

# Final Year Project Report

Full Unit - Final Report

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## Structural Bioinformatics Framework using a MapReduce formalism

Vinay Kakkar

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**Supervisor:** Hugh P. Shanahan



Department of Computer Science  
Royal Holloway, University of London

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# Abstract

Structural bioinformatics is a rapidly growing field that aims to understand biological processes at the molecular level. In this dissertation, we present a novel framework for structural bioinformatics using a MapReduce formalism. Our framework allows for efficient processing of large-scale structural data by distributing computation across a cluster of computers. We demonstrate the effectiveness of our approach through benchmarking and comparisons with current implemented solutions. Our framework not only provides faster computation but also offers improved scalability and fault tolerance, making it a valuable tool for large-scale structural bioinformatics analyses. Furthermore, we highlight the potential of our framework for facilitating collaboration and data sharing among researchers, which is crucial for advancing our understanding of complex biological systems. Overall, our proposed framework presents a significant step forward in the field of structural bioinformatics, enabling the efficient and scalable analysis of complex structural data.

## 1 Motivation

Structural bioinformatics plays a critical role in the development of new drugs, as well as in understanding the molecular basis of biological processes. It involves the analysis and interpretation of large amounts of complex data, including the three-dimensional structures of proteins, DNA, and other macromolecules. As the size and complexity of these data sets continue to increase, the need for efficient and scalable computational tools for their analysis and processing becomes ever more pressing.

One promising approach to addressing this challenge is the use of MapReduce, a programming model for large-scale data processing in distributed computing environments. MapReduce enables the parallel processing of large data sets across multiple nodes in a cluster, which can significantly improve the speed and efficiency of data analysis tasks.

In this report, we present a structural bioinformatics framework that utilizes a MapReduce formalism for the efficient analysis of large-scale structural data. Our framework is based on a distributed computing architecture that allows for the parallel processing of structural data, including protein structures, protein-protein interactions, and other molecular structures. We describe the key components of our framework, including the data preprocessing, map, and reduce phases, as well as the parallel algorithms and data structures used to implement these phases.

We demonstrate the effectiveness of our framework through a series of experiments on real-world structural data sets, showing that our approach can significantly reduce the time and resources required for complex bioinformatics tasks. Our results highlight the potential of MapReduce-based approaches for accelerating the analysis of large-scale structural data in the field of bioinformatics and facilitating breakthroughs in drug development and biological research.

## 2 Aims and Objectives

### 2.1 Aim provided in the Project Description

”The aim of this project is to provide a framework where large numbers of protein structures can be analysed using a user-provided executable in the MapReduce formalism.”

#### Aim

Analysing protein structures can provide insights into their properties and interactions with other molecules. However, analysing large numbers of protein structures can be computationally intensive, requiring significant computational resources and time. Thus, the aim is to provide a framework that allows for the analysis of large numbers of protein structures using a MapReduce formalism.

**MapReduce:** is a programming model for processing and generating large data sets. Allowing for parallel processing of data across multiple nodes in a cluster, making it well-suited for analysing large datasets.

More specifically the aim is to develop a framework that can take a user-provided executable and apply it to a large number of protein structures in a parallel, distributed manner using MapReduce. This will enable researchers to perform complex analyses on large datasets of protein structures more efficiently, saving time and computational resources. Essentially enabling researchers to analyse large numbers of protein structures more quickly and effectively than before.

#### Motivation

The analysis of protein structures is a critical area of research, providing insights into their properties and interactions with other molecules. However, as mentioned before the computational demands of analysing large datasets of protein structures can be a significant obstacle, requiring substantial time and resources. To address this challenge, this project aims to develop a framework that utilizes the MapReduce programming model to enable researchers to efficiently analyse large numbers of protein structures using a user-provided executable. By doing so, this framework has the potential to revolutionize the field, allowing researchers to perform complex analyses on massive datasets in a parallel, distributed manner. This will ultimately save valuable time and computational resources, ultimately enabling researchers to make more rapid progress in their investigations of protein structures and potentially leading to new discoveries in the field.

## 2.2 Objectives

To achieve the aim of the project, the following objectives have been formulated:

1. Develop a software framework that supports the MapReduce formalism and can process large numbers of protein structures.
2. Implement a distributed computing system using MapReduce to parallelize protein structure analysis across multiple computing nodes.
3. Optimize the software framework to reduce the processing time required for protein structure analysis.
4. Design an interface that allows users to manipulate the PDBs that are being passed into the executable for protein structure analysis.
5. Ensure the software framework is scalable and can handle increasingly large datasets.
6. Ensure the software framework can be easily updated to keep pace with advancements in protein structure analysis techniques and computing technology.
7. Provide documentation and user support to enable researchers to use the software framework effectively

## MapReduce Framework for Protein Analysis

Develop a software framework that supports the MapReduce formalism and can process large numbers of protein structures.

With the increase in the amount of data being generated in Structural Bioinformatics, there is a need for frameworks that can process large amounts of data efficiently. The MapReduce programming model is a powerful tool for processing large datasets across a distributed computing cluster. It allows for parallel processing of data by dividing it into smaller chunks, which are then processed in parallel across multiple computing nodes. Thus, developing a software framework that supports MapReduce allows for the efficient processing of large numbers of protein structures, as it can handle the distributed processing of data across a cluster of computers.

## Parallelized Protein Analysis with MapReduce

Implement a distributed computing system using MapReduce to parallelize protein structure analysis across multiple computing nodes.

Protein structure analysis is a computationally intensive task that can take a long time to complete on a single computer. By implementing a distributed computing system using MapReduce, the analysis can be parallelized across multiple computing nodes, significantly reducing the time required to process the data. This allows for faster analysis of protein structures, which is essential in research fields such as drug discovery.

## Optimization for Faster Protein Analysis

Optimize the software framework to reduce the processing time required for protein structure analysis.

Even with a distributed computing system, the processing time required for protein structure analysis can be significant. Thus, the software framework must be optimized to reduce the processing time required for protein structure analysis. This can be achieved by improving the efficiency of the algorithms used in the analysis, minimizing the amount of data transferred between nodes, and optimizing the hardware used in the computing cluster. Optimizing the software framework ensures that protein structure analysis can be done as quickly and efficiently as possible.

## User Interface for Protein Analysis

Design an interface that allows users to manipulate the p dbs that are being passed into the executable for protein structure analysis.

As the framework revolves around using and manipulating PDB file. An interface for manipulating the input data is necessary to provide users with the flexibility to customize their protein structure analysis. This interface should be intuitive and easy to use, allowing researchers to select the protein structures they want to analyze and adjust the analysis parameters as needed. Providing this interface ensures that researchers can tailor the analysis to their specific needs and allows for greater flexibility in the types of analysis that can be performed.

## Scalable Data Handling for Protein Analysis

Ensure the software framework is scalable and can handle increasingly large datasets.

As the size of protein structure datasets continues to increase, it is essential that the software framework can scale to handle this growth without a significant decrease in performance. The system should be designed to handle datasets of various sizes and be able to scale up or down based on the demands of the analysis. This ensures that the software framework can handle the ever-increasing amounts of data generated by modern research techniques.

## Upgradable Framework for Advanced Analysis

Ensure the software framework can be easily updated to keep pace with advancements in protein structure analysis techniques and computing technology.

As the field of protein structure analysis continues to evolve, it is important that the software framework can keep pace with the latest advancements. This means that the framework should be designed to be modular and extensible, allowing for new analysis techniques to be easily integrated. It should also be designed to take advantage of the latest computing technologies, high-performance computing clusters. This ensures that the software framework can remain relevant and effective as new research techniques are developed.

## User Support for Protein Analysis Software

Provide documentation and user support to enable researchers to use the software framework effectively.

To ensure that researchers can use the software framework effectively, comprehensive documentation and user support are essential. This includes detailed documentation on how to install and configure the software, as well as guides on how to use the various functions provided by the framework. Providing documentation and user support helps to ensure that researchers can use the framework for their research purposes and obtain accurate results from their protein structure analyses.

## Further Objective: Integrate API for PDB access

Integrate a PDB file download and search from the RCSB API for efficient data access.

The PDB is a crucial resource for structural bioinformatics, and researchers need to be able to access and search for specific PDB files relevant to their research. By integrating an API that allows users to download and search for PDB files, researchers will have an easier time accessing and analysing the relevant structural data. This will increase the efficiency of the research process and allow researchers to make more informed decisions. Additionally, making this data more accessible to researchers can also promote collaboration and sharing of data, furthering scientific progress in the field of structural bioinformatics.

### 3 Protein Structures, PDB Bank and Big Data Frameworks

#### 3.1 Protein Structures

##### Introduction

Amino acids are molecules that when combined forms proteins. All of the 20 amino acids, see table 1 have in common a central carbon atom which is attached to a hydrogen atom, an amino group, and a carboxyl group [BT98].

Proteins are responsible for catalysing most of the chemical reactions in cells. They can function as enzymes catalysing a wide variety of reactions important for life and thus also important for the structure of living systems such as proteins involved in the cytoskeleton. The size of protein can vary [Zve08].

**Definition 3.1 (Catalysing)** *Catalysing is to make a chemical reaction happen or happen more quickly by acting as a catalyst.*

**Definition 3.2 (Cytoskeleton)** *A dynamic network of interlinking protein filaments present in the cytoplasm of all cells [Zve08].*

##### Primary, Secondary, Tertiary and Quaternary Structure

Please refer to 1 for a visual representation.

The **primary structure** of a peptide or protein is the linear sequence of its amino acids. It is read and written from the amino-terminal to the carboxyl-terminal end. [SFB04].

The **secondary structure** refers to the local arrangement of a peptide chain. Where several common secondary structures have been identified in proteins [SFB04].

The **Tertiary structure** is a three-dimensional structure of a protein the formation is built up of bonds and interactions that serve to change the shape of the overall protein [God22].

The **quaternary structure** of a protein is built-up of several protein chains/subunits. Each of the subunits has its primary, secondary, and tertiary structure [OR15].

## Considering Protein structure on several different levels

The fold of the protein plays part in determining the way the protein will function, and also whether it will function correctly. As there are Protein structures on different levels we need to consider the analysis of protein structure by experimental techniques such as X-ray crystallography, nuclear magnetic resonance, and RNAseq which show that proteins adopt distinct structural elements [Zve08].

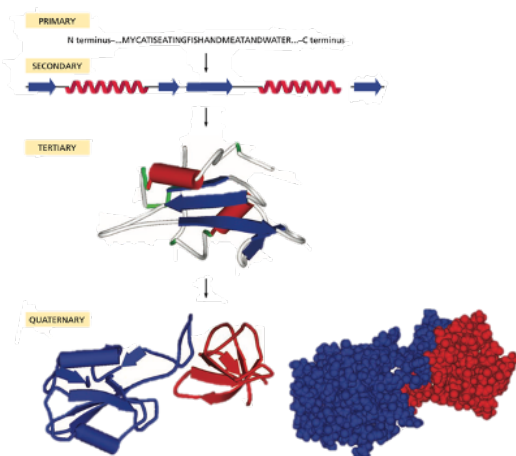


Figure 1: From the sequence alone, the primary structure to secondary structure, to tertiary structure(3D), to finally quaternary structure found when several tertiary structures form a multisubunit complex [Zve08].

## Amino Acids

Sequence of amino acids will build up the linear protein chain [Zve08]. Amino acids are different from each other due to their side chains and due to this the functional properties of various different proteins are different [Zve08]. You can see the amino acids grouped here.

| Amino acid    | Three-letter code | One-letter code |
|---------------|-------------------|-----------------|
| Glycine       | Gly               | G               |
| Alanine       | Ala               | A               |
| Valine        | Val               | V               |
| Leucine       | Leu               | L               |
| Isoleucine    | Ile               | I               |
| Proline       | Pro               | P               |
| Phenylalanine | Phe               | F               |
| Methionine    | Met               | M               |
| Tryptophan    | Trp               | W               |
| Cysteine      | Cys               | C               |
| <hr/>         |                   |                 |
| Asparagine    | Asn               | N               |
| Glutamine     | Gln               | Q               |
| Serine        | Ser               | S               |
| Threonine     | Thr               | T               |
| Tyrosine      | Tyr               | Y               |
| <hr/>         |                   |                 |
| Aspartic acid | Asp               | D               |
| Glutamic acid | Glu               | E               |
| <hr/>         |                   |                 |
| Histidine     | His               | H               |
| Lysine        | Lys               | K               |
| Arginine      | Arg               | R               |

Table 1: The 20 amino acids. The amino acid name, the three-letter code, and the one-letter code are given. The Amino acids are split up into Nonpolar, Polar, Acidic and Basic respectfully

### 3.2 Large Scale Expression

Gene expression begins when genes are transcribed into messenger RNAs, which are then translated to produce proteins.

Total gene expression in cultured cells or a tissue sample can be detected in three main ways:

1. DNA microarray technology.
2. Two-dimensional Gel electrophoresis or Chromatography.
3. RNAseq

Both DNA microarray technology and Two-dimensional Gel electrophoresis, produce enormous amounts of raw data [Zve08] due to this, many proteins currently evade high-resolution structure determination.

