nodes were available. The range of the scaling used was 1 node of 32 cores (to process 121 million conformations) to 32 nodes with a total of 1024 cores (to process the full dataset) and analysed 1, 2, 4, 8, 16, and 32 nodes - in all cases, the number of ligand conformations processed per core was 3.8 million.

It is not stated in their paper how many map processes were running per core, but it is assumed that it is 1 map step per core. Whilst they demonstrated their method was amenable to scaling, they observed an appreciable decrease in parallel efficiency with the increase in cores, from 99.1% down to 43.8% for 64 cores (2 nodes) and 1024 (32 nodes), respectively. This appears to be due to the increased overhead in communications between the processes as the number of processes increases (communications to computation ratio).

A similar application [72] was developed on the Hadoop platform that partitions the results of molecular dynamics simulations. The trajectories of atom positions, velocities and energies as a function of time are clustered, as large datasets. This method yields important information about the most probable conformations of proteins in ensembles. Their system employs the GROMOS algorithm [73], which is not inherently parallel, by implementing it as a series of map and reduce functions so as to utilise Hadoop. They tested their parallelised MapReduce implementation of the GROMOS algorithm on a Hadoop cluster comprising of 1 master and 3 slave nodes, each comprising two hexa-core Xeon E5645 CPUs, 2GB of RAM and 2 TB of disk space. They observed up to 10 and 7 times speed-up (over using sequential GROMOS) of the first and second phases, and final two phases of their algorithm, respectively.

A docking application also relevant to our discussion, although not implemented in Hadoop, has been developed by [74] using a scientific workflow management tool, SciCumulus deployed on AWS. Their system employed molecular docking, using AutoDock4 and Vina, on their platform to explore both drug discovery and scalability. Their drug discovery objective was the identification of putative drug ligands that bind to Cysteine Proteases of Protozoan genomes utilising 10,000 protein-ligand complexes. This aims to facilitate the development of drugs for the Neglected Tropical Diseases (NTDs). In investigating the scaling-up of the computational task, they utilised up to 32 heterogeneous nodes (containing varied numbers of cores) to include a total of 128 Amazon AWS EC2 cores. They observed an almost linear relationship between number of nodes and speed, but this plateaued as they approached the maximum of 32 nodes, suggesting less benefit in scaling beyond this. They point out that this is likely due to more complicated load balancing in the set of heterogeneous nodes. The result of their docking experiments using the Cysteine Protease-ligand complexes identified 287 and 355 putative ligands for AutoDock4 and Vina, respectively. It is important to note, however, that these potential drug molecules have, on average, RMSDs greater than 2−3Å (Angstroms) which is the maximum accepted value for a useful result.

3.2 Structural Alignment

The alignment of proteins by structure, as opposed to by sequence, is a computational technique used to identify homologous polymer structures within proteins that may be conserved between proteins. The technique facilitates the study of the structural and evolutionary relationships of proteins with low sequence similarity [75, 76]. A variety of algorithms have been developed to perform structural alignment of proteins, such as:

- **Pairwise Alignment**: This technique compares two protein sequences and identifies regions of similarity. The alignment is performed using dynamic programming algorithms, such as the Needleman-Wunsch algorithm, to find the optimal match between the two sequences.

- **Multiple Alignment**: This technique extends pairwise alignment to multiple sequences, often using the Consensus Tree method or the Structure-Shape-based method to determine the optimal alignment.

- **Template-based Alignment**: This technique uses a three-dimensional model of a known protein structure as a template to align and model other similar proteins.

- **Distance Geometry**: This technique aligns proteins based on their pairwise distance measurements, which can be obtained from crystallographic structures or NMR spectroscopy.

These techniques are crucial for understanding the structural and functional relationships between proteins, which can provide insights into the evolutionary history and function of proteins in various biological processes.
as, for example, STRUCTAL (Structural Analysis Algorithm), DALI (Distant Alignment) [77], CE (Combinatorial Extension) [78], VAST (Vector Alignment Search Tool) [75], and FATCAT (Flexible structure Alignment by Chaining Aligned fragment pairs allowing Twists) [79], SSAP (Sequential Structure Alignment Program) [80], and MUSTANG (Multiple Sequence Alignment Algorithms) [81]. The technique has been applied to the study of protein binding sites, and solvent exposed surfaces (these effect the energetics of protein-ligand conformations) [82, 83, 84].

