A framework for large scale phylogenetic analysis

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Abstract

With growing exchanges of people and merchandise between countries, epidemics have become an issue of increasing importance and huge amounts of data are being collected every day. Hence, analyses that were usually run in personal computers are no longer feasible. It is now common to run such tasks in High-performance computing (HPC) environments and/or dedicated systems. On the other hand, we are often dealing in these analyses with graphs and trees, and running algorithms to find patterns in such structures. Hence, although graph oriented databases and processing systems can be of much help in this setting, as far as we know there is no solution relying on these technologies to address large scale phylogenetic analysis challenges. This work aims to develop a modular framework that exploits such technologies, namely Neo4j. We address this challenge by proposing and developing a framework which allows representing large phylogenetic networks and trees, as well as ancillary data, that supports queries on such data, and allows the deployment of algorithms for inferring/detecting patterns and pre-computing visualizations, as a Neo4j plugin. This framework is innovative and brings several advantages to the phylogenetic analysis process, like the management of the phylogenetic trees, which will avoid having to compute them again, and the use of multilayer networks, that will make the comparison between them more efficient and scalable. The experimental evaluation results showcase that it can be very efficient in the mostly used operations and that the supported algorithms comply with their time complexity.

Key Words: Phylogeny; data processing; data storage; graphs; database;
Resumo

Na atualidade, trocas de pessoas e mercadorias entre diferentes países tem vindo a aumentar. Como consequência as epidemias tornaram-se uma preocupação maior, resultando na recolha de grandes quantidades de dados todos os dias. As análises que normalmente eram executadas em computadores pessoais já não são viáveis. Agora é comum executar essas análises em ambientes HPC e/ou sistemas dedicados. Por outro lado, nessas análises lidamos frequentemente com gráficos, árvores e com execuções de algoritmos para encontrar padrões nestas estruturas. Embora existam base de dados orientadas a grafos e sistemas de processamento que podem ajudar neste tema, não conhecemos nenhuma solução baseada nestas tecnologias para lidar com os desafios da análise filogenética em larga escala. O objetivo deste projeto é o desenvolvimento de uma plataforma que explore estas tecnologias, nomeadamente o Neo4j. Nós abordamos este desafio com a proposta e o desenvolvimento de uma plataforma que permita a representação de grafos e árvores filogenéticas de maior dimensão, bem como dados auxiliares, que suporta consultar esses dados e que permita a execução de algoritmos, para inferir/detectar padrões e pré-computar visualizações, como plugins do Neo4j. Esta plataforma é inovadora e traz vantagens para a análise filogenética, como por exemplo, o armazenamento dos grafos, que evita ter que computá-los novamente, e o uso de redes multi camadas, que torna a comparação entre eles mais eficiente. A análise dos resultados experimentais mostra que a plataforma pode ser muito eficiente nas operações mais utilizadas e que os algoritmos suportados obedecem à sua complexidade de tempo.

Palavras Chave: Filogenia; processamento de dados; armazenamento de dados; grafos; base de dados;
Contents

List of Figures viii
List of Tables ix
List of Acronyms xi

1 Introduction 1
  1.1 Objectives . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 2
  1.2 Document Structure . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 2

2 Background 3
  2.1 Graph Databases . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 3
    2.1.1 Use Cases . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 4
    2.1.2 RDBMS Comparison . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 4
    2.1.3 Graph Databases . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 5
  2.2 Large Scale Phylogenetic Analysis . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 7
    2.2.1 Phylogenetic Data . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 8
    2.2.2 Microbial Typing . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 8
    2.2.3 Algorithms . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 10
  2.3 Discussion . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 12

3 Proposed Solution 13
  3.1 Functional Requirements . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 13
  3.2 Non Functional Requirements . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 13
  3.3 Use Cases . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 14
  3.4 Architecture . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 15
    3.4.1 Data Model . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 15
    3.4.2 Client Server . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 20
    3.4.3 API . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 20
    3.4.4 Plugin . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 22
  3.5 Technology . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 24
    3.5.1 Spring Framework . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 24
    3.5.2 Google Identity Provider . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 25
    3.5.3 Neo4j . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 26
  3.6 Discussion . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 27
## 4 Implementation

4.1 Structure .................................................. 29

4.2 API ......................................................... 29
   4.2.1 Interface .............................................. 30
   4.2.2 Formatters ........................................... 31
   4.2.3 Errors ............................................... 31
   4.2.4 Security ............................................. 32
   4.2.5 Controllers ......................................... 34
   4.2.6 Services ............................................. 34
   4.2.7 Repositories ........................................ 36

4.3 Plugin ...................................................... 37
   4.3.1 Algorithms ......................................... 38
   4.3.2 Procedures ......................................... 39
   4.3.3 Services ............................................. 40
   4.3.4 Repositories ........................................ 40