### Structural mass spectrometry

Structural mass spectrometry is a powerful approach used to determine the 3D structure of biological proteins it has nearly an unlimited size constraint and speed. Although the data provided by mass spectrometry is vague for full high-resolution structure elucidation, structural mass spectrometry can be used to examine the size, solvent accessibility, and topography of proteins [LLV18] [LZG20].

We can have computational methods that aid experimental technique intending to elucidate protein structures [SL20] [LWL<sup>+</sup>20]. Software packages can be used to combine data with advanced structure sampling and scoring techniques. Computational tools for protein structure modeling, include the Rosetta software suite [LWL<sup>+</sup>20] [ALFJ<sup>17</sup>], I-TASSER [YYR<sup>15</sup>], Phyre2 [KMY<sup>15</sup>], Integrative Modeling Platform [RLW<sup>12</sup>], HADDOCK [DBB03], and MODELLER [EWMR<sup>06</sup>] [BL22].

## Large Scale Gene Expression

Genome DNA microarray experiments produce large amount of data can be computationally heavy on where methods can yield alternative conclusions from increasing the computational effort.

The goal of these experiments is to determine biological or functional meaning from the lists of genes, either by:

1. Identify critical genes that are responsible for a biological effect.
2. Find patterns within the genes that point to an underlying biological process.

[Zve08]

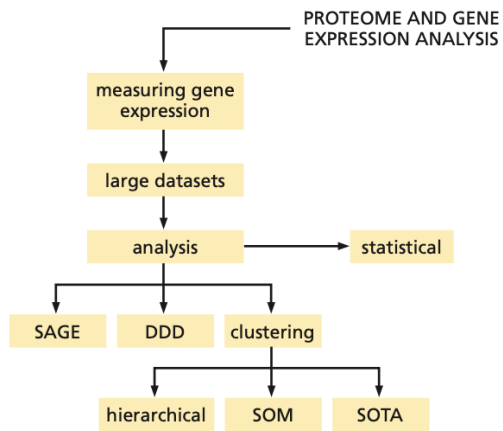


Figure 2: Describing Common experimental aspects of gene expression and of the analysis of the resulting data [Zve08].

## Serial analysis of gene expression

Serial analysis of gene expression is the alternative compared to microarrays when trying to investigate patterns of gene expression.

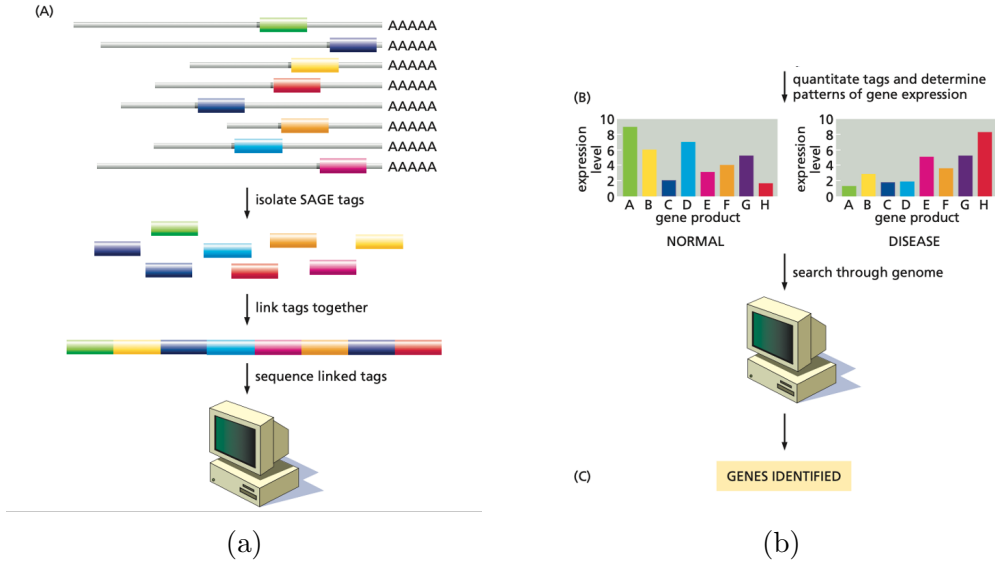


Figure 3: *An outline of the SAGE method for comparing levels of gene expression. (A) Short sequence tags. The sequence tags are isolated and are linked together to produce long DNA molecules that can be cloned and sequenced. (B) Once sequenced, each tag can be calculated, resulting in a value that gives the expression level of the corresponding transcript [Zve08].*

A short sequence contains enough information to uniquely identify a gene. The sequence tags from the total cellular RNA can be linked together to form long DNA molecules. The total number of times a particular tag is observed approximates the expression level of the corresponding gene. The data produced by SAGE include a list of the tags with their corresponding counts, providing a digital output of cellular gene expression. Which allows the user to specify which organ is to be investigated. Libraries consisting of gene lists organized by the various types of tissues or cell lines are provided for further choice. The output from SAGE provides the SAGE tag, the UniGene ID, the gene description, and color and letter-coded differences in expression levels [Zve08].

## Clustered gene expression data

Clustered pattern data obtained from gene expression microarrays/genome bioinformatics can be used as a tool to identify new transcription factors or other cell-regulatory proteins.

The clustered genes/proteins can be analyzed. Leading to a vast collection of data from many gene/protein expression experiments being available on the Web [Zve08].

## Large Scale Protein Expression

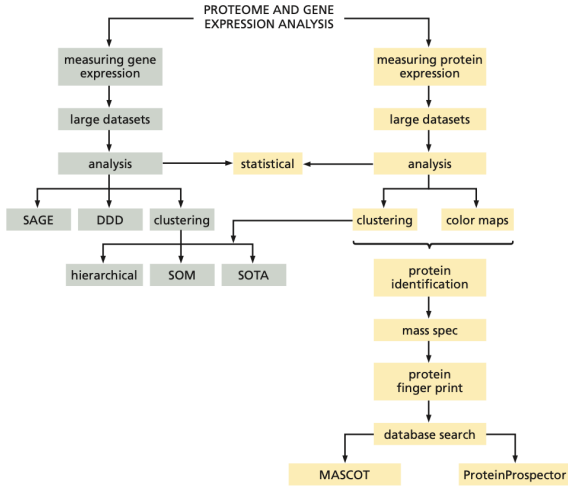


Figure 4: Describing some experimental aspects of protein expression and of the analysis of the resulting data. [Zve08].

For functional protein, mRNAs need to be translated, whilst the protein products can change which influence their function. For this reason we can measure and analyse different proteins.

There is more proteins than there are genes in a genome. Transcripts can be spliced in various ways to give different mRNAs, providing different protein products, from the same gene. However, proteins that can be modified after translation giving more different protein products.

Protein expressions can vary in an organism depending on the origin and it will also differ between the separate stages of an organism's life cycle and under different environmental conditions [Zve08].

**Definition 3.3 (proteome)** *The proteome refers to all the proteins that make up an organism at a specific point in time and under specific conditions.*

## RNAsseq

The transcriptome is important for revealing the molecular constituents of cells and tissues, interpreting the functional aspects of the genome, also for understanding development and disease [WGS09].

Many methods deduce and quantify the transcriptome, including hybridization or sequence-based approaches. For example, hybridization-based approaches involve incubating fluorescently labeled cDNA with microarrays or commercial high-density oligo microarrays [WGS09].

However, these methods have several limitations, such as:

- Dreliance upon existing knowledge about genome sequence.
- Limited dynamic range of detection owing to both background.
- High background levels owing to cross-hybridization [OM06] [RRG07].
- saturation of signals.

**Definition 3.4 (transcriptome)** *The transcriptome is the complete set of transcripts in a cell, and their quantity, for a specific developmental stage or physiological condition.*

Sequence-based approaches directly determine the cDNA sequence such as Tag-based methods which include SAGE, CAGE [KKN<sup>+</sup>06], MPSS [RBL<sup>+</sup>02].

Each approach is high throughput and can provide precise, gene expression levels. However, a significant portion of the short tags can not be uniquely mapped to the reference genome [WGS09].

RNA-Seq RNA sequencing has clear advantages over existing approaches it uses deep sequencing technologies where a population of RNA is converted to a library of cDNA fragments with adaptors attached to one or both ends. Each molecule is then sequenced in a high-throughput manner to obtain short sequences from one or both ends [WGS09].

## Bioinformatic Difficulties with Predictions on Proteins

It is difficult to define the precise ends of the helices(The secondary structure of proteins is made up of a-helices and b-strands) for structures found in globular proteins that are not perfectly regular. Making it one step more difficult when trying to predict these structures [Zve08].

To Note:

- Several different types of b-sheet are found in protein structures.
- Turns, hairpins, and loops connect helices and strands.
- Any chain between two regular structures is referred to as a loop.
- Mostly a loop will contain a turn (or even several).

In antibody recognition, immunoglobulins employ loops at the edge of a b-sheet. All immunoglobulin structures with the same overall chain fold, but it is the difference at these loops that results in different results. Loops take up one of a limited number of structures called canonical forms. This type of classification is another reason why trying to predict both the structure and function of the protein is difficult [Zve08].

**Definition 3.5 (Immunoglobulin)** *Immunoglobulins are heterodimeric proteins composed of two heavy and two light chains. Types of white blood cells that helps the body fight infection [SC10].*

## Alpha Fold

AlphaFolds' goal is to predict the 3D coordinates of all heavy atoms for a given protein using the primary amino acid sequence and aligned sequences of homologues as inputs [JEP<sup>+</sup>21].

Mutations in proteins can lead to misfolding which is often associated with disease states, for example, Alzheimer's and Parkinson's which is one of the challenges for alphaFold [Fel].

The output is a file containing the 3D coordinates for every non-hydrogen atom in the protein. whilst showing the confidence levels for every amino acid residue, providing the reliability of the predicted structure [Fel].

## Bioinformatics with Alpha Fold

In July 2021, AlphaFold was developed by DeepMind and was made available to the public [TAW<sup>+</sup>21].

Where it tries to solve the issue of invariant protein structures that are under translations and rotations [BP03].

AlphaFold is trained on protein chains from the PDB using the input sequence to query databases of protein sequences to generate a multiple sequence alignment [JEP<sup>+</sup>21]. Although we still do not exactly know how a protein sequence folds and alpha fold do not help in figuring this out its impact will likely be in accelerating and improving the production of new medications [NZLJ22].

## AlphaFold 2

The CASP14 was recently held which is a blind trial that critically assesses techniques for protein structure prediction [DITS22], AlphaFold2 was entered and out-performed all competitors.

Recently, RoseTTAFold was developed, trying to implement similar principles. Since then, other end-to-end structure predictors have emerged using different principles such as fast multiple sequence alignment processing in DMPFold218 and language model representations.[BPE22].

We use the root mean square deviation, to calculate the similarity between the two structures, AlphaFold models had an accuracy of 0.96 compared to 2.80 which was the second-best score. AlphaFold models also had a high level of accuracy in predicting the position of residue side chains when the protein backbone prediction was accurate [DITS22] [JEP<sup>+</sup>21].

### 3.3 The Protein Data Bank and the File Formats

#### Protein Data Bank

The Protein Data Bank was established at Brookhaven National Laboratories [BKW<sup>+</sup>77] in 1971 as an archive for biological macromolecular crystal structures [BWF<sup>+</sup>00].

**Definition 3.6 (Macromolecular)** *Macromolecular is any very large molecule, usually with a diameter ranging from about 100 to 10,000 angstroms*

It is an information source for data retrieved from atomic structures, crystallography, and three-dimensional structures of biomolecules, including nucleic acids and proteins [BG21].

At the time this was the first open-access digital data resource in biology which started with just seven protein structures [BBB<sup>+</sup>22b].

Various groups such as the Protein Data Bank in Europe, Protein Data Bank Japan help manage the Protein Data Bank archive. Current wwPDB members also include the ElectronMicroscopy Data Bank and the Biological Magnetic Resonance Bank [BBB<sup>+</sup>22b].

Protein Data Bank China has recently joined the wwPDB as an Associate Member with its role as wwPDBdesignated PDB Archive Keeper. Where they are responsible for weekly updates of the archive and safeguarding both digital information and a physical archive of correspondence [BBB<sup>+</sup>22a].

The management of PDB must comply with FAIR (the acronym depicts: Findable, Accessible, Interoperable, Reusable) and FACT [vdABH17] guiding principles for scientific data [WDA<sup>+</sup>16] [WSHB20].

| The FAIR Guiding Principles |   |
|-----------------------------|---|
| To be Findable:             | <ul style="list-style-type: none"> <li>F1. (meta)data are assigned a globally unique and persistent identifier</li> <li>F2. data are described with rich metadata (defined by R1 below)</li> <li>F3. metadata clearly and explicitly include the identifier of the data it describes</li> <li>F4. (meta)data are registered or indexed in a searchable resource</li> </ul>  |
| To be Accessible:           | <ul style="list-style-type: none"> <li>A1. (meta)data are retrievable by their identifier using a standardized communications protocol <ul style="list-style-type: none"> <li>A1.1 the protocol is open, free, and universally implementable</li> <li>A1.2 the protocol allows for an authentication and authorization procedure, where necessary</li> </ul> </li> <li>A2. metadata are accessible, even when the data are no longer available</li> </ul> |
| To be Interoperable:        | <ul style="list-style-type: none"> <li>I1. (meta)data use a formal, accessible, shared, and broadly applicable language for knowledge representation.</li> <li>I2. (meta)data use vocabularies that follow FAIR principles</li> <li>I3. (meta)data include qualified references to other (meta)data</li> </ul>  |
| To be Reusable:             | <ul style="list-style-type: none"> <li>R1. meta(data) are richly described with a plurality of accurate and relevant attributes <ul style="list-style-type: none"> <li>R1.1. (meta)data are released with a clear and accessible data usage license</li> <li>R1.2. (meta)data are associated with detailed provenance</li> <li>R1.3. (meta)data meet domain-relevant community standards</li> </ul> </li> </ul>   |

Table 2: The guidelines to what builds up the FAIR principles [WDA<sup>+</sup>16]

## Aims and Objectives of PDB

Enzymology, electron microscopy, computational chemistry small molecule crystallography, biochemistry, biophysics, macromolecular crystallography and nuclear magnetic resonance spectrometry all help the aims and goals of the PDB archive [BG21].