Structural alignment algorithms are usually computationally complex, [85] present a method which runs at best in approximately Polynomial time, but they also point out that approximations are often used. There is also a need to apply such techniques at scale. For example, Hadoop has been employed by [84] who implemented structural alignment for binding site prediction. Using test sets of 200 and 48 ligand-protein complexes, they were able to achieve 93% and 98% accuracy, respectively, and were able to improve the efficiency of the experiment by exploiting parallelisation. A service for structurally aligning pairs of proteins has also been implemented for the Hadoop platform by the developers of Cloud-PLBS [86] (discussed in the previous section 3.1). It utilises the same distributed architecture as Cloud-PLBS, that is, individual Hadoop nodes running within their own VM, and each VM running a map and reduce process. As with Cloud-PBS, a web-interface is used to enable the user to provide input of two PDB files by their PDB-ID. To perform the structural alignment they state their system uses the DALI and VAST algorithms. Whilst the authors do detail the basis of RMSD (Root Mean Square Deviation) in structural alignment algorithms, and discuss refinement methods in their paper, they claim to implement these algorithms in MapReduce. As previously noted with Cloud-PLBS, there is no source code available, and the corresponding web service is unavailable. It is highly likely that, given the complexity of these algorithms and that there are already implementations available, the same method used in Cloud-PLBS - execution of an external program within a map step - is employed.

A similar bioinformatics SaaS (Software-as-a-Service) for structural alignment of proteins, has been developed for the Microsoft Azure platform - Cloud4Psi developed by [87]. Their service utilises three newer algorithms that are implemented in the BiJava project, and which are derived from CE (jCE), and FATCAT (jFATCAT-rigid and jFATCAT-flexible) [88]. They tested their system on a subset of 1,000 PDB structures for both scalability and reproducibility. For scalability, two different scaling methods were compared: horizontal-scaling (i.e. by adding more nodes to the system) and vertical-scaling (i.e. by using nodes with more CPU cores). They found that, whilst both scaling methods increased the n-fold speed-up for each of the three algorithms, both suffered a decrease in the performance gains - for horizontal scaling, this was found to be the result of increased disk I/O due to multiple nodes utilising a shared VHD (Virtual Hard-Disk), and for vertical scaling this was due to an increase in processes on the same node (due to higher specification of each node), resulting in increased CPU utilisation. As the horizontal scaling method suffered less from this effect, it was the method chosen. For reproducibility of results, they found that each of the three algorithms produced the same results independent of cluster configuration and scaling used.

3.3 Other Structural Bioinformatics applications using Hadoop

Large-scale processing of molecular data is desirable in both applications. Such techniques facilitate the in-silico study of vast arrays of molecular compounds and macromolecular structures that are available from large molecular databases, which are also increasing in size and diversity [89, 90, 91, 92, 53, 7]. A number of scalable Structural Bioinformatics methods have been provided by the bioinformatics Group at UCL (University College London) as web-based services through their Protein Analysis Workbench [93]. These are accessible via SOAP (Simple Object Access Protocol), and XML-RPC (Extensible Markup Language-Remote Procedure Call) protocols. Importantly, the most commonly used methods have also been deployed as Java packages specifically for the Hadoop platform. This includes PSIPRED for protein structure prediction [94], GenTHREADER for protein fold recognition method using genomic sequences [95] and Disopred [96] for predicting protein disorder. However, other Workbench tools such as BioSerf, MEMSAT, DomPred, MetSite and FFPred appear not to have been deployed as Java packages for Hadoop and the Hadoop package is not available in their most up to date https://github.com/psipred suggesting that it is not being updated.

4 Conclusions

This purpose of this review is to give an insight into the impact that Hadoop and the MapReduce formalism has in Structural Bioinformatics. This is a appropriate moment to consider this as there have been a number of different applications of Hadoop in the field and Hadoop (and the wide variety of different applications that have been built on it) has become more stable and accessible. Spark is very likely to supplant Hadoop. Researchers working in Structural Bioinformatics have not yet begun to apply Spark to their own research and the same challenges (determining the trade off between entirely rewriting a legacy application to make use of the paradigm or using the formalism to launch the legacy application, understanding the algebraic formalism based on key-value pairs, benchmarking with traditional approaches such as batch schedulers and optimising configuration) apply there as well.

As noted previously the adoption of Hadoop is not a trivial step, particularly for a Structural Bioinformatics lab that already has extensive experience in using traditional batch schedulers running on a local cluster. Rewriting stable code to make them use the MapReduce formalism at the lowest level would require substantial effort though using MapReduce to call executables requires much less effort and can still make use of the fault-tolerance and data locality features of Hadoop.