4.4 Discussion ................................................ 40

## 5 Experimental Evaluation

5.1 Performance Testing ....................................... 44

5.2 Load Testing ............................................... 46

5.3 Discussion ................................................ 47

## 6 Final Remarks

6.1 Conclusions ............................................... 49

6.2 Future Work ............................................... 50

References .................................................... 51

Appendices ..................................................... 55
## List of Figures

| Figure | Description | Page |
|--------|-------------|------|
| 2.1    | Example of alleles. | 8    |
| 2.2    | Example of ancillary details. | 9    |
| 2.3    | Example of Multilocus Sequence Typing (MLST) profiles. | 10   |
| 2.4    | Example of Single Nucleotide Polymorphism (SNP) profiles. | 10   |
| 2.5    | Example of a distance matrix. | 10   |
| 2.6    | Example of an evolutionary hypothesis. | 11   |
| 3.1    | Use cases. | 14   |
| 3.2    | Graph data model architectural view. | 16   |
| 3.3    | Example representation of phylogenetic data. | 17   |
| 3.4    | Example representation of typing data. | 18   |
| 3.5    | Example representation of algorithms data. | 19   |
| 3.6    | Client-Server architectural view. | 20   |
| 3.7    | Layered architectural view. | 21   |
| 3.8    | Decomposition architectural view. | 22   |
| 3.9    | Uses architectural view. | 23   |
| 3.10   | Layered architectural view. | 23   |
| 3.11   | Decomposition architectural view. | 24   |
| 3.12   | Uses architectural view. | 25   |
| 3.13   | Authentication process using an identity provider. | 26   |
| 4.1    | Formatter generalization architectural view. | 32   |
| 4.2    | Security generalization architectural view. | 33   |
| 4.3    | Controller generalization architectural view. | 34   |
| 4.4    | Service generalization architectural view. | 35   |
| 4.5    | Repository generalization architectural view. | 37   |
| 4.6    | Plugin generalization architectural views. | 38   |
| 4.7    | Algorithms generalization architectural view. | 39   |
| 5.1    | Performance testing time plot. | 45   |
| 5.2    | Performance testing memory plot. | 46   |
| 5.3    | Load testing time plot, simulating several parallel users. | 47   |
| 1      | Example of a workflow system. | 57   |
List of Tables

2.1 Finding extended friends in a relational database versus efficient finding in Neo4j. Adapted from [46]. ................................................................. 5
2.2 Databases comparison summary. ...................................................... 5
5.1 Performance testing time results in milliseconds. .............................. 44
5.2 Performance testing memory results in megabytes. ............................ 45
5.3 Load testing, simulating several parallel users, time results in milliseconds. 47

1 Community Detection Algorithms. ...................................................... 57
2 Centrality Algorithms. ....................................................................... 58
3 Path finding Algorithms. ..................................................................... 59
4 Similarity Algorithms. ........................................................................ 60
5 Link Prediction Algorithms. ................................................................. 60
List of Acronyms

**HPC** | High-performance computing
---|---
**RDBMS** | Relational Database Management System
**CRUD** | create, read, update, and delete
**OLTP** | Online Transaction Processing
**ACID** | atomicity, consistency, isolation, durability
**RDF** | Resource Description Framework
**DPT** | data processing tools
**NGS** | Next-Generation Sequencing
**DNA** | Deoxyribonucleic acid
**MLST** | Multilocus Sequence Typing
**MLVA** | Multiple-Locus Variable Number Tandem Repeat Analysis
**SNP** | Single Nucleotide Polymorphism
**CSV** | comma-separated values
**ST** | Sequence Type
**eBURST** | Electronic Based Upon Related Sequence Types
**goeBURST** | Globally Optimized eBURST
**API** | application programming interface
**HTTP** | Hypertext Transfer Protocol
**AOP** | aspect-oriented programming
**APOC** | Awesome Procedures on Cypher
**SQL** | Structured Query Language
**IoC** | inversion of control
**JSON** | JavaScript Object Notation
OGM object graph mapper

REST representational state transfer
Chapter 1

Introduction

Phylogenetics is the study of the evolutionary history and relationships among individuals or groups of organisms, which aims to produce diagrammatic hypothesis about the history of the evolutionary relationships of a group of organisms known as phylogenetic tree. The relationships are inferred through the analysis of the traits of the individual or group, that is, by applying computational algorithms, methods, and programs to the phylogenetics data.

With the growing exchanges of people and merchandise between countries, epidemics have become an issue of increasing importance. The computational phylogenetics were mostly performed in personal computers and desktops. That is, the data was loaded to the client computer through a text file, the algorithms were executed, and the results of the algorithms executions were then displayed to the user. However, this kind of analysis is not feasible anymore, since huge amounts of data are being collected every day, and there are certain operations that require a considerable amount of memory or time. Instead, it is now common to run such tasks in high performance computing environments and/or dedicated systems. Therefore, there is a need to find a better way to store and maintain the data rather than in personal computers and desktops.

In large scale phylogenetic analysis of microbial population genetics, it is often needed to sequence, and type the information of the organisms, and afterwards to apply a set of phylogenetic inference methods to produce a diagrammatic hypothesis about the history of the evolutionary relationships of a group of organisms. The computation and analysis of microbial population genetics often produce graphs and trees, which have many relationships [52]. As graph databases naturally apply to these data structures and are optimized to perform queries and operations over them, that is, they are designed specifically to deal with highly connected data, it should be possible to store them in a graph database.

Although graph oriented databases can be of much help in this setting, as far as it is known there is no solution relying on these technologies to address large scale phylogenetic analysis challenges. Thus, a study on which database engine better addresses the needs of this challenge can provide new insights and lead to innovative approaches, comparing graph databases such as Neo4j [20], Titan Aurelius [2], JanusGraph [13], Dgraph [9], Allegrograph [22], and Apache Rya [41]. The comparisons made in this context suggest that Neo4j offers the most interesting set of features and capabilities, such as relying on an architecture based on native graph storage and processing engine, having an active community, and allowing to extend itself with plugins to support any graph algorithm. Therefore, Neo4j is the graph database system that is used to address the large scale phylogenetic analysis challenge.
1.1 Objectives

The objective of this project is to develop a modular framework for large scale phylogenetic analysis that exploits a graph oriented database technology which allows to access the phylogenetics data, without needing to load it into the clients computers. This framework should have a data model that allows the representation of large phylogenetic networks and trees, as well as the ancillary data. It should support queries on such data, and allow the deployment of algorithms for inferring/detecting patterns and for pre-computing visualizations.