**Definition 3.7 (Enzymology)** *Enzymology is the branch of biochemistry aiming to understand how enzymes work*

**Definition 3.8 (Electron Microscopy)** *Electron microscopy is a technique for obtaining high resolution images of biological and non-biological specimens.*

PDBs provide open access to nearly 200 000 archived, validated, and biocurated experimentally determined three-dimensional structures of biological macromolecules.3D structures archived in the PDB have enabled important scientific breakthroughs by basic and applied researchers [Bur21]. Open access to PDB data without restrictions on usage has also aided structural bioinformatics in areas such as computational biology.

## Recent Project

A project was undertaken to change the information management services for RCSB.org. The idea was to have developed a primary place for studying 3D biostructures by extending RCSB.org web portal functionality to support parallel delivery of more than one million CSMs publicly available from AlphaFold DB and ModelArchive together [BBB<sup>+</sup>22a].

## Covid

During the COVID-19 pandemic, more than 2000 structures associated with the agent of the coronavirus disease were released and have become accessible to global users for free. The properties of these structures give us this opportunity to find out the ligand binding sites, the spatial conformation of ligands, protein-to-protein interactions, and amino acid substitutions regarding different viral proteins. Moreover, chemical, functional and energetic characteristics can also be gained to describe the potential capabilities of each molecule. These properties might aid us to determine the potential drug targets for drug design and vaccine preparation [LZD<sup>+</sup>20].

## PDB Currently

As of 2022, the PDB has a vast number of 3D biostructures, eukaryotic protein structures exceeded 105 000. Bacterial protein structures were also numerous, totaling nearly 66 000. Archaeal protein structures were the least numerous totaling 5500. However the PDB coverage is decidedly limited, with mouse protein structures being most numerous at 8000 structures [BB21].

We have powerful tools developed by RCSB PDB for searching and analysis which include structure, sequence, sequence motif, structure motif, and visualization [BBB<sup>+</sup>22a].

Upon reaching the RCSB.org home page, users can query, organize, visualize, analyse, compare, and explore PDB structures and CSMs side-by-side. Searching 3D structure information can encompass PDB structures and CSMs or be limited to PDB structures only. Either PDB structures or CSMs can be excluded from the search results. The two types of structure information accessible via RCSB.org are clearly distinguished from each other. Top bar searching and data delivery for PDB structures and CSMs [BBB<sup>+</sup>22a].

The screenshot shows the RCSB.org homepage with three main search sections:

- Top Bar or Basic Search:** A search bar at the top with dropdowns for "Include CSM" (On, Off) and "Include Annotations" (On, Off). It also includes "Search" and "Advanced Search Query Builder" buttons.
- Advanced Search:** A detailed search interface with fields for "Basic Attributes" (e.g., UniProt ID, PDB ID, Resource ID), "Advanced Search by term, author, ligand name, ID, structure and chemical properties, sequences, structures, motifs, chemical formula", and dropdowns for "Include CSM" and "Include Annotations".
- Browse:** A section titled "ATC Browser" which lists categories of drugs based on the Anatomical Therapeutic Chemical (ATC) classification system. It includes a "Browse by drug class, enzyme classification (E.C.), source organism, molecular function, structure classification (e.g., SCOP, CATH) and more" link.

Figure 5: Search options at RCSB.org include Top Bar or Basic Search; Advanced Search; and Browse Annotations [BBB<sup>+</sup>22a].

Five examples of search options available from every RCSB.org web page:

- Top Bar Search options:** Shows a search bar with "Turn On to include CSM" and "Search" button.
- Drop down autosuggestions:** Shows a dropdown menu for "Insulin receptor" with options like "Insulin receptor", "Insulin receptor Sultate homolog", "Insulin receptor substrate 1", etc.
- Boolean operators:** Shows a table of operators: OR (+), AND (&), NOT (-), and parentheses ((), ||). Descriptions explain how they combine search terms.
- Amino acid sequence search:** Shows a search bar with "Query: Paste protein, DNA, or RNA sequence" and "Search" button.
- Documentation search:** Shows a search bar with "Query: Documentation from left hand pull-down option - Type query for phrase search or Click on term to search for a word" and "Search" button.

Figure 6: Top Bar or Basic Search options available from every RCSB.org web page. Examples of searching for 3D structures using (A) simple text string insulin receptor; (B) drop down autosuggestions based on the text string insulin receptor; (C) Boolean operators to combine insulin + receptor (+ = AND); or (D) an amino acid sequence. (E) Searching RCSB.org documentation using a text string biological assembly [BBB<sup>+</sup>22a]

## Recent RCSB PDB data architecture improvements

In 2020, RCSB PDB had an upgrade of its delivery architecture [RDL<sup>+</sup>21] at RCSB.org [BBB<sup>+</sup>21].

The legacy monolithic data delivery application was changed into a distributed deployment of individual microservices, each with a single responsibility.

Data access services provide both Representational State Transfer and GraphQL API access to a data warehouse hosted in a MongoDB document-oriented database. Originally, advanced Search QueryBuilder functionality encompassed text, PDB data attributes, 3D structure, sequence, biopolymer sequence motif, and chemical similarity. Every search function is implemented as an independent service.

A separate search service is responsible for launching each search function, combining and delivering their integrated results to public programmatic search APIs. When each service has a single responsibility, we have greater flexibility in scaling the deployment of services in response to changes in user load and significant reductions in the time required to develop, test, and deploy new features. The Sequence Motif search function has been extended with a new 3D Structure Motifsearch capability [BBR20].

## Recent advances in RCSB PDB data integration

RCSB PDB integrates the content of each expertly biocurated Entry with information from more than 50 external data resources.

Integrated external data needs to follow a data schema that defines the organization of the RCSB PDB data warehouse. Finally, it is available to RCSB PDB front-end services, public data access APIs, and our text search indexing service [BBB<sup>+</sup>22b].

## Recent PDBx/mmCIF data standard improvements

The PDBx/mmCIF data standard is maintained by the wwPDB organization in collaboration with wwPDBPDBx/mmCIF Working Group domain experts recruited from the scientific community. The PDBx/mmCIF web resource supports browse and search access to standard terminology. The Working Group includes developers for many of the widely used structure determination software systems, who ensure that data produced by these programs comply with the PDBx/mmCIF data standard, generating complete and correct data files for PDB deposition. The wwPDB and the Working Group collaborate on developing terminologies for new and rapidly evolving methodologies such as Free Electron Laser, 3DEM, Serial Crystallography, and X-ray, whilst improving representations for existing data content. Most recently, the Working Group has focused on modernizing content descriptions for processed X-ray diffraction data, including extensions describing anisotropic diffraction limits, unmerged reflection data, and new quality metrics of anomalous diffraction data. Deposition and delivery improve our ability to assess experimental data quality, and every PDB data consumer's ability to Find and Reuse relevant PDB Entries [BBB<sup>+</sup>22b].

| External Resources            |  |
|-------------------------------|--|
| AlphaFold DB                  | Computed Structure Models by AlphaFold2  |
| ATC                           | Anatomical Therapeutic Chemical (ATC) Classification System from World Health Organization   |
| Binding MOAD                  | Binding affinities   |
| Binding DB                    | Binding affinities   |
| BMRB                          | BMRB-to-PDB mappings   |
| Cambridge structural Database | Crystallographic small molecule data from the Cambridge Crystallographic Data Centre   |
| CATH                          | Protein structure classification- Class, Architecture, Topology/fold, and Homologous superfamily   |
| ChEMBL                        | Manually curated database of bioactive molecules with drug-like properties   |
| CSD                           | Cambridge Structural Database: Validated and curated small-molecule organic and metal-organic crystal structures from the Cambridge Crystallographic Data Centre |
| DrugBank                      | Drug and drug target data  |
| ECOD                          | Evolutionary Classification of Protein Domains   |
| EMDB                          | 3DEM density maps and associated metadata  |
| ExplorEnz                     | IUBMB Enzyme nomenclature and classification   |
| Gencode                       | Human and Mouse Gene annotations   |

Table 3: Some of the External Resources Integrated Into RCSB PDB

## Future and struggles of PDB

### Future

As the PDB archive has started its 52nd year, it gives open access to analyses of structures and much more to: basic and applied researchers, educators, and students spanning fundamental biology, biomedicine, bioenergy, bioengineering, and biotechnology, with key points that help many communities that use this facility. Firstly It delivers Data In and Data Out services efficiently to a user base that is now numbering many millions worldwide. Secondly, it has wwPDB partners that process, validate, and biocurate the growing number of increasingly complex PDB depositions received. Manages and safeguards the growing PDB archive in its role as wwPDB designated Archive Keeper. Thirdly it enables searching, visualization, exploration, and analysis of experimentally-determined PDB structures integrated with more than one million CSMs through its web portal. [BBB<sup>+</sup>22a].

### Struggles

Even after all the advancements PDB has gone through there are still additional challenges lying ahead which include:

- Rapid growth in public-domain CSMs of individual polypeptide chains, already numbering >200 million at the time of writing.
- Anticipated advances in AI/ML-based prediction of structures of multi-protein complexes.
- Continued development of biomolecular structure determination methods using X-ray Free Electron Lasers, revealing the microscopic details of chemical reactions in real time.
- Growth in the number and complexity of atomic-level cryoelectron microscopy structures of macromolecular machines.
- Integration of PDB structures and CSMs with complementary information coming from correlative light microscopy and related imaging methods across length scales ranging from atoms to small molecules to individual biomolecules to macromolecular assemblies to organelles to cells and ultimately tissues
- Merging of the PDB-Dev prototype archiving system for integrative methods structures with the PDB archive
- Federating other biodata resources, such as the SmallAngle Scattering Database and the Proteomics Identification Database, with the PDB, EMDB and BMRB core archives jointly managed by the wwPDB partnership

[BBB<sup>+</sup>22a].

## File Formats

The PDB archive holds a few different types of file types that hold data such as atomic coordinates and other information describing proteins and other biological macromolecules. Depending on what the data is created from it can fall into a different category.

### PDB Data

The main information in the PDB archive is coordinate files for biological molecules. These files list the atoms in each protein and their 3D coordinates.

These files are available in several formats:

- PDB
- mmCIF
- XML

The header section of the text summarizes the protein, citation information, and the details of the structure solution, which is then followed by the sequence and a long list of the atoms and their coordinates. It also contains the experimental observations used to determine atomic coordinates [Goo].

## .pdb Files

The PB format consists of a collection of records that describe the atomic coordinates, chemical and biochemical features, and experimental details of the structure determination [WF03].

Each item of data in the PDB format is assigned to a one of PDB record types (HEADER. SOURCE. REMARK, etc.). The ATOM records the atomic coordinate data [WF03].

PDB format has been extended with new REMARK records. For example, REMARK 3 that encodes refinement information has been modified and extended for each new refinement program and program version [WF03].

The PB format uses fixed-width fields to represent data, so we have limits on the size of certain items of data. For example, we can't have more than 99,999 atoms and polymer chain can be only one character. This means some structures are divided into multiple files [WF03].

|      | Amino Acid |     |                                    | Chain name |   | Sequence Number |       |       | Coordinates |   |   |  | (etc.) |
|------|------------|-----|------------------------------------|------------|---|-----------------|-------|-------|-------------|---|---|--|--------|
|      | Element    |     |                                    |            |   |                 |       |       | x           | y | z |  |        |
| ATOM | 1          | N   | ASP                                | L          | 1 | 4.060           | 7.307 | 5.186 | ...         |   |   |  |        |
| ATOM | 2          | CA  | ASP                                | L          | 1 | 4.042           | 7.776 | 6.553 | ...         |   |   |  |        |
| ATOM | 3          | C   | ASP                                | L          | 1 | 2.668           | 8.426 | 6.644 | ...         |   |   |  |        |
| ATOM | 4          | O   | ASP                                | L          | 1 | 1.987           | 8.438 | 5.606 | ...         |   |   |  |        |
| ATOM | 5          | CB  | ASP                                | L          | 1 | 5.090           | 8.827 | 6.797 | ...         |   |   |  |        |
| ATOM | 6          | CG  | ASP                                | L          | 1 | 6.338           | 8.761 | 5.929 | ...         |   |   |  |        |
| ATOM | 7          | OD1 | ASP                                | L          | 1 | 6.576           | 9.758 | 5.241 | ...         |   |   |  |        |
| ATOM | 8          | OD2 | ASP                                | L          | 1 | 7.065           | 7.759 | 5.948 | ...         |   |   |  |        |
|      | \\         |     | Element position within amino acid |            |   |                 |       |       |             |   |   |  |        |

Figure 7: Showing contents of a PDB file for the Atom values [AAB<sup>+19</sup>].