Questions have been raised about whether the focus on scalability is masking other significant overheads for such systems and a metric has been developed to examine this [97]. The impact of this for Bioinformatics applications has not been explored in the literature and is an interesting and open question in this area.

This article has focused on

- determining the breadth of cases where Hadoop has been used,
- how well it scales and
- how dependent the installation of Hadoop is on its configuration (which is indicative of the difficulty one would have in installing Hadoop).

In the first instance we see a range of traditional high-throughput applications in Structural Bioinformatics (e.g. docking and structural alignment) where Hadoop has been employed.

With respect to scaling, the publications reviewed here indicate that some adjustment of parameters have been made, but these largely focused on how the applications scale with the number of nodes. They show that performance is linear though the gains in performance tend to reduce as the systems are scaled up. In molecular docking, [68] observed a fall in parallel efficiency of 55.3% (99.1% - 43.8%) when scaling from 2 nodes to 32 nodes. In structural alignment, [84] observed that performance degraded slightly after 8 mappers was increased to 30. Furthermore, this trend has also been observed on the MS Azure platform we have discussed for comparison - in scaling Cloud4Psi, also a structural alignment application [87], observed that horizontal scaling resulted in performance degradation.
as a result of an increase in nodes sharing a virtual disk, and that vertical scaling resulted in performance degradation as a result of increased CPU utilisation (due to more processes running per node).

The platform, and method of distribution, also dictate performance and scalability. In observing two identical docking operations on the DUD database, one using a 1.088 core Hadoop cluster [62], and the other using 15,408 cores with a mixed parallel MPI implementation of AutoDock [65], the Hadoop cluster took 69 hours, and the MPI implementation completed within 24 hours. Whilst this is certainly a result of the number of cores, the MPI system scaled better, with only very slight degradation in performance after 6,000 CPU cores. Although this comparison involved the same docking operation and dataset using different platforms, currently, there are no comparisons of performance between Hadoop and batch-schedulers on the same cluster apparatus in the literature.

The comparison of vertical and horizontal scaling in Cloud4Psi indicates significant change in performance, so configuration is important. As noted with the Cloud4Psi example, there can be a significant variation depending on configuration. Performance gains across applications, therefore, are dependent on configuration.

As Hadoop platforms stabilise, the significant advantage of its employment is of using a platform where the computation is expressed explicitly in terms of an algebra. This makes building workflows easier by allowing the developer to concentrate on the calculation, rather than the process such as WDL [41]. Nonetheless, there is a significant gap in take up as such systems remain difficult to deploy.

The Hadoop ecosystem is rapidly evolving and such gaps will reduce over time. Spark will improve the situation further. On the other hand, it will also require regular release upgrades. These can be potentially difficult to deploy, and often add new components to the ecosystem which increase the potential to introduce bugs that may affect different areas of the system.

To address this, organisations such as Cloudera and Hortonworks [98, 99] provide supported Hadoop-stack releases, and cloud-service providers such as Amazon offer managed-services, for example Elastic Map Reduce (EMR) [100]. Implementing systems on the Hadoop platform, as discussed, also requires specialist programming knowledge of MapReduce, and if the relevant Hadoop cluster is maintained in-house, specialist skills in maintaining an Hadoop cluster are also required. For this reason, managed services are often utilised by enterprise companies because such systems have been deployed and tested by technical experts and therefore mitigates risks, and dispenses the need to employ or train in-house skilled personnel to maintain a Hadoop cluster.

In summary we see a complex pattern of results suggesting that while Hadoop is promising as a platform for Structural Bioinformatics, the benefits are uneven. The use of Spark should produce better and more consistent performance and we look forward to future work in this regard.

Key Points
- There exist a number of implementations using Hadoop of high-throughput analyses, e.g. ligand-protein docking and structural alignment.
- In general these deployments are based on using standard executables which are called from the MapReduce formalism rather than rewrites of the software.
- Scalability is complex and configuration-dependent though there is some evidence that MPI implementations scale better than Hadoop.
- The ease to which Hadoop can be used for Structural Bioinformatics will improve through further innovations such as Spark, easy to deploy instances of cloud platforms and the Workflow Languages such as WDL, CWL, and Nextflow.

Author Description:
Dr. Jamie Alnusri completed his PhD from University of London. His research interests are Next Generation sequencing, Distributed Computing, High Performance Computing, Computational Biology, and Bioinformatics. He currently works in Scientific Computing at the Institute of Cancer Research, London.

Dr. Hugh P. Shanahan is a Reader at the department of Computer Science, Royal Holloway, University of London. His research interests are Next Generation Sequencing, Metagenomics and High Throughput Computing.

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