1.2 Document Structure

This document is composed by several chapters, namely by the Background, Proposed Solution, Implementation, Experimental Evaluation, and Final Remarks chapters. The Background chapter starts by explaining the concept of a graph database and presents several use cases where a graph database is used to store highly connected data. Then, several graph databases are identified and discussed to understand which database suits better. Moreover, it presents a platform that performs phylogenetic analysis, explains how it is accomplished and presents the different types of phylogenetic data. The Proposed Solution chapter enumerates several requirements and use cases for this project. It proposes an architecture for a framework, to cope with the phylogenetic large scale analysis challenge, exploiting several architectural views. Additionally, it describes what technologies shall support this framework, providing several key characteristics of each. The Implementation chapter emphasizes the architectural choices by exemplifying how to extend some parts of the framework. Furthermore, it presents implementation decisions and explains why they were embraced. The Experimental Evaluation chapter explains the types of tests, the operations, the datasets, and the system settings used to evaluate the framework. Afterwards, it provides an analysis of the results obtained from these tests. Finally, the Final Remarks chapter presents the conclusions of this project, and showcases how it can be extended in future work.

This project is publicly available at https://github.com/Brunovski/phyloDB and it can be deployed using Docker [21]. The repository of the project provides the architecture documentation, the deployment and usage definitions, and examples of how to deploy and use the framework.
Chapter 2

Background

This chapter provides an overview of the graph databases, and the large scale phylogenetic analysis of microbial population genetics.

The data that is managed during the large scale phylogenetic analysis of microbial population genetics may have many relationships, and as graph databases are designed specifically to deal with highly connected data, it should be possible to store and manage this data within a graph database. Therefore, the concept of a graph database is introduced, and several use cases of using a graph database system to store and manage similar data are presented. Furthermore, the problems that may result from using a Relational Database Management System (RDBMS) to handle the same type of data are discussed. To support this discussion, an experiment that compares a RDBMS with a graph oriented database is analysed. Due to RDBMS not being suitable for large scale phylogenetic analysis, some metrics are defined to better understand which graph database engine is the most suitable. Hence, several graph databases engines are compared using those metrics.

The study of microbial population genetics is composed of several processes, such as the alignment of organisms sequences, the application of a typing methodology and the execution of a set of phylogenetic inference methods. Hence, a platform which performs this kind of analysis is presented. Afterwards, since the usual processes of the study of microbial population genetics constitutes scientific workflows, the definition and examples of scientific workflows are presented. Moreover, the workflow system which is used by the platform is introduced. Each one of the specified processes are detailed to understand how the study of microbial population genetics is accomplished. Finally, throughout the explanation of all these concepts and processes, the several types of data that exist or may appear during an execution of the overall process are also explained.

2.1 Graph Databases

A graph database management system is an online database management system with create, read, update, and delete (CRUD) methods that expose a graph data model. Graph databases are generally built for use with Online Transaction Processing (OLTP) systems. Accordingly, they are normally optimized for transactional performance, and engineered with transactional integrity and operational availability in mind[44, 45]. This type of database addresses the problem of leveraging complex and dynamic relationships in highly connected data. Graph databases offer appealing characteristics, such as performance and flexibility.
Regarding the performance of graph databases, it tends to remain relatively constant, even as the dataset grows, because the queries only use the respective portion of the graph. In terms of flexibility, a graph database allows adding new nodes, labels, and relationships, to an existing structure, without disturbing the existing queries and application functionality. A graph database exposes a graph data model, by storing the data in a graph format. A graph is a set of nodes that represent entities and a set of edges that represent the relationships that connect them. Graphs and trees are often produced in the computation and analyses of phylogenetic data. Therefore, it should be possible to use a graph oriented database, which naturally applies to these data structures and are optimized to perform queries and operations over them.

2.1.1 Use Cases

The most common graph database use cases include contexts such as social data, recommendation algorithms, geospatial operations, and network and data centre management. All of these use cases share a common context characteristic, that is they all deal with highly connected data, just like in the context of large scale phylogenetic analysis. In each use case, using a graph data model and the specific characteristics of the graph database, allows to generate competitive insight and significant business value. For instance, in social applications, such as Facebook, using a graph database allows to understand who interacts with whom, and how people are connected. This allows the social network application to generate insight of the aspects that influence individual behaviours.

2.1.2 RDBMS Comparison

The use of a RDBMS to store connected, semi-structured datasets is not suitable, as they struggle when attempting to model several depths of relationships. In spite of relationships existing in relational databases, they only exist as a mean of joining tables. In this context it is often needed to disambiguate the semantics of the relationships that connect entities, as well as qualify their weight or strength. Relational relations do nothing of this kind. Another drawback of using a RDBMS occurs when the amount of data increases significantly, and the structure of the dataset becomes more complex and less uniform. To cope with this, the relation model has to include large join tables, null-checking logic, and sparsely populated rows. Therefore, as the number of relations increases, it degrades the performance of the system and makes it difficult to evolve the existing database in response to changing business needs.

To exemplify that RDBMS are not adequate for this type of context, an experiment was conducted using a relational store and Neo4j. The objective of this experiment was to test the execution time of both database systems when querying for friends of friends in a social network context with different levels of depth of friends, to the maximum level of depth of five. Summarizing the experiment, the idea is that given any two persons chosen at random, find a path that connects them which is at most five relationships long.