## .mmCIF Files

Mmcif is a dictionary-based approach to data extracted from crystallographic experiments [WF03].

It includes all the data we can find in a pdb file. Also, we have sufficient data names so that the experimental section of a structure paper can be written automatically and to facilitate the development of tools i.e. computer programs could easily access and validate mmCIF data files [WF03].

## .xml

XML builds from a PDB Exchange dictionary. Although presented in very different syntaxes, the PDB Exchange and XML representations use the same logical data organization. [WIN<sup>+</sup>05].

The dictionary data block is mapped to the standard top-level XML schema element, and the data file data block is mapped to a datablock element. Category or table definitions in the Exchange dictionary are described as XML complex types. The category definition. [WIN<sup>+</sup>05].

| PDB Exchange data dictionary<br>attributes | XML schema mapping   |
|--|--|
| Data block                                 | Root level <i>schema element</i>   |
| Category groups                            |  |
| Categories                                 | <i>complexType</i>   |
| Definition                                 | <i>annotation and documentation elements</i>                                       |
| Examples                                   | <i>annotation and documentation elements</i>                                       |
| Primary keys                               | <i>attributes</i> of the data category   |
| Items                                      | <i>elements</i> of the data category   |
| Definition                                 | <i>annotation and documentation elements</i>                                       |
| Examples                                   | <i>annotation and documentation elements</i>                                       |
| Data types                                 | <i>simpleType</i>  |
| Range restrictions and allowed values      | <i>restrictions</i> within <i>simpleType</i> or <i>unions</i> of <i>simpleType</i> |
| Mandatory data code                        | Element attributes <i>minOccurs</i> and <i>nillable</i>                            |
| Parent-child relationships                 | <i>key/keyref</i> elements   |
| Interdependency/exclusivity                |  |
| Units of measurement                       | Additional <i>fixed attributes</i>   |
| Subcategory membership                     |  |

Figure 8: Summary of the correspondences between PDB Exchange data dictionary and XML schema metadata [WIN<sup>+</sup>05].

## Visualizing Structures

PDB files can be viewed from text editors but we can also use a browsing or visualization program. RCSB PDB allows you to search and explore the information, including information on experimental methods and the chemistry and biology of the protein. Visualization programs allow to read of the PDB file and, display the protein structure generating custom pictures of it. These programs can contain analysis tools that allow you to measure distances and bond angles, and identify interesting structural features [Goo].

## Reading Coordinate Files

Before exploring structures in the PDB archive we need some prior understanding of the coordinate files. For example, we can find a diverse mixture of biological molecules, small molecules, ions, and water which can get confusing we can use the names and chain IDs to help sort these out. In structures determined from crystallography, atoms are annotated with temperature factors that describe their vibration and occupancies that show if they are seen in several conformations. NMR structures often include several different models of the molecule [Goo].

## Potential Challenges

There are some things to note as you could fall into some challenges when browsing through the PDB archive. Many structures, particularly those determined by crystallography, only include information about part of the functional biological assembly. One thing to note is that the PDB can aid with this. Another note is many PDB entries are missing portions of the molecule that were not observed in the experiment. These include structures that include only alpha carbon positions, structures with missing loops, structures of individual domains, or subunits from a larger molecule. In addition, most of the crystallographic structure entries do not have information on hydrogen atoms [Goo].

### 3.4 Hadoop spark and pyspark

#### What is Hadoop

Hadoop is an open-source framework for writing and running distributed applications that process large amounts of data. Key aspects making it valuable such 1. Accessible 2.Robust 3. Scalable 4.simple [Lam10].

HDFS is used in haddop which is a filesystem and a MapReduce engine. With one master node and many worker nodes. The master node provides instructions to the worker nodes and computations are performed on the worker nodes. [HRJ17].

#### Mapper

Input key/value pairs are mapped to a set of key/value pairs. The mapper then sorts the key-value pairs by the keys. Partitioners are mainly responsible for providing intermediate key/values to the reducers [PBN12] [HRJ17].

#### Reducer

Firstly, the reducer combines data having the same key from different map functions. The values having the same key are reduced to a smaller set of values and output is produced [HRJ17].

## What is Spark

Apache Spark is a popular open-source platform for large-scale data processing used for iterative machine learning tasks [MBY<sup>+</sup>16].

Spark is a cluster computing system providing APIs in Java, Scala, Python (pySpark), and R, along with an optimized engine that supports general execution graphs. Moreover, Spark is efficient at iterative computations so it is suited for the development of large-scale machine learning applications [MBY<sup>+</sup>16].

Spark is a quick and general engine used for analysing large-scale data stored across a cluster of computers. Spark uses in-memory cluster computing which is its most important feature for increasing the processing speed of an application. It combines SQL streaming and complex analytics [HRJ17].

## Spark Architecture

There are five core components that make Spark so powerful and easy to use. The core architecture of Spark consists of the following layers:

- Storage
- Resource management
- Engine
- Ecosystem
- APIs

[Sin22].

## Storage

Before using Spark, data must be made available in order to process it. This data can reside in any kind of database. Spark offers multiple options to use different categories of data sources, to be able to process it on a large scale. Spark allows you to use traditional relational databases as well as NoSQL, such as Cassandra and MongoDB [Sin22].

## Resource Management

The next layer consists of a resource manager. As Spark works on a set of machines (it also can work on a single machine with multiple cores), it is known as a Spark cluster. Typically, there is a resource manager in any cluster that efficiently handles the workload between these resources. The two most widely used resource managers are YARN and Mesos. The resource manager has two main components internally [Sin22]:

- Cluster manager
- Worker

It's kind of like master-slave architecture, in which the cluster manager acts as a master node, and the worker acts as a slave node in the cluster. The cluster manager keeps track of all information pertaining to the worker nodes and their current status. Cluster managers always maintain the following information [Sin22]:

- Status of worker node (busy/available)
- Location of worker node
- Memory of worker node
- Total CPU cores of worker node

The main role of the cluster manager is to manage the worker nodes and assign them tasks, based on the availability and capacity of the worker node. On the other hand, a worker node is only responsible for executing the task it's given by the cluster manager [Sin22].

The tasks that are given to the worker nodes are generally the individual pieces of the overall Spark application. The Spark application contains two parts [Sin22]:

- Task
- Spark driver

The task is the data processing logic that has been written in either PySpark or Spark R code. It can be as simple as taking a total frequency count of words to a very complex set of instructions on an unstructured dataset. The second component is Spark driver, the main controller of a Spark application, which consistently interacts with a cluster manager to find out which worker nodes can be used to execute the request. The role of the Spark driver is to request the cluster manager to initiate the Spark executor for every worker node [Sin22].

## Engine and Ecosystem

The base of the Spark architecture is its core, which is built on top of RDDs (Resilient Distributed Datasets) and offers multiple APIs for building other libraries and ecosystems by Spark contributors. It contains two parts: the distributed computing infrastructure and the RDD programming abstraction. The default libraries in the Spark toolkit come as four different offerings [Sin22].

### Spark SQL

SQL being used by most of the ETL operators across the globe makes it a logical choice to be part of Spark offerings. It allows Spark users to perform structured data processing by running SQL queries. In actuality, Spark SQL leverages the catalyst optimizer to perform the optimizations during the execution of SQL queries. Another advantage of using Spark SQL is that it can easily deal with multiple database files and storage systems such as SQL, NoSQL, Parquet, etc [Sin22].

### MLlib

Training machine learning models on big datasets was starting to become a huge challenge, until Spark's MLlib (Machine Learning library) came into existence. MLlib gives you the ability to train machine learning models on huge datasets, using Spark clusters. It allows you to build in supervised, unsupervised, and recommender systems; NLP-based models; and deep learning, as well as within the Spark ML library [Sin22].

### Structured Streaming

The Spark Streaming library provides the functionality to read and process real-time streaming data. The incoming data can be batch data or near real-time data from different sources. Structured Streaming is capable of ingesting real-time data from such sources as Flume, Kafka, Twitter, etc [Sin22].

### Graph X

This is a library that sits on top of the Spark core and allows users to process specific types of data (graph dataframes), which consists of nodes and edges. A typical graph is used to model the relationship between the different objects involved. The nodes represent the object, and the edge between the nodes represents the relationship between them. Graph dataframes are mainly used in network analysis, and Graph X makes it possible to have distributed processing of such graph dataframes [Sin22].

## Programming Language APIs

Spark is available in four languages. Because Spark is built using Scala, that becomes the native language. Apart from Scala, we can also use Python, Java, and R [Sin22].

## Spark Execution

Any Spark application spins off a single driver process (that can contain multiple jobs) on the master node that then directs executor processes (that contain multiple tasks) distributed to a number of worker nodes shown 9.

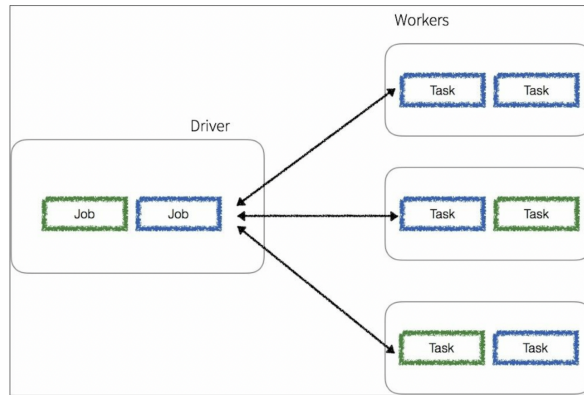


Figure 9: Think of something to say [DL17].

The driver process determines the number and the composition of the task processes directed to the executor nodes based on the graph generated for the given job. Note, that any worker node can execute tasks from a number of different jobs [DL17].

## Spark vs Hadoop

| Hadoop Map Reduce  | Spark   |
|--|---|
| For Applications that repeatedly reuse the same set of data, map reduce is very inefficient. | Spark uses in-memory processing, reusing it for faster computation.                           |
| MapReduce is quite faster in batch processing.   | As memory size is limited, it would be quite slower in batch processing of huge data set.     |
| Data is stored in disk for processing.   | Data is stored in main memory. As it is an inmemory computation engine entire data is copied. |
| Difficulty in processing and modifying data in real time due to its high latency.            | Used to process and modify data in real time due to its low latency.                          |
| Predominantly used to process from bygone datasets.  | Predominantly used for streaming, batch processing and machine learning                       |
| For fault tolerance, MapReduce uses replication.   | For fault tolerance, Spark uses RDDs.   |
| It merges and partitions shuffle files.  | It does not merges and partition shuffle files.   |
| Primarily disk based computation.  | Primarily RAM based computation.  |

Table 4: Showing the differences between haddop and spark [HRJ17].

| Number of words | Hadoop (Sec) | Spark(Sec) |
|-----------------|--------------|------------|
| 100             | 79           | 28.841     |
| 1000            | 91           | 31.185     |
| 10000           | 96           | 35.181     |
| 100000          | 103          | 36.969     |
| 1000000         | 116          | 39.569     |

Table 5: Comparision of Execution time for wordcount program [HRJ17].

| Number of words | Hadoop (Sec) | Spark(Sec) |
|-----------------|--------------|------------|
| 5               | 2.541        | 0.9030     |
| 10              | 3.370        | 1.459      |
| 50              | 6.420        | 2.840      |
| 100             | 9.383        | 3.452      |
| 200             | 10.100       | 5.749      |

Table 6: Comparison of Execution time for logistic regression program [HRJ17].

Summarising the results shows Spark to be quicker in both experiments. Spark also provides an API for python which will be very helpful in this project seeing its easy nature to be able to read files and work with text-based files. Therefore I have decided to work with Pyspark for this project.

## Software Architectural Bottlenecks

HDFS has scheduling delays in the architecture which results in cluster nodes waiting for new tasks as the access pattern is periodic.HDFS client code, serializes computation and I/O instead of decoupling and pipelining those operations. [SRC10].

**Definition 3.9 (HDFS)** *The Hadoop Distributed File System (HDFS) is a distributed file system designed to run on commodity hardware*

## Portability Limitations

Some performance-enhancing features in the filesystem are not available such as bypassing the filesystem page cache and transferring data directly from the disk into user buffers. Thus, HDFS implementation runs less efficiently and has higher processor usage than would otherwise be necessary [SRC10].

### 3.5 Analysis of existing systems that solve similar tasks

There are a few systems that solve similar tasks these include:

- SparkMD: This is a Spark-based framework for molecular dynamics simulations. It uses a distributed version of the GROMACS simulation engine to enable large-scale simulations of protein systems.
- BioSpark: This is a Spark-based framework for processing and analyzing large-scale genomics and proteomics datasets. It provides a set of APIs for analyzing DNA sequences, protein structures, and other biological data.
- Pysparkling: This is a Python-based library for parallelizing computations using Spark. It can be used for a range of bioinformatics analyses, including sequence alignment, protein structure prediction, and gene expression analysis.
- SparkProt: This is a Spark-based framework for protein structure prediction. It uses a combination of machine learning algorithms and structural bioinformatics tools to predict the 3D structure of proteins from their amino acid sequences.
- DeepChem: This is a deep learning-based framework for drug discovery and development. It uses Spark to parallelize computations and can be used for a range of tasks, including protein-ligand docking, virtual screening, and compound synthesis planning.

## 4 Software Engineering

### 4.1 Objective Implementation

Out of all the objectives listed in section 2.2 I will go through each one and show my implementation, thoughts, issues and any technical decisions i had to make whilst trying to achieve the objective.

#### **MapReduce Framework for Protein Analysis**

Develop a software framework that supports the Mapreduce formalism and can process large numbers of protein structures.

To implement this we need to break down the objective into two parts the first part is to build a framework that supports a mapreduce formalism and the second part is to ensure that the build can process large number of protein structures.

##### **1. Framework that supports a mapreduce formalism**

The first decisions i have to make is to decide what software to use which can enable me to use a mapreduce formalism I concluded with two options, hadoop and apache spark. Please refer to my Literature review on Hadoop spark and pyspark in summary spark is newer and performs quicker whilst having an API for python. Which ultimately led me to choose spark over hadoop.

## Implementing PySpark

**Install Java:** PySpark requires Java to run. You can download and install the latest version of Java from the official website (<https://www.oracle.com/java/technologies/javase-downloads.html>).

**Install Apache Spark:** PySpark is built on top of Apache Spark, so you need to install Apache Spark first. You can download the latest version of Apache Spark from the official website (<https://spark.apache.org/downloads.html>). Choose a pre-built package for Apache Spark and select the version that matches your installed Java version. Extract the downloaded package to a location of your choice.

**Install Python:** PySpark requires Python 3.x to run. You can download and install the latest version of Python from the official website (<https://www.python.org/downloads/>).

Set environment variables: To use PySpark, you need to set environment variables for Java and Spark. On MAC OS and linux open the terminal and run the following commands:

```
export JAVA_HOME=/Library/Java/
JavaVirtualMachines/JDKversion/Contents/Home
export PATH=$JAVA_HOME/bin:$PATH
export SPARK_HOME=<path_to_spark_folder>
export PATH=$SPARK_HOME/bin:$PATH
```

On windows open command prompt and run the following commands:

```
setx JAVA_HOME "C:\Program Files\Java\jdk-16.0.2"
setx PATH "%PATH%;%JAVA_HOME%\bin"
setx SPARK_HOME "C:\spark-3.2.0-bin-hadoop3.2"
setx PATH "%PATH%;%SPARK_HOME%\bin"
```

Replace 'JDK version' with the version of Java you installed and 'path to spark folder' with the path to the folder where you extracted Apache Spark.

**Install PySpark:** You can install PySpark using pip. Open the terminal and run the following command:

```
pip install pyspark
```

**Finally Test PySpark:** You can test PySpark to check if all is set up correctly by opening the Python shell and running the following commands:

```
from pyspark.sql import SparkSession
spark = SparkSession.builder.appName("test").getOrCreate()
df = spark.range(10)
df.show()
```

The above code creates a `SparkSession`, which is the entry point to Spark, a distributed computing framework that supports a MapReduce formalism.

In particular, the following code creates a `SparkSession` with the name "test". The `SparkSession` is the entry point to Spark and provides a way to interact with Spark's distributed data processing capabilities. Once the `SparkSession` is created, the code generates a `DataFrame` using the `range()` method, which creates a `DataFrame` with a single column named "id" and ten rows with values from 0 to 9. Finally, the code prints the contents of the `DataFrame` using the `show()` method. This displays the contents of the `DataFrame` in a tabular format. Overall providing a Framework that supports a mapreduce formalism.

## 2. Process large numbers of protein structures

The second part of the objective is to ensure that we can process large numbers of protein structures. To do this we need to optimize our Spark job and cluster configuration. There are actions that fall outside of our ability to execute, while others can be considered during the planning phase before coding, and some can be implemented directly into the code itself.

### Can not implement

**Cluster Configuration:** I initially aimed to use a YARN implementation which allows you to dynamically share and centrally configure the same pool of cluster resources between all frameworks that run on YARN. [goy18] These are the issues i ran across:

1. Configuration: Yarn requires additional configuration beyond what is required for a standalone Spark cluster. You need to configure Yarn-specific settings such as memory allocation and container sizes, as well as Spark-specific settings such as executor and driver memory.
2. Resource Management: Yarn is responsible for managing resources for multiple applications running on a shared cluster. This can make it more difficult to manage resource allocation and ensure that each application has access to the resources it needs.
3. Monitoring: Because Yarn is managing resources for multiple applications, it can be more difficult to monitor the performance of individual applications. You need to look at both Yarn-level metrics and Spark-level metrics to get a complete picture of performance.
4. Troubleshooting: When issues arise in a Yarn-managed cluster, it can be more difficult to troubleshoot the problem. You need to determine whether the issue is related to Yarn or Spark, and then diagnose the problem accordingly.

Due to a smaller-scale workload and a simpler, more flexible, and more cost-effective option, a standalone Spark cluster may be a better fit.

**Data Serialisation:** Serialisation is the process of converting data structures into a format that can be easily transmitted over the network. By default, PySpark uses Python's pickle library for serialisation, which can be slow for large datasets. You can improve serialisation performance by using more efficient serialisation formats like Avro or Parquet. However i came across some issues:

Serialisation and deserialisation can introduce performance overhead due to the immense size and complexity of protien structure analysis data being serialized. While both are widely used, they may not have the same level of tooling and support as other formats, like CSV or JSON or txt. This can make it more difficult to work with data in these formats. Finally, both can add complexity to your data processing pipeline. For example, you may need to write custom serialization and deserialization logic, or configure your data processing systems to work with these formats.

Thus, I have decided to use Python's pickle library for serialization.

## Before implementing the code

**Hardware Configuration:** Ensure that your cluster has sufficient CPU, RAM, and disk resources to handle the workload. You can also use SSDs for faster disk access and faster I/O operations.

Thus, we need to ensure that the device running the framework meets the minimum criteria. To do this we need to look at the minimum requirements for running PySpark.

- A 64-bit operating system e.g., macOS, Linux, or Windows
- At least 8 GB of RAM
- A multi-core processor e.g., Intel Core i5 or i7

## Whilst Implementing code

These are some features of pyspark that we can implement into our code to ensure that optimize our Spark job and cluster configuration.

**Caching:** Caching is an optimization technique that involves storing intermediate results in memory to avoid recomputation. Caching is particularly useful when you have iterative algorithms that reuse the same data multiple times. You can use the cache function in PySpark to cache Resilient Distributed Datasets or DataFrames.

**Partitioning:** Partitioning the input data can significantly improve performance, especially for large datasets. Partitioning involves splitting the data into smaller chunks and processing them in parallel.

| Partitioning Function | Description  |
|-----------------------|--|
| hash()                | Hashes the keys and partitions the data based on the hash value of the key. It is the default partitioning function in PySpark.  |
| range()               | Partitions the data based on the range of the key values. For example, if the key values range from 1 to 100, and there are 4 partitions, the first partition will contain keys from 1 to 25, and so on. |
| round robin()         | This function partitions the data in a round-robin fashion. For example, if there are 3 partitions, the first partition will contain the first record, the fourth record, the seventh record, and so on. |
| custom()              | This function allows you to define your own partitioning logic by implementing the getPartition(key) method.   |

Table 7: Showing examples of pyspark partitioning functions and what they do.

### Amended code to handle large amounts of Protein data

```
proteinfiles = spark.read.format("text").load(path)
processedproteins = protein_files.rdd.map(processprotein)
```

These are two lines of code that will allow the ability to handle large amounts of protein data.

Where proteinfiles holds the information of the protein structure. We then would pass this into a map function that will conduct processes on the protein structure data. Finally performing a .collect() function will save the results of the processed data into processedproteins.

Ultimately using this with conjunction of the code used to test pyspark we end up with a framework that supports the Mapreduce formalism and can process large number of protein structures.

## Parallelized Protein Analysis with MapReduce

Implement a distributed computing system using MapReduce to parallelize protein structure analysis across multiple computing nodes.

To implement a distributed computing system using MapReduce to parallelize protein structure analysis across multiple computing nodes using PySpark, we need complete the following steps:

1. Setup a PySpark cluster with multiple worker nodes.
2. Write a Map function to parse the protein structure data and extract the relevant features.
3. Write a Reduce function to combine the results from the Map function.
4. Load the protein structure data into PySpark RDDs.
5. Create a function that can process protein structures.
6. Apply the Map function to each protein structure RDD in parallel.
7. Apply the Reduce function to combine the results from the Map function.
8. Save the output to a distributed file system.

The following code has been taken from my lines program which is intended to print the number of lines each pdb file contains which complete the steps above

```
from pyspark.sql import SparkSession
from pyspark.sql.types import StringType
import os
from pathlib import Path
import tempfile

def main(spark):
    directory = ("/PROJECT/Programs/Lines/PDBsDirectory/*")
    tempdirectory = "/PROJECT/Programs/Lines/PDBsfromRDD"

    rddkeyvalue = spark.sparkContext.wholeTextFiles(directory)

    def numberoflines(filename, filecontent):
        with tempfile.NamedTemporaryFile() as tmp:
            tmp.write(filecontent)
            os.system("wc -l "+ tmp.name)

    r = rddkeyvalue.map(lambda x: numberoflines(x[0], x[1]))
    r = r.collect()
if __name__ == '__main__':
    spark = SparkSession.builder.appName('PDB').getOrCreate()
    main(spark)
```

## PySpark cluster setup

The SparkSession object is a higher-level interface to create a Spark application with SparkContext, which provides a unified entry point to interact with the Spark cluster.

We use the builder API to create a SparkSession instance and set the application name to "PDB". The getOrCreate method creates a new SparkSession instance or reuses an existing one if one already exists. This creates a SparkSession with all the necessary configurations and settings for connecting to the Spark cluster.

When we execute the code, the SparkSession instance is created and the worker nodes are launched, enabling distributed computing across the cluster.

## Map function

The numberoflines function takes a folders path and the values within the protien structure and creates a temp file in which it writes the contents of the protien structure. The temp file is then used to get the number of lines with the shell command "wc -l 'tempfilename'".

Once the numberoflines function has been defined, we can apply it to each protein structure file in parallel using the Map function in Spark.

## Reduce function

The code provided does not include an explicit implementation of a Reduce function. However, you can write a Reduce function using PySpark's reduce function, which would result in the same responce if needed to return the number of lines.

## PySpark RDDs

The code provided, loads protein structure data by using the wholeTextFiles function to read in all the files in a directory as an RDD of key-value pairs, where the key is the filename and the value is the contents of the file.

## Save to distributed file system

The code provided does not include any code to save the output to a distributed file system. However, you can use PySpark's saveAsTextFile function to save the output as text files in a distributed file system.

## Issues and Technical decisions

With the code provided the process we are conducting is returning the number of lines for each pdb file provided. I originally set out to use a user executable called hoppscore. Which is a script written for the purpose to provide a score when passed in a pdb files. However, I came across some issues with setting the script up.

Here is an example of what an example hoppscore execution would look like:

```
hoppscore -r 1.7 -g 12 sample.pdb
```

Output is sent to STDOUT

```
hoppscore -r 1.7 -g 12 sample.pdb > sample.pdb.out
```

Program output is sent to STDOUT and looks like this:

```
!F – Favored , A – Allowed , U – Unfavored , D – Disallowed
!F > 612.7, 183.3 < A > 612.7, 0 < U > 183.3, D=0
!Avg: 183.290662650602, Sigma: 858.791452289234
!Input File: test.pdb
!Reference Database Resolution Limit: 17
!phi–psi pairs: 1
! SS AA FILENAME RES F/Avg Favored?
!
C D test.pdb 2 10.19 F 2
S A test.pdb 3 34.83 F 2
G P test.pdb 4 99.17 F 2
G F test.pdb 5 5.93 F 2
G E test.pdb 6 1.01 A 1
.
.
.
!
Sum: 111
Score: 1.23
```

However when attempting to install the program i failed to do so. The readme files show how to install the tool in different operating systems however i failed to do so when following the steps and there is no external help or documentation online for this tool. This can be because the program was last updated in 10/04/2013 showing that it has not been an active analysis tool recently. Thus, i decided to change the protein analysis function to return the number of lines instead.

## Optimization for Faster Protein Analysis

Optimize the software framework to reduce the processing time required for protein structure analysis.