The conclusion of the experiment is that for a social network containing about 1,000,000 people, each with approximately 50 friends, the results strongly suggest that graph databases are the best choice for connected data, as shown in Table 2.1. The most interesting fact of this experiment is that at depth four the relational database exhibits crippling latency, making it practically useless for an online system. Neo4j’s timings have deteriorated a little too, but the latency here is at the periphery of being acceptable for a responsive online system. Finally, at depth five, the relational database simply takes too long to complete the query, however Neo4j returns a result in around two seconds.
2.1.3 Graph Databases

The RDBMS systems might not be then the best choice to address the context of large scale phylogenetic analysis challenge. Although graph oriented databases can be of much help in these contexts, as far as it is known there is no solution relying on these technologies to address this challenge. Therefore, the set of graph database systems that were selected for comparison are Neo4j, Titan Aurelius [2], JanusGraph [13], Dgraph [9], Allegrograph [22], and Apache Rya [41].

Neo4j is one of the most popular graph database management systems, and it is used in several companies such as Ebay and Microsoft. It is implemented in Java and accessible from software written in other languages using the Cypher query language [15]. Titan Aurelius is a scalable graph database optimized for storing and querying graphs. It is built over backend storage frameworks such as Apache HBase [10] or Apache Cassandra [25]. JanusGraph, which debuted in 2017, is a database optimized for storing and querying large graphs with billions of edges and vertices distributed across a multi-machine cluster. It is based on the source Java code base of the Titan graph database project and is supported by the likes of Google, IBM and the Linux Foundation. Dgraph, which is written in Go, is a distributed graph database offering horizontal scaling and atomicity, consistency, isolation, durability (ACID) properties. It is built to reduce the disk seeks and minimize network usage footprint in cluster scenarios. It automatically moves data to rebalance cluster shards. AllegroGraph is a commercial graph database which supports many programming languages, and is based in uniform machine architectures. Apache Rya is a scalable system for storing and retrieving the Resource Description Framework (RDF) data in a cluster of nodes. It introduced a new serialization format for storing the RDF data, an indexing method to provide fast access to data, and query processing techniques for speeding up the evaluation of SPARQL queries. RDF is based on the idea of making statements about resources in the form of `<subject, predicate, object>` expressions called triples.

To better understand which graph databases are best suited for our context, several of their capabilities and features are compared in Table 2.2. These include the underlying storage and processing engine, writing scalability, storage capacity, semantics extensibility, built-in algorithms, visualization tools, data processing tools (DPT) integration, and source code and activeness.

| Database             | Storage PE | Scalability | Nodes   | Extensible Algorithms | Visualization | DPT | Source | Active |
|----------------------|------------|-------------|---------|-----------------------|---------------|-----|--------|--------|
| Neo4j                | N          | N           | C       | Yes                   | Yes           | Yes | Open   | Yes    |
| Titan Aurelius       | N          | N           | C       | ?                     | No            | No  | Yes    | No     |
| JanusGraph           | N          | N           | C       | 2^59                  | No            | 2   | Yes    | Open   |
| Dgraph               | N          | N           | C       | ?                     | No            | No  | Yes    | Open   |
| AllegroGraph         | N          | NN          | SS      | 1 Billion             | Yes           | 0   | No     | Closed |
| Apache Rya           | NN         | NN          | C       | ?                     | No            | 0   | No     | Open   |

**Keys:** PE - Processing Engine, N - Native, NN - Non Native, SS - Single Server, C - Cluster

Table 2.2: Databases comparison summary.
**Underlying Storage and Processing Engine** The underlying storage can be a native graph storage, that is optimized and designed for storing and managing graphs. Alternatively, it can serialize the graph data into a relational database, an object-oriented database, or some other general-purpose data store. The processing engine can either be native, if the database exposes a graph data model through CRUD operations and uses index-free adjacency, or non-native, if it uses global indexes. A database engine that relies on index-free adjacency is one where each node maintains direct references to its adjacent nodes, and each node acts as a micro-index to other nearby nodes, which is much cheaper than using global indexes. It means that query times are independent of the total size of the graph, and are instead proportional to the amount of the graph data searched. A non-native graph database engine, in contrast, uses global indexes to link nodes together. These indexes add a layer of indirection to each traversal, thereby incurring greater computational cost.

Neo4j and Dgraph are built over a native graph storage and native processing engine system. Titan Aurelius and JanusGraph are based on a native graph processing engine, but they are built over backend storage frameworks such as Apache HBase [10] or Apache Cassandra [25], therefore not providing a native graph storage. AllegroGraph has a native graph storage and a non-native graph processing engine. Finally, Apache Rya is not based either on a native graph storage or native graph processing engine.

**Writing Scalability** The writing scalability refers to the possibility of the database to scale, depending on the amount of write operations workload that it is subjected to. The appealing form of scalability is horizontal, as it allows to scale by adding more resources to support the database. AllegroGraph is based on a single server architecture, and all the remaining database engines support cluster architectures that allows them to scale horizontally.

**Storage Capacity** Storage capacity is the space that a database provides to store the data, which in this case is organized as nodes and relationships. AllegroGraph allows to store up to 1 billion nodes, while Neo4j supports the storage of 34 billion nodes [42]. Titan Aurelius and JanusGraph allows to store up to 1 quintillion relationships and half as many nodes. And finally, Dgraph and Apache Rya do not specify a storage capacity limit.

**Semantics Extensibility** Semantics extensibility is the possibility to extend the database functionalities by writing custom code, which can then be invoked. Neo4j and AllegroGraph provide mechanisms that allow the extension of the database semantics, by implementing custom functions or procedures, whereas the remaining database systems do not.