I will now present the first version of my lines programs and explain features i changed so that the framework will be more optimized for reduce the processing time required for protein structure analysis.

```

from pyspark.sql import SparkSession
from pyspark.sql.types import StringType
import os
import shutil
from pathlib import Path

def main(spark):
    directory = ("/Programs/Lines/PDBsDirectory")
    tempdirectory = "/PROJECT/Programs/Lines/PDBsfromRDD"

    for file in os.listdir(os.fsencode(directory)):
        filename = os.fsdecode(file)
        if filename.endswith(".DS_Store"):
            continue
        rdd = spark.read.text("/PDBsDirectory/" + filename)
        rdd = rdd.map(lambda x: x[0])
        if os.path.exists(tempdirectory):
            shutil.rmtree(tempdirectory)
        rdd.saveAsTextFile(tempdirectory)
        os.system("wc -l " + tempdirectory + "/part-00000")

if __name__ == '__main__':
    # This creates a local cluster
    spark = SparkSession.builder.appName('PDB').getOrCreate()
    main(spark)

```

### Issues with code

We have instantiated a for loop which loops through all the files in the PDBsDirectory the code then reads each file in the PDBsDirectory directory and loads it into an RDD using spark.read.text ignoring any files that end with DSStore which is one of the outputs returned from saveAsTextFile. This operation can be time-consuming for large files because it involves reading the entire file into memory as a single string. This is not efficient for processing large files.

The code then saves the RDD to the PDBsfromRDD directory using rdd.saveAsTextFile but we need to check if this directory exists already, deleting it if so, as for the next pdb file in the PDBsDirectory we will need to cast saveAsTextFile to this which requires that folder to not exist in the first place. SaveAsTextFile function can be expensive because it involves writing the entire RDD to disk.

Finally if there is a lot of files in PDBsDirectory the the for loop will need to go through each one whilst also running shutil.rmtree which is not effeciant as there are some operations being run each iteration which we can remove.

## Improvement Using wholeTextFiles()

wholeTextFiles() can be used instead of textFile() to read all files within a directory as a single RDD. wholeTextFiles() reads the files in the specified directory and returns an RDD where each element is a tuple of (filename, contents), where filename is the name of the file and contents is the entire contents of the file as a single string.

This eliminates the need to have a for loop to go through the directory as we can read each file in the directory using wholeTextFiles() and a wildcard '\*' in the directory path for example directory = "/path/\*" which will select all the files in that path.

As we have the name and content of the file we can remove saveAsTextFile() as we can pass in the file name into the map function thus casting the shell command on the file.

This will result in the following code:

```
from pyspark.sql import SparkSession
from pyspark.sql.types import StringType
import os
import shutil
from pathlib import Path

def main(spark):
    start = time.time()
    directory = ("/PROJECT/Programs/Lines/PDBsDirectory/*")

    rddkeyvalue = spark.sparkContext.wholeTextFiles(directory)

    def numberoflines(k):
        os.system("wc -l "+ k[40:])

    rddkeyvalue.map(lambda x: numberoflines(x[0])).collect()

if __name__ == '__main__':
    # This creates a local cluster
    spark = SparkSession.builder.appName('PDB').getOrCreate()
    main(spark)
```

## Integrate API for PDB access

Integrate a PDB file download and search from the RCSB API for efficient data access.

As the software framework is dependent on what files are currently loaded into the correct folders this objective is important as it is an easy way for the researcher to download pdb files without having to access the website.

The objective is to work with the RCSB protein data bank website using there provided api. Please refer to my Literature review of the website for a better understanding of what it holds and the features it provides. In summary the RCSB Protein Data Bank website is a comprehensive resource that provides access to 3D structural data of proteins, nucleic acids, and complex assemblies. It is a valuable tool for researchers in the fields of biochemistry, biophysics, and molecular biology.

The RCSB site provides many apis including:

- Data API
- Search API
- ModelServer API
- VolumeServer API
- 1D Coordinate Server

The features that are required for this objective and are also the main features that power rcsb.org are, File Download Services which, serves to download the pdb file of a provided protein structure. Search API serves to find out what identifiers match a certain search condition.

## File Download Services

The file download service provides the ability to download a file that the website contains given the correct parameters. This is key as it will allow the reasercher to use the api to download any required files without needing to access the website.

The following url is an example on how to download a .pdb file in this case 4hhb.pdb:

<https://files.rcsb.org/download/4hhb.pdb>

There are a few different options when downloading files from the website:

| PDB entry files                        |              |                                 |
|--|--------------|---------------------------------|
| File Format                            | Compression  | Example URL                     |
| Biological Assembly File in PDB        | Uncompressed | /download/1hh3.pdb1             |
| Biological Assembly File in PDB        | Compressed   | /download/1hh3.pdb1.gz          |
| Biological Assembly File in PDBx/mmCIF | Uncompressed | /download/5a9z-assembly1.cif    |
| Biological Assembly File in PDBx/mmCIF | Compressed   | /download/5a9z-assembly1.cif.gz |
| PDB                                    | Compressed   | /download/4hhb.pdb.gz           |
| PDB                                    | Uncompressed | /download/4hhb.pdb              |
| PDBx/mmCIF                             | Compressed   | /download/4hhb.cif.gz           |
| PDBx/mmCIF                             | Uncompressed | /download/4hhb.cif              |
| XML                                    | Compressed   | /download/4hhb.xml.gz           |
| XML                                    | Compressed   | /download/4hhb.xml              |

Depending on what is put after the download in the URL we will get a different download. The table highlights some of the main File formats and examples on how to use the URL to download them.

### Implementing Download Service into Python

Using this we can retrieve the data for a given pdb file name and write it into a file and save it to the path given:

```
import requests

apicall = 'https://files.rcsb.org/download/{}.pdb'.format(pdbfile)

response = requests.get(apicall)

with open('path' + pdbfile + '.pdb', 'wb') as f:
    f.write(response.content)
```

The RCSB site holds an extremely vast amount of pdb files we can actually see all current PDB ids by going to:

```
https://data.rcsb.org/rest/v1/holdings/current/entry\_ids
```

This brings in the need for the search api. As it is not good to assume that the user will have the name for each file they would like to download.

## Search API

The Search API accepts HTTP GET or POST requests with JSON payloads. In GET method, search request should be sent as a URL-encoded query string in json parameter:

```
https://search.rcsb.org/rcsbsearch/v2/query?json={search-request}
```

A search request is a complete specification of what should be returned in a result set. The search request is represented as a JSON object. The building blocks of the request are:

| Context                    | Description  |
|----------------------------|--|
| return type (Required)     | Specifies the type of the returned identifiers, e.g. entry, polymer entity, assembly, etc.   |
| query (Optional)           | Specifies the search expression. Can be omitted if, instead of IDs retrieval, facets or count operation should be performed. In this case the request must be configured via the request options context.  |
| request options (Optional) | Controls various aspects of the search request including pagination, sorting, scoring and faceting. If omitted, the default parameters for sorting, scoring and pagination will be applied.  |
| request info (Optional)    | Specifies an additional information about the query, e.g. query id. It's an optional property and used internally at RCSB PDB for logging purposes. When query id is sent with the search request, it will be included into the corresponding response object. |

Table 8: Aspects that a valid json for the search api on RCSB can hold.

Here is an example valid json and the response for the results shown when searching for thymidine kinase:

```
"query": {
    "type": "terminal",
    "service": "full_text",
    "parameters": {
        "value": "thymidine kinase"
    }
},
"return_type": "entry"
```

This is the response received back:

```
"query_id": "f3424f3f-a69a-4b70-96ee-df9fb8db7558",
"result_type": "entry",
"total_count": 57547,
"result_set": [
    {
        "identifier": "2UZ3",
        "score": 1.0
    },
    {
        "identifier": "2B8T",
        "score": 0.9175522850503486
    },
    {
        "identifier": "2WVJ",
        "score": 0.898559256390395
    },
    {
        "identifier": "2J87",
        "score": 0.602571649883811
    },
    {
        "identifier": "2VTK",
        "score": 0.3646165762974438
    },
    {
        "identifier": "1XBT",
        "score": 0.2990240123934934
    },
    {
        "identifier": "1KIM",
        "score": 0.24848954298993028
    },
    {
        "identifier": "1E2J",
        "score": 0.18949651432997677
    },
    {
        "identifier": "1P6X",
        "score": 0.08545313710302091
    },
    {
        "identifier": "2QQE",
        "score": 0.0
    }
]
```

We get the pdb identifier and the score of how relatable it is to the search value. This is helpful as now we can in theory download each result as we have the identifier associated to them. For example: <https://files.rcsb.org/download/2UZ3.pdb>

## Implementing Search API in Python

Before starting the python program we need to create a valid json file. Using the examples provided from the RCSB API documentation found here <https://search.rcsb.org/basic-queries> We can set up a json file that looks like the following:

```
"query": {
    "type": "terminal",
    "service": "full_text",
    "parameters": {
        "value": "vinay"
    }
},
"return_type": "entry"
```

We now need to create a following python program that can use the following json file and execute a get request to the api. The following program was created:

```
import requests
import json
import urllib.parse
from os.path import exists
import sys

def main(folder, value):

    jsonfile = 'Search.json'
    directory = (path)

    with open(file=jsonfile, mode="r") as jsonFile:
        data = json.load(jsonFile)

        data['query']['parameters']['value'] = value

    with open(file=jsonfile, mode="w") as jsonFile:
        json.dump(data, jsonFile)

    #fix to make the values in data
    #not use ' qoutes but use "" qoutes instead
    data = json.dumps(data)

    newdata = urllib.parse.quote(data)
    apicall =
        'https://search.rcsb.org/rcsbsearch/v2
        /query?json={}'.format(newdata)

    result = requests.get(apicall)
    result = result.json()

    for x in result["result_set"]:
        print(x)
```

```

if __name__ == '__main__':
    globals()['sys.argv[1]'](*sys.argv[2], sys.argv[3])

#Example run line: python3 searchpdbfiles.py main PDBsDirectory2 vinay

```

In this program, we first define the URL of the RCSB PDB API endpoint. We then open and change the value of the search filter to what is provided and load the json file from a query dictionary to a JSON string using json.dumps(). The urllib.parse.quote() method is used to URL-encode the search query JSON string. The script sends a GET request to the API using the requests.get() method with the apicall URL. Finally retrieving the results and printing them.

### Issues with creating a valid JSON

I came across an issue when creating a valid JSON to pass into the Search API. Currently the JSON file is only allowing the search for one value. However there is a vast number of other filters and search options a researcher can use when browsing the pdb bank. Therefore I decided to consolidate with my supervisor to determine some useful filter options for the researcher to use whilst working with the framework. Two useful filters included:

- R Factor
- Date of Deposition

This required multiple filters in one JSON query the value, R factor and Date of Deposition. This was found difficult to create a valid JSON, as the documentation uses high level biology terms which made it hard to look for exactly what I needed. I first broke down the problem by looking at how to translate these filter terms into a valid JSON format. I found that:

- reflns.pdbx Rmerge I obs is a measure of the quality of a protein structure, specifically the average R-merge value of the observed reflections. R-merge is a statistical parameter that is used to evaluate the agreement between multiple observations of a diffraction spot. A lower value of reflns.pdbx Rmerge I obs generally indicates a better quality protein structure.
- rcsb accession info.initial\_release\_date is the date when a protein structure was first released to the RCSB PDB. This is the date when the protein structure was deposited in the PDB for the first time, and it is assigned by the depositor. The date is in the format YYYY-MM-DD.

The documentation provides a 'Open in Editor' button which allows an easy and effective way of editing json and sending it through to the api for a response.

Figure 10: Showing the User interface of the editor for the RCSB Search API.

It provides the user with error messages if the entered JSON on the left hand side is incorrect. However this was found to be not helpful as the messages were vague. Seeing this was not helping I decided to investigate the RCSB PDB bank website more specifically looking at the API requests being sent on the website by investigating them through the advanced search using the Network tab.

I first created a search using the advanced search button that fulfilled the filters I am attempting to create in json format.

Figure 11: Payload for when filtering through value date of deposition and rfactor

This payload provided me with a valid JSON with correct filters i require however, this JSON had lots of extra components that i did not require. So using the editor, documentation and valid json provided by the payload i created a valid json that filters on the three criterias Value, Rfactor and Date of Deposition which is the following:

```
"query": {  
  "type": "group",  
  "logical_operator": "and",  
  "nodes": [  
    {  
      "type": "group",  
      "logical_operator": "and",  
      "nodes": [  
        {  
          "type": "terminal",  
          "service": "full_text",  
          "parameters": {  
            "value": "covid"
```

```

        }
    },
{
    "type": "terminal",
    "service": "text",
    "parameters": {
        "attribute": "reflns.pdbx_Rmerge_I_obs",
        "operator": "greater",
        "negation": false,
        "value": 0.2
    }
}
],
},
{
    "type": "terminal",
    "service": "text",
    "parameters": {
        "attribute": "rcsb_accession_info.initial_release_date",
        "operator": "less",
        "value": "2019-08-20"
    }
}
],
},
"return_type": "polymer_entity"

```

## Explaining the Query

**query:** This is the main component of the JSON, which specifies the search criteria. It consists of three parts:

1. **type:** This specifies that the query is a group query, which means that it consists of multiple sub-queries that are combined using a logical operator.
2. **logical operator:** This specifies that the logical operator used to combine the sub-queries is "and". This means that all the sub-queries must be true for a protein structure to be included in the search results.
3. **nodes:** This is an array of sub-queries, each of which is also a group query with two sub-nodes. The first sub-node is a query to search for the term "covid" in any text field and the second sub-node is a query to filter for protein structures with a reflns.pdbx Rmerge I obs value greater than 0.2. The second group query is then combined with a third sub-node, which is a terminal query to filter for protein structures with an initial release date less than 2019-08-20.