**Built-in Algorithms** Built-in algorithms are the algorithms which are provided by the database to run over the stored data. Neo4j has community-driven libraries available, that combined with the officially available algorithms, provide 31 implemented algorithms to use over the graphs stored in the system. Finally, JanusGraph provides 2 built-in algorithms, while the remaining database engines do not provide any.

**Visualization Tools** Graph visualization tools allow to visualize the data which are stored on the database. Neo4j, JanusGraph and Dgraph allow to integrate visualization tools to visualize the data which is stored in the database. For instance, Neo4j allows to visualize data with tools that connects directly to the database such as Neovis [7] and Popoto [23]. The remaining databases do not support the visualization of the data with visualization tools.
DPT Integration Data processing tools allow to collect and manipulate data into a desired form. Dgraph does not provide the possibility of integrating the database with a data processing tool. However, all the other databases provide such functionality. For example, Neo4j allows to integrate with Apache Spark.

Source Code and Activeness The source code of a project or tool can either be open or closed source, depending on whether the code is available for the public or not. Activeness stands for the status of a project, which can be active, if it is still in development, or concluded if not. Thereby, Neo4j, JanusGraph, Dgraph, and Apache Rya are open source and active, AllegroGraph is closed source and also active, and Titan Aurelius is open source and concluded.

Considering these graph database system comparisons, it can be recognized that Neo4j offers the most interesting set of capabilities and features, and also has an active community. The most interesting set of capabilities and features will be identified and detailed in Section Proposed Solution.

2.2 Large Scale Phylogenetic Analysis

In large scale phylogenetic analysis projects, such as INNUENDO, the performance of epidemiological surveillance is necessary, with the data generated also being used to study microbial population genetics. In the particular case of INNUENDO, it provides an effective indicator based surveillance system of priority-listed pathogens, which is fundamental in combating and controlling food-born diseases. This system is able to monitor the geographical location, spread, type and genomic variation of pathogens to rapidly detect the emergence of food-borne outbreaks. It is based on a framework that contains an analytical platform and standard procedures for the use of whole-genome sequencing in the surveillance, outbreak detection, and investigation of food-born pathogens in the context of small countries with limited resources.

The analysis of this type of data is based on the need to pass files through a series of transformations, called a pipeline or a scientific workflow. A scientific workflow is a framework which aims to compose and execute a series of computational or data manipulation steps over some data. An example of a workflow system is shown in Figure 1.

The early forms of workflow systems are based on scripts and the make utility. However, for phylogenetic analysis it is desired that these frameworks should be able to accommodate production pipelines consisting of both serial and parallel steps, complex dependencies, varied software and data file types, fixed and user-defined parameters and deliverables. Thereby, there are many workflow frameworks of different types, that were built specifically to solve some mentioned problems. The INNUENDO project relies on a workflow framework named FlowCraft. FlowCraft is a python engine that automatically builds pipelines by assembling previously made and ready to use components, which are modular pieces of software or scripts. Hence, FlowCraft is used to assemble a set of transformations to perform the study of microbial population genetics. The alignment of genetic sequences, the application of a typing methodology, and finally the execution of a set of phylogenetic inference methods, comprise the typical sequence of steps in microbial population genetics studies. The alignment task deals with small parts of the organism genome, specifically with locus and alleles. Subsequently, by applying a typing methodology several data types arise such as isolates, allelic profiles, and ancillary data. Finally, the last task applies inference and visualization algorithms, which generate distances between allelic profiles, and visualization coordinates for each allelic profile. Each one of these concepts are explained in the following subsections.
2.2.1 Phylogenetic Data

The input of Flowcraft is the organism information that comes from the laboratories. Hence, given such information that comes as biological samples, a Next-Generation Sequencing (NGS) process is applied to obtain the genetic sequences. Then, alignment tools or assembly tools are executed to assemble the genomes [4, 24]. This process can be defined as the reconstruction of the organism genome system with random small parts of it to determine the nucleic acid sequence, that is, the order of nucleotides in the Deoxyribonucleic acid (DNA). It allows to map genomes of new organisms, finish genomes of previously known organisms, or to compare genomes across multiple samples. It can be performed by two different processes, the matching and the assembly. In the matching process, the alignment is accomplished by building the current genome based on a previously built genome of the same organism. In the assembly process, all the needed combinations of the alleles are made to obtain the correct form of the genome. These processes can be compared as building a puzzle, where the matching has the solution for the puzzle available, while the assembly does not.

Locus and Alleles  The sequences assembled in the alignment process may occupy a given position of a locus and define distinct alleles of that locus. A locus is a specific location in the chromosome, and every unique sequence, either DNA or peptide depending on the locus, is defined as a new allele. An allele can also be defined as a viable DNA coding sequence for the transmission of traits, and it is represented with a number identifying the allele and string containing the sequence. An example of this data is presented in Figure 2.1 where the allele with identifier 1 belongs to locus aroE, and the sequence of the allele is the following string with the nucleic acid sequence, which is a succession of letters that indicate the order of nucleotides. It is expected that, the alleles are represented through files which follow the FASTA format. FASTA is a text-based format for representing either nucleotide sequences or amino acid sequences, in which nucleotides or amino acids are represented using single letter codes. This format is represented in Figure 2.1.

>aroE_1
GAAGCGAGTGACTTGCGACAAACAGTGCCCAATGTTGCTGCTACAGATGTTGTCATC
GCGCGCTTGGATTTGGCTGACACCAGTGGCAATTGATATTTAATTGATAT
CTAGACAAGTTACAGACGGACAGGTTGTATATTAGTGATATTGAT
>aroE_2
GAACCAGTGACTTGCGACAAACAGTGCCCAATGTTGCTGCTACAGATGTTGTCATC
GCGCGCTTGGATTTGGCTGACACCAGTGGCAATTGATATTTAATTGATAT
CTAGACAAGTTACAGACGGACAGGTTGTATATTAGTGATATTGAT

Figure 2.1: Example of alleles.

2.2.2 Microbial Typing

After the alignment phase, a typing methodology is applied to identify or fingerprint each organism based on the genes that are presented in almost all organisms, which are named as conserved genes. Bacterial identification and characterization at subspecies level is commonly known as microbial typing. This process provides the means to execute phylogenetic inference methods, which then produces a hypothesis about the history of the evolutionary relationships about a group of organisms. There are several typing methodologies, such as the MLST [48, 31], Multiple-Locus Variable Number Tandem Repeat Analysis (MLVA) [28], and SNP [8]. These are available for a multitude of bacterial species, and are being used globally in epidemiological microbial typing and bacterial population studies.
Isolate  The main goal of the typing methods is the characterization of organisms existing in a given sample. However, some microorganisms from the sample collected need to be isolated to be characterized. Thus, each organism isolated from the microbial population becomes an isolate.

Ancillary Data  An isolate can be associated with typing information and ancillary details. Ancillary details include information about the place where the microorganism was isolated, the environment, the host, and other possible contextual details. These details are usually represented through comma-separated values (CSV) formatted files, where the tab character is used to separate values. An example of this data is shown in Figure 2.2, which shows that the ancillary detail line with id 1 is related to the allelic profile with id 1, and is composed by some information that was extracted from the isolate with the identifier AU2523. This information include the species, that is *A. denitrificans*, and the isolation localization, which was in North America, more precisely in USA.

| id | ST | isolate  | species   | country | continent    |
|----|----|----------|-----------|---------|--------------|
| 1  | 1  | AU2523   | *A. denitrificans* | USA     | North America|
| 2  | 2  | AU8059   | *A. denitrificans* | Unknown |              |
| 3  | 3  | AU8060   | *A. denitrificans* | France  | Europe       |
| 4  | 4  | AU8080   | *A. insolitus*    | USA     | North America|
| 5  | 5  | ACH26    | *A. insolitus*    | USA     | North America|

Figure 2.2: Example of ancillary details.

Allelic Profile  One of the most popular methodologies is the **MLST** which is an unambiguous procedure for characterizing isolates of bacterial species using the sequences of internal fragments. This methodology types several species of microorganisms, and when applied, the set of alleles identified at the loci are considered to define a Sequence Type (ST), a key identifier for this methodology, that can also be defined as an allelic profile. The chosen loci are usually different for each species, although some species may share some or even all loci in their MLST schemas. The number of chosen loci can vary and be greater or smaller than the seven loci more commonly adopted. The generated sequences are compared to an allele database and for each gene, the different sequences are assigned as distinct alleles and, for each isolate, the alleles at each of the loci define the allelic profile, also known as ST in this methodology. These allelic profiles are represented through files which follow a tabular format. Usually, it is a delimited text file that uses a tab character to separate values. The **MLVA** methodology follows a similar concept, in the sense that by applying the methodology it is obtained sequences which are identified by some identifier, and it has several columns representing each element that characterizes the sequence. The **SNP** methodology differs from the latter, since the data which is originated by it follows a sequential format, where each sequence is normally constituted by an identifier and a set of numbers. A value of 0 in any locus represents the allele that was mostly found on that locus, while a 1 represents any other possible allele.

Examples of several allelic profiles, created by applying the **MLST** and **SNP** methodologies, are presented in Figure 2.3 and in Figure 2.4 respectively. In Figure 2.3 it can be interpreted that the profile with ST identifier 1 is identified by the loci *nuSA*, *rpoB*, *eno*, *gltB*, *lepA*, *nuoL*, and *aroe*. The number below each locus is the allele identifier, which ends up mapping to the different alleles, that were described before. For example the locus *aroe*, with the allele identifier with value 1 maps for the first entry of the Figure 2.1. A ST having the locus *aroe* with the allele identifier with value 2 maps to the second entry. **MLVA** allelic profiles would have a similar representation.
|   |   |   |   |   |   |   |   |   |
|---|---|---|---|---|---|---|---|---|
| ST | nusA | rpoB | eno | gltB | lepA | muoL | aroe |
| 1  | 1   | 26  | 2   | 2   | 59  | 8    | 1    |
| 2  | 1   | 26  | 2   | 4   | 59  | 2    | 1    |
| 3  | 1   | 26  | 2   | 2   | 62  | 8    | 2    |
| 4  | 1   | 26  | 7   | 2   | 59  | 3    | 2    |
| 5  | 1   | 27  | 1   | 1   | 62  | 9    | 1    |

Figure 2.3: Example of MLST profiles.

In Figure 2.4 it can be interpreted that the profile with identifier 1 is characterized by the following sequence, which contains a set of 0’s and 1’s that represent if this profile is characterized by the alleles that were the mostly found on that locus.

1 0100000111101000101010101010101100111010010101010101010101010100
2 1111010010101100100101010101001000001010010001010101010100

Figure 2.4: Example of SNP profiles.

### 2.2.3 Algorithms

Succeeding the typing process follows the execution of a phylogenetic inference method to the results. A phylogenetic inference method is the application of computational algorithms, methods, and programs to phylogenetic data that allows to produce a diagrammatic hypothesis about the history of the evolutionary relationships of a group of organisms. There are several types of phylogenetic inference methods, such as distance matrix methods, maximum parsimony, maximum likelihood, and Bayesian inference. However, in INNUENDO the inference methods which are applied are based on distance matrix methods [17, 33, 55].