**Finally, return type:** This specifies the type of entity that should be returned in the search results. In this case, the return type is "polymer entity", which means that the search results will only include polymer entities, which are protein or nucleic acid molecules with a defined sequence.

## User Interface for Protein Analysis

Design an interface that allows users to manipulate the pdbs that are being passed into the executable for protein structure analysis.

The framework is dependent on what files are present in the correct folder for the functions to work. This creates the need for a user interface for an easy approach to manipulate these folders to the researchers needs. Before beginning the user interface i needed some functions that will be used when interacting with the user interface. Here is a list of all the functions i would like the user interface to inherit:

- Clear all PDB files in folders
- Remove One or more specifc pdb file in folders
- Return all values within folder
- Add one or more specific pdb file in folders
- Search pdb file using API
- Search and download pdb file from API into folder

After implementing these functions in python we need to create a user interface that can inherit these functions.

## Implementing Functions For User Interface

There is no need to implement Search pdb file using API as we already have please refer to Intergrate API for PDB access objective implentation.

### Clear all PDB files in folders

This is the following program as the folder should only contain pdb files we dont need to specify what type of files to remove:

```
import os
import sys

def main(folder_path):
    # iterate through all files and subdirectories in the provided folder
    for root, dirs, files in os.walk(folder_path):
        for file in files:
            # delete the file
            os.remove(os.path.join(root, file))

if __name__ == '__main__':
    globals().__setitem__(sys.argv[1], sys.argv[2])

#Example run line: python3 emptypdbfolder.py main PDGsDirectory1
```

## Return all values within folder

This is the following program as the front end has not been created yet I have left the return as print but is subject to change:

```
import os
import sys

def main(folderpath):
    for file in os.listdir(folderpath):
        print(file)

if __name__ == '__main__':
    globals()[sys.argv[1]](sys.argv[2])
#Example run line: python3 getcurrentpdbfiles.py main PDBsDirectory1
```

## Search and download pdb file from API into folder

This is the following program:

```
def main(folder, value):
    jsonfile = 'Search.json'
    directory = (path)
    with open(file=jsonfile, mode="r") as jsonFile:
        data = json.load(jsonFile)
    data['query'][ 'parameters'][ 'value'] = value
    with open(file=jsonfile, mode="w") as jsonFile:
        json.dump(data, jsonFile)
    data = json.dumps(data)
    newdata = urllib.parse.quote(data)

    apicall = 'https://search.rcsb.org/rcsbsearch/v2/query?json={}'.format(newdata)

    result = requests.get(apicall)
    result = result.json()
    listofresults = []
    for x in result["result_set"]:
        listofresults.append(x['identifier'])
    #Everything above is for getting the values of the search
    for pdbfile in listofresults:
        if exists(directory + pdbfile + '.pdb'):
            print(pdbfile + ': already exists')
            continue

    apicall = 'https://files.rcsb.org/download/{}.pdb'.format(pdbfile)

    response = requests.get(apicall)
    with open(directory + pdbfile + '.pdb', 'wb') as f:
        f.write(response.content)
if __name__ == '__main__':
    globals()[sys.argv[1]](sys.argv[2], sys.argv[3])
#Example run line: python3 getpdbfiles.py main PDBsDirectory2 vinay
```

**Technical Decisions: User Interface for Protein Analysis**

Unfortunatly i did not manage to complete this objective and i will discuss in more details in section 5. I am depriortizing this objective due to time constraints. I also believe that a better user interface is not a key objective when trying to achieve the aim of the project. The functions are currently giving the reasercher the ability to be able to use them through the terminal or command prompt giving some form of user interface to the reasercher when using the framework tool. The user interface is more of a addition feature to the framework making it easier to use. I plan to create a readme file which will in detail explain the functions of the program what they do and what they return and how to use them in the terminal.

So i have the made the technical decision to prioritize in finishing all other objectives before moving back on to finishing this objective.

## Scalable Data Handling for Protein Analysis

Ensure the software framework is scalable and can handle increasingly large datasets.

There are several things we can consider to ensure the software framework is scalable and can handle increasingly large datasets:

1. Partitioning data
2. Distributed computing
3. Efficient algorithms
4. Testing and validation

I will be using my TMalign program to show how I have implemented each one of these:

```
def main(spark):
    directory1 = ("PROJECT/Programs/TMalign/PDBsDirectory1/*")
    directory2 = ("PROJECT/Programs/TMalign/PDBsDirectory2/*")

    rddkeyvalue1 = spark.sparkContext.wholeTextFiles(directory1)
    rddkeyvalue2 = spark.sparkContext.wholeTextFiles(directory2)

    rdd = rddkeyvalue1.cartesian(rddkeyvalue2)

    def TMalign(tuple1, tuple2):
        with tempfile.NamedTemporaryFile() as tmp1,
             tempfile.NamedTemporaryFile() as tmp2:
            tmp1.write(tuple1[1])
            tmp2.write(tuple2[1])
            os.system("./TMalign " + tmp1.name + " " + tmp2.name)

    rdd.map(lambda tuple: TMalign(tuple[0], tuple[1])).collect()

if __name__ == '__main__':
    # This creates a local cluster
    spark = SparkSession.builder.appName('PDB').getOrCreate()
    main(spark)
```

## Partitioning data

The cartesian() operation results in an RDD with a number of partitions equal to the number of partitions in the two input RDDs multiplied together. By default, Spark uses hash partitioning for cartesian() operation, which hashes the keys of the input RDDs and distributes them evenly across partitions to ensure a roughly equal workload per partition.

## Distributed computing

As we are using MapReduce, this can help with scalability as the data is being broken down and performed in parallel. This can be seen with the map step where each node will be given a pair of pdb files which need tmalign run on. Overall the processing workload is spread out across multiple machines, allowing for faster processing and the ability to handle larger datasets. Additionally, if one machine fails or experiences an error, the rest of the system can continue running without interruption.

## Efficient algorithms

Some issues with the a earlier version of the tmalign program:

```
def main(spark):
    directory1 = ("~/PROJECT/Programs/TMalign/PDBsDirectory1")
    directory2 = ("~/PROJECT/Programs/TMalign/PDBsDirectory2")
    tempdirectory1 = ("~/PROJECT/Programs/TMalign/PDBsfromRDD1")
    tempdirectory2 = ("~/PROJECT/Programs/TMalign/PDBsfromRDD2")

    for file in os.listdir(os.fsencode(directory1)):
        filename = os.fsdecode(file)
        if filename.endswith(".DS_Store"):
            continue
        rdd = spark.read.text(filename).rdd.map(lambda x: x[0])
        if os.path.exists(tempdirectory1):
            shutil.rmtree(tempdirectory1)
        rdd.saveAsTextFile(tempdirectory1)

        for file in os.listdir(os.fsencode(directory2)):
            filename = os.fsdecode(file)
            if filename.endswith(".DS_Store"):
                continue
            rdd = spark.read.text(filename).rdd.map(lambda x: x[0])
            if os.path.exists(tempdirectory2):
                shutil.rmtree(tempdirectory2)
            rdd.saveAsTextFile(tempdirectory2)
            os.system("./TMalign part-00000 part-00000")

    if __name__ == '__main__':
        # This creates a local cluster
        spark = SparkSession.builder.appName('PDB').getOrCreate()
        main(spark)
```

Here we are going through each file in the first folder and saving each file into a rdd. We then save this rdd into a tempfile before iterating through to the next file in the folder we have another for loop for the second folder. This is to loop through and repeat the steps above for the second folder. As a result we end with two temp files which we can run tmalign on iterating through all the files in the second folder before moving on to the next file in the first folder.

This is not effeciant as we have nested for loops and we are having to read through the second file eberytime we go the the next file in the first folder.

Introducing wholeTextFiles() method allowed the program to only need to read each folder once. We then get the values from each key-value pair and perform the cartesean product on them. Giving us an rdd where each value in the first folder is paired with each value in the second folder. This helped utilize the memory and compute resources of a cluster efficiently which improved overall performance.

**Testing and validation:** Whilst developing the framework, using python unittests i was able to test and validate its performance on different sizes of datasets also looking at catching unexpected errors occurring. This helped identify areas where the code got improvements and ensured that it can handle increasingly large datasets. Here is an example Unittest i have created:

```
class Testgetpdbfiles(unittest.TestCase):

    def test_input_values(self):
        os.mkdir(self.directory)
        values = ["covid", "v", "vinay"]

        api_url='https://files.rcsb.org/download'

        for value in values:

            result = tempgetpdbfiles(self.directorywithoutmain,
                                     value, 'main/Search.json', api_url)

            self.assertIsNone(result)
            file_size = os.path.getsize(self.directory)
            self.assertGreaterEqual(file_size, 0)
            for filename in os.listdir(self.directory):

                file_path = os.path.join(self.directory,
                                         filename)

                try:
                    if os.path.isfile(file_path) or
                       os.path.islink(file_path):
                        os.unlink(file_path)
                    elif os.path.isdir(file_path):
                        shutil.rmtree(file_path)
                except Exception as e:

                    print(f'Failed to delete {file_path}.
                          Reason: {e}')

            os.rmdir(self.directory)
```

This is a test to check if the function returns the correct output for different input values. When using the getpdbfiles function (Search and Download pdb files off of the RCSB API) The test case aims to validate that the function works as expected by checking the following:

1. The function should not return any value (None) if it executes successfully.
2. The size of the downloaded files should be greater than or equal to zero.
3. The downloaded files should be deleted after the function execution.

The test case creates a directory and uses it as a target directory for the downloaded files. It then defines a list of values that the function will use to download the files. The loop iterates over the values and calls the tempgetpdbfiles() function with each value. After that, it asserts that the function returns None and checks the size of the directory where the files are saved. It also checks that all downloaded files have been deleted after the function execution.

Finally, the test case removes the target directory created at the beginning. If all the assertions pass without any errors, the test case passes, indicating that the function works as expected for the given input values. If any of the assertions fail, the test case fails, indicating that there is an issue with the function's implementation.

## Issues with setting up UnitTests

When setting up Unitests for my python programs i ran across an issue to do with relative path when running the code and running the unittest cases. Here is what my current directory looks like:

```
PROJECT
  /main
    /PDBsDirectory1
      1a0aA.pdb
    /PDBsDirectory2
      invalid.json
      search.json
      main.py
      test_main.py
```

When running my function directly from /main for example "python3 main.py getpdbfiles PDBsDirectory1 vinay" This would work as the relative folder path is correct and the line of code within the function getpdbfiles "directory = (""/Users/vinaykakkar/Desktop/PROJECT/main/" + folder + "")" would instansiate the correct directory. However when i ran the unittest for test main.py from /main passing in the value PDBsDirectory1 did not work as it is running the code from /PROJECT which was enforcing the test to fail as the relative path was not correct. Due to this a work around is to create a temp function that replicates the original functions. For any changes that occured to the function due to the test cases both functions were ammended equally.

## Example Bug Caught

An example change is shown where we use tempfiles over saveAsTextFile. As in the eleair version we did not have a key value pair we needed to use the saveAsTextFile method which spits out the contents of the rdd for us to be able to then run into the tmalign function. However we now have the contents of the pdv so we can write these to a temp file where once used the directory is closed and not used again. This saves reasources as we are not creating and deleting a permanent directory for each pdb file each iteration.

## Upgradable Framework for Advanced Analysis

Ensure the software framework can be easily updated to keep pace with advancements in protein structure analysis techniques and computing technology.

In order to achieve this objective we need to consider implementing a modular design that allows for easy integration of updates and improvements. We can also incorporate version control tools such as Git to manage changes in the codebase and ensure that the latest updates are available to users.

### Modular Design

Within my software I have ensured that:

- Each module should perform one specific task or functionality independently
- Modules should have well-defined interfaces for communication with other modules
- Modules should be relatively self-contained, i.e., they should not have dependencies on other modules
- Each module have unit tests to ensure its correct functionality
- Modules should be designed to be reusable

### Git

Within git these actions make sure that the software framework can be easily updated to keep pace with advancements:

- We have a repository where the framework code will be uploaded to and managed.
- After each update or modification to the codebase, commit the changes to the repository with a clear commit message describing the changes made.
- Merge code to the repository's main branch after verifying that it is stable.
- Publish documentation in the form of a README file or wiki, providing detailed information about how to interact with the software and its requirements.

By implementing modular design and using Git best practices, you can ensure that the codebase remains a high-quality, reliable, and scalable software framework with the latest updates available to users.

## User Support for Protein Analysis Software

Provide documentation and user support to enable researchers to use the software framework effectively

To complete this objective i need to provide a readme file that completes these points:

- Overview of framework
- Installation
- Usage
- Examples

To do this i have created a readme file that fulfils each point. The Readme file will Provide a brief overview of your software framework, including its purpose and functionality. Provide clear and concise instructions on how to install your software framework, including any prerequisites that must be installed. Describe how to use your software framework, including any input/output file formats and command line options. Provide examples that demonstrate the use of your software framework, including sample input files and the expected output.

## 5 Project Framework

Within this section i will be issuing a guide to how to setup the framework whilst providing function and non-functional requirements. I will also creating a video showing the functions and framework running. I will also be walking through the repository and explaining the files stored in there. Finally, i will be going through a setup guide to how to install the program.

### 5.1 Directory

```

PROJECT
  /Notes
  /Programs
    /Lines
    /TMalign
    /TempFileTest
  /ProofofConcepts
    /PDBontaCluster
    /TmAlign
  /Python_Tutorial
    /ApiTests
    /Cluster_Tutorial
    /MapReduce_Tutorial
    /TestProject
  /Reports
    /Interim_Report
    /Project_Report
    /Protein_Data_Bank
    /Protein_Structures
    /Template
    /latex
/main

```

PROJECT holds all of the work i have conducted.

Notes contains all of the information i collected on the required background reading. I used these notes to help create the literature review. Python Tutorials folder contains all the tutorials i completed in which helped me understand the basics of features. This includes using API's, setting up clusters, performing mapreduce functions and Test Project used to help me understand the basics of pyspark.

ProofofConcepts folder contains two programs PDBontaCluster which proves distributing protein structure files amongst MapReduce cluster and, TmAlign which proves we can MapReduce using a single type of executable for analysing protein structures.

Programs contains three programs performing two executables Lines returns the number of lines and TmAlign performs its script on two protein structures. TempfileTest was a test to use a temp file on the Lines program.

Reports contains all of the reports i have created for this project. Including my Interim Report, Project Report, Report on Protein Data Bank, Report on Protein Structures, a template for the Project Report, and finally, a latex test for setting up purposes.

Lastly we have our main folder, this folder contains all of the available functions in separate

python files, a main python file that contains all of the available functions in one file, unit tests that have tests created to be run on the main python file and finally two benchmarking python programs intended for liens and TMalign.

## 5.2 Framework

The python file in PROJECT/main/main.py contains the framework with all the available functions.

### Functional Requirements

- The code should be able to count the number of lines in each file of a given directory.
- The code should be able to align two sets of files using the TMalign algorithm.
- The code should be able to search for PDB files on the RCSB PDB database using a given value.
- The code should be able to download PDB files from the RCSB PDB database for a given value and store them in a specified directory.

### Non-Functional Requirements

- The code should be written in Python3.
- The code should use the PySpark library for processing large datasets.
- The code should be easily understandable and maintainable by developers.
- The code should have proper error handling and should raise appropriate exceptions when errors occur.

### Setup Guide

**Install Git:** If Git is not already installed you can download it from the Git website (<https://git-scm.com/downloads>) and follow the installation instructions.

**Clone the Repository:** Clone the repository by running the following command: `git clone https://gitlab.cim.rhul.ac.uk/zfac214/PROJECT/-/tree/main`

**Install Required Dependencies:** Navigate to the README.md which lists the dependencies required for the program. Install any necessary dependencies using the appropriate package manager for example pip install.

**Run the Program:** Once all the dependencies are installed run the program using the appropriate command you can see the correct commands using the README.md file.

### 5.3 Example Results

It is important to note that the program must be run within the /main folder and that the folders used within the framework must be stored in the /main folder.

I will provide some examples of running functions showing how to run the function and the expected result. The available functions include:

- linecount
- tmalign
- searchpdbfiles
- getpdbfiles
- emptypdbfolder
- getcurrentpdbfiles

The current formula to run any of these functions available in the main.py file is the following:

```
python3 main.py <Function name> <Parameters>
```

To see what parameters are needed for the function please refer to the README file in the /main folder.

For the following examples please note that the /main folder contains two folders PDBsDirectory1 and PDBsDirectory2 which contain the following .pdb files respectfully: 1a0aA, 1apq, 1aqb and 1aqt, 1aua, AF-A0A5E8GAP1-F1-model.

#### **linecount**

An example for linecount function where the parameter takes the folder of which contains all the pdb files:

```
python3 main.py linecount PDBsDirectory1
```

This will result in the following output:

```
main % python3 main.py linecount PDBsDirectory1
499 PDBsDirectory1/1a0aA.pdb (0 + 1) / 1]
416 PDBsDirectory1/1apq_.pdb
1426 PDBsDirectory1/1aqb_.pdb
```

The output shows the number of lines for each file within the PDBsDirectory1 folder. We can identify what file it is as it displays the files name.

## tmalign

An example for tmalign function where the parameter takes two folders which contains the pdb files:

```
python3 main.py tmalign PDBsDirectory1 PDBsDirectory2
```

This will result in the following output please note that each file PDBsDirectory1 will be run through the tmalign script with each file in PDBsDirectory2 thus in this example we get nine results so i will only display the first one:

```
main % python3 main.py tmalign PDBsDirectory1 PDBsDirectory2
```

```
Name of Chain_1: /var/folders/bl/4cf/T/1a0aA.pdb1iu4dqlf  
(to be superimposed onto Chain_2)
```

```
Name of Chain_2: /var/folders/bl/4cf/T/1aqt_.pdbjzohaiuy
```

```
Length of Chain_1: 63 residues
```

```
Length of Chain_2: 135 residues
```

```
Aligned length= 27, RMSD= 1.51,
```

```
Seq_ID=n_identical/n_aligned= 0.074
```

```
TM-score= 0.36843
```

```
(if normalized by length of Chain_1, i.e., LN=63, d0=2.71)
```

```
TM-score= 0.18532
```

```
(if normalized by length of Chain_2, i.e., LN=135, d0=4.32)
```

```
(You should use TM-score normalized  
by length of the reference structure)
```

```
(": denotes residue pairs of  
d < 5.0 Angstrom, ". denotes  
other aligned residues)
```

---

MK

---

RES H

KHAEQARRNRALAVLHELASLIPAEWKQQNVSAAPSKATTVEAACRYIRHLQQNGST

.:: ::::::::::::::::::::

—STYHLDVVSAEQQMFSGLVKEIQVTGSEGEGLGIYPGHAPLLTAIKPGMIRIVKQH  
GHEEFIYLGGILEVQPGNVTVLADTAIRGQDLDEARAMEAKRKAEEHISSSHGDVD  
YAQASAELAKIAQLRVIELTKK—

Total CPU time is 0.00 seconds

The output provides information about the alignment between the two protein structures.

The quality of the alignment is then assessed using various metrics. The TM-score is a measure of structural similarity between the two proteins, with higher TM-score values indicating better alignment.

The output also provides a visual representation of the alignment, with colons.

Finally, the total CPU time required for the alignment is also reported.

### **searchpdbfiles, getpdbfiles, emptypdbfolder, getcurrentpdbfiles**

As all the functions follow the same pattern please refer to the README file to determine what parameters the function expects it provides what parameters it needs and example executions whilst also explaining what to expect as an output from the function. I will provide an example output for each of the remaining functions:

**searchpdbfiles** is used to return results from the RCSB API. It is to replicate what would be returned if we search on the website:

```
main % python3 main.py searchpdbfiles vinay
[{'identifier': '1DGO', 'score': 1.0}]
```

Over here we search with the key word vinay and get a result with a protein structure with the identifier 1DGO. We can check if this is correct but searching the same value on the RCSB website.

**getpdbfiles** provides a download feature for the every identifier returned by the api which is determined by the value:

```
main % python3 main.py getpdbfiles PDBsDirectory1 vinay
```

We search with the keyword vinay as we have seen the identifier returned is 1DGO. We also mention what folder we would like to download it into. Thus we get the file 1DGO.pdb downloaded into the folder PDBsDirectory1.

**getcurrentpdbfiles** returns a list of all the pdb files in a folder:

```
main % python3
>>> from main import getcurrentpdbfiles
>>> getcurrentpdbfiles("PDBsDirectory1")
['PDBsDirectory1/1DGO.pdb', 'PDBsDirectory1/1a0aA.pdb',
 'PDBsDirectory1/1aqb_.pdb', 'PDBsDirectory1/1apq_.pdb']
```

We have the correct files being displayed here.

**emptypdbfolder** is created to remove and empty a folder provided:

```
main % python3 main.py emptypdbfolder PDBsDirectory1
OriginalPDBscopy
```

After running this function it emptied the PDBsDirectory1 folder.

## 6 Project Achievements

A list of all the things achived in my project:

- PySpark
- Cluster Setup
- Key Value Pair
- TMalign
- API Service Download
- API Get
- Benchmarking

### 6.1 The Project

The project was quite successful all the objectives were implemented in a detail manner expect for the user interface. Overall the reasercher is able to search and download pdb files from the framework. The reasercher is also able to call out two user executables on the pdbs they require more specifically these are implemented useing a mapreduce formalism. Whilst having some functions that help manipulate the folders holding the pdb files. Ultimatlty, at the end of this project we have a software framework that suppots the Mapreduce formalism and can process large numbers of protein structures. Whilst being able to do so by parallelising protein structure analysis across multiple computing nodes. Keeping in mind that the framework is optimized to reduce the processing time required for protein structure analysis.

Finally the software framework is scalable and can handle increasingly large datasets and can be easily updated to keep pace with advancements in protein structure analysis techniques and computing technology.

### 6.2 Project Reflection

All set out obejectives were completed in order and occording to plan. However seting up the clusetr with correct rdd and key value pairs took much more time then expected as at first i was trying to use just the data of the pdb file as the rdd. For this reason i didnt have enough time to create a good interface keeping at a bash level of interaction for the reasercher. Due to this i have tried to prioteise a detailed reademe file which helps the reasercher understand the functions within the framework however a more user frienly user interface would be benifitil for eaxmple when running the functions that implement the api to search or download pdbs a better user interafce can be created that would be easier to use such as to set up the pdb files the reasercher would like to run the executables on to.

### 6.3 conclusions or results analysed or discussed

Need to include benchmarking here

## 7 Professional Issues: 10 marks

### 7.1 Should be a topic relevant to the project undertaken.

Structural bioinformatics involves the use of complex data analysis techniques and tools that require a high level of accuracy and reliability. As such, it is important to consider ethical issues related to data privacy, ownership, and sharing. It means that there could potentially impact on public health.

Public health is an important consideration in the field of structural bioinformatics, as the analysis of biological molecules at a molecular level can lead to a greater understanding of diseases and potential treatments. For example, structural bioinformatics has been used to develop new drugs, vaccines, and therapies that can improve public health outcomes.

Structural bioinformatics combines computational and experimental techniques to study the structure and function of biological molecules at a molecular level, requiring the analysis of large amounts of complex data and the use of advanced tools and techniques to extract meaningful information.

Given the sensitive nature of the data involved, ethical issues related to data privacy, ownership, and sharing are important.

Data privacy is a crucial aspect of structural bioinformatics as it involves the handling of personal data. Researchers must ensure that the data they collect is kept confidential and used only for the purpose for which it was collected. Appropriate measures must also be taken to protect the data from unauthorized access or disclosure. These measures can consist of researchers must ensure that any sensitive data they collect (such as genomic data) is kept confidential and not disclosed to unauthorized parties. Appropriate measures must be taken to protect the data from unauthorized access, such as using encryption or secure storage methods. This means that researchers must take appropriate measures to prevent data breaches and ensure that the data they collect is secure. If a breach does occur, researchers must take prompt action to minimize the harm to individuals and organizations whose data has been compromised.

Ownership of data is another ethical issue that must be carefully considered. Researchers must respect the rights of the individuals or organizations that provide the data and ensure that they are given appropriate credit for their contributions. The potential commercial value of the data must also be considered to ensure that researchers do not exploit it for their own gain. To do so Researchers must respect the rights of the individuals or organizations that provide the data, and ensure that they are properly credited for their contributions. For example, Researchers need to obtain informed consent from participants before collecting their data, and clearly explain the purpose and potential risks of the study. Participants should also be informed about their rights to access and control their data. If the data has commercial value, researchers must consider the ethical implications of using it for their own gain, and whether they need to obtain consent or provide compensation to the data providers. Another important aspect to ownership is transparency such that researchers should be transparent about their data collection and analysis methods, and make their findings publicly available whenever possible. This can help ensure that the scientific community and the public at large are aware of the potential benefits and risks associated with structural bioinformatics research.

Sharing of data is a fundamental aspect of structural bioinformatics, as it allows researchers to collaborate and share knowledge. However, sharing of data must be done in a responsible and ethical manner, with appropriate safeguards in place to protect the privacy and confidentiality

of the data. When sharing data, researchers should establish data sharing agreements that outline the terms and conditions of the data sharing arrangement, including who has access to the data and how it can be used. Researchers must ensure that appropriate safeguards are in place to protect the privacy and confidentiality of the data, such as using data use agreements or anonymization techniques. Such that, anonymization techniques can be used to protect the privacy of individuals and organizations whose data is being used in structural bioinformatics research. Researchers can remove identifying information from the data, such as names or addresses, to prevent it from being traced back to specific individuals.

Policies are important to consider too where we have:

Data access policies: Institutions and organizations that collect and share data should establish data access policies that outline who can access the data and under what circumstances. These policies can help ensure that the data is used in a responsible and ethical manner.

Data retention policies: Institutions and organizations should establish data retention policies that dictate how long data can be stored and when it should be destroyed. This can help ensure that data is not used for unintended purposes or retained longer than necessary.

In conclusion, as the Structural Bioinformatics Framework is examined within the context of a MapReduce formalism, it is crucial to consider the ethical implications of this field. The potential impact of structural bioinformatics on public health is significant, but the responsible management of data privacy, ownership, and sharing must be prioritized to ensure the ethical and responsible advancement of this field.

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