**Distances** Distance matrix methods rely on the genetic distance between the sequences being classified. The distances are often defined as the fraction of mismatches at aligned positions, with gaps either ignored or counted as mismatches. An example of a distance matrix, resultant from applying the hamming distance between each profile represented in Figure 2.3, is shown in Figure 2.5.

$$
D = \begin{bmatrix}
0 & 2 & 2 & 3 & 5 \\
2 & 0 & 4 & 4 & 5 \\
2 & 4 & 0 & 3 & 5 \\
3 & 4 & 3 & 0 & 6 \\
5 & 5 & 5 & 6 & 0 \\
\end{bmatrix}
$$

Figure 2.5: Example of a distance matrix.

**Inference Algorithms** An inference algorithm is then executed to compute the diagrammatic hypothesis based on the distance matrix previously calculated. The diagrammatic hypothesis calculated comes in the form of a graph or a tree, where each node represents the allelic profile, and the relationships between them are quantified by the distances. Figure 2.6 is an example of such evolutionary hypothesis. The semantics of these relationships depends on the algorithm used. The Globally Optimized eBURST (goeBURST) algorithm [16] is an example of an implementation of these algorithms. It is a globally optimized implementation of the electronic Based Upon Related Sequence Types (eBURST) algorithm [11] that identifies alternative patterns.
of descent for several bacterial species. This algorithm can be stated as finding the maximum weight forest or the minimum spanning tree, depending on weight definition.

![Graph](image)

Figure 2.6: Example of an evolutionary hypothesis.

The `goeBURST` algorithm uses the Kruskal algorithm, described in Algorithm 1, to achieve its goals. It relies on this algorithm since it can obtain the optimal tree, with respect to the defined partial order on the set of relationships between organisms. However, in case of a tie when comparing the weight of the relations, it should consider the number of relations at distance one, two and three, the occurrence frequency of the relation, and lastly the sequence type identifier.

When executing many inference algorithms over the same allelic profiles, the concept of multilayer networks, which consists of using the same nodes to represent different graphs, becomes appealing. That is, due to the several algorithm outputs being expressed over the same context, it becomes easier to compare them. For example, if the `goeBURST` algorithm and any other algorithm are run, then the nodes of both resulting graphs would be the same, but for each algorithm it would produce different relationships.

**Algorithm 1**  
Kruskal algorithm.

**Input:** A graph $G$ whose edges have distinct weights.  
**Output:** $F$ is the minimum spanning forest of $G$.

**Initialization:** Create a graph $F$ (a set of trees), where each vertex in the graph is a separate tree. Create a set $S$ containing all the edges in the graph.

**Loop:** While $S$ is not empty and $F$ is not yet spanning do:

1. Remove an edge with minimum weight from $S$.
2. If the removed edge connects two different trees then add it to the forest $F$, combining two trees into a single tree.

**Finalize:** Return the hierarchy $F$.

**Visualization Algorithms**  
After executing an inference algorithm, a visualization algorithm, such as Radial Static Layout [3], which is described in Algorithm 2 or GrapeTree [57], is executed to compute the optimal coordinates for each node of the received graph or tree. Afterwards, the coordinates are provided to a render framework which then presents each profile and relationship to a user interface. The Radial Static Layout algorithm is a method of displaying tree structures in a way that expands outwards, radially. It is one of many ways to visually display a tree and can be used when the size of the node is not considered.
Algorithm 2 Radial Static Layout algorithm.

Input: \( T = (V, E, \delta) \).
Output: Coordinates \( x, y : V \rightarrow R^+ \) for the nodes.
Data: Queue \( Q \), leafcount: \( V \rightarrow N^+ \{ \text{from a previous postorder traversal} \} \).

Initialization: \( r \leftarrow \text{root}(T) \); rightborder\( (r) \leftarrow 0 \); wedgesize\( (r) \leftarrow 2\pi \); \( x(r) \leftarrow y(r) \leftarrow 0 \).

Loop: While \( !Q.\text{empty}() \) do:

1. \( v \leftarrow Q.\text{delete.first}() \).
2. \( \eta \leftarrow \text{rightborder}(v) \).
3. Loop: For Each child \( w \) of \( v \) do:
   (a) \( Q.\text{insert}(w) \)
   (b) \( \text{rightborder}(w) \leftarrow \eta \).
   (c) \( \text{wedgesize}(w) \leftarrow 2\pi \times \text{leafcount}(w) \div \text{leafcount}(r) \)
   (d) \( \alpha \leftarrow \text{rightborder}(w) + \text{wedgesize}(w) \div 2 \)
   (e) \( \alpha \leftarrow \text{rightborder}(w) + \text{wedgesize}(w) \div 2 \)
   (f) \( x(w) \leftarrow x(v) + \cos(\alpha) \times \delta((v, w)) ; y(w) \leftarrow y(v) + \sin(\alpha) \times \delta((v, w)) \)
   (g) \( \eta \leftarrow \eta + \text{wedgesize}(w) \)

2.3 Discussion

This chapter provided an overview of the graph databases, the phylogenetic analysis process and the types of data related to it. It started by presenting the definition of a graph database, and several use cases where a graph database can be useful. Additionally, it discusses the problems which may arise of utilizing a RDBMS to manage data that can be represented by graphs, by analysing an experiment. To understand which graph database could be suitable, several graph database engines were compared. Then, the large scale phylogenetic analysis process is presented emphasizing the phylogenetic data and algorithms that are used by it. There are several types of data such as, the alleles, the sequence allelic profiles, the isolates, the ancillary data, the relations between each profile, and the optimal coordinates for the visualization rendering. Some of these data can be represented by dataset files of different formats, such as MLST, MLVA or SNP data. Since graph databases have been applied in contexts where there is highly connected data, it should be possible to use a graph database to store this types of data.
Chapter 3

Proposed Solution

This chapter defines the functional and non functional requirements, the uses cases, the architecture, and the technologies that shall compose the proposed solution.

The proposed solution relies on a framework that allows to store and manage the data resulting from the phylogenetic analysis in a graph database, execute inference and visualization algorithms over such data, and more. There are several uses cases for a framework like this, for instance the possibility to query data and to load datasets. To provide functionalities like those, the framework is composed by different components. These components are described with several architectural views, such as a data model, client server, layered and decomposition. These views are complemented with the respective reasoning. At last, these components must rely on some technology to be executed. Hence, the technologies that shall support the different components of the framework are specified.

3.1 Functional Requirements

Functional requirements state what the system must do, and how it must behave or react to. The functional requirements that were identified for the framework of large scale phylogenetic analysis are as follows:

- Store in a database the data resulting from the phylogenetic analysis considering multilayer networks;
- Development of a framework which allows to perform queries over the data stored in the database;
- Loading of several datasets with different formats, such as CSV and FASTA, into the database;
- Execution of inference and visualization algorithms, like goeBURST and Radial Static Layout respectively, over the data stored in the database;
- Support of authentication and authorization.

3.2 Non Functional Requirements

Non-functional requirements or quality attributes requirements are qualifications of the functional requirements or of the overall product. The non-functional requirements that were identified for the framework of large scale phylogenetic analysis are as follows:
The framework should be complemented with documentation that provides different types of views.

- The framework code should be modular, in a way that it allows to easily reuse and extend the code.

### 3.3 Use Cases

A use case is a written description of how users will perform tasks on a system. It outlines, from the user perspective, the behaviour of the system as it responds to a request. The use cases identified shall allow to reason about who are the users of the framework, their objectives, the actions they are able to perform, and how the framework shall respond to each action. The framework should allow its users, which are other applications, to query data, load datasets, execute algorithms, and obtain results. Each of these operations start by verifying if the request is authenticated, the permissions of the user, and the request. Each use case is represented in Figure 3.1.

![Figure 3.1: Use cases.](image)

**Query Data**  The framework should allow its users to query any type of data that is stored on the database. For example, it should be possible to find all isolates for which it already exists typing information under a specific typing method schema. Thus, the user should perform the several requests, that identifies the resources to obtain. Once the request is received by the framework it should return the data gathered from the database. The several types of data may include profiles, isolates, and others.

**Load Dataset**  The framework should allow its users to load several datasets into the database. For example, it should be possible to load a file containing the profiles of a taxonomic unit. Thus, the user should perform a request sending the dataset to be loaded. Once the request is received by the framework, it should parse the data into database entities, store them, and return a code identifying if the operation was successful. The datasets may come from several file formats, such as FASTA, CSV, and others.

**Execute Algorithm**  The framework should allow its users to run an algorithm, either for inference or visualization, over the data that is stored in the database. For example, it should be possible to run the goeBURST algorithm to calculate the minimum spanning tree, given a set of profiles, and to run the Radial
Static Layout algorithm to calculate the respective visualization coordinates. Thus, the user should perform a request identifying which algorithm it wants to run. Once the request is received by the framework, it should schedule the execution of the algorithm and return a code identifying if the operation was scheduled.

**Obtain Algorithm Result** The framework should allow its users to obtain the resulting graph from an algorithm execution, after it has been completed. Thus, the user should perform a request identifying which algorithm execution result to retrieve. Once the request is received by the framework, it should return the result stored in the database.

### 3.4 Architecture

The software architecture of a system is the set of structures needed to reason about the system, which comprise software elements, relations among them, and properties of both. There are several views to document an architecture. Their purpose is to provide insight of how the system is structured as a set of implementation units, in contrast to others that aim to understand how the system is structured as a set of elements that have run-time behaviour and interactions [5]. Thus, the architecture of the proposed framework should be represented by several views. However, only the most relevant and generic shall be presented in this document, noting that all the others are available in the project repository.

#### 3.4.1 Data Model

The data model style [6] describes the information structure in terms of data entities and their relationships. It is used to perform impact analysis of changes, to enforce data quality by avoiding redundancy and inconsistency, and to guide the implementation of modules that access the data. It should be designed as a graph, since the domain of large scale phylogenetic analysis is composed of highly connected data. Therefore, the data model is described as a connected graph of nodes and relationships, where the nodes represent domain entities, and the relationships represent how the different nodes relate. By using a graph data model extensibility should be achieved by itself. That is, it should be possible to extend the system just by adding a set of nodes and relationships. However, it must be ensured that the domain is not violated, since the graph data model does not impose a strict schema. The data model of this project is shown in Figure 3.2.

This data model incorporates versioning and soft deletes concerns to mitigate some problems that occur nowadays, such as the impossibility to delete a wrongly inserted profile after executing an analysis that generates a graph containing it. Such profiles can not be removed, because the generated graphs would then become invalid. In this case, by considering a versioning and soft delete strategy, these removals should be possible, since the graphs would be linked to the statuses of the profiles and not to the profiles themselves.

The versioning strategy to achieve this behaviour is to separate each object from its state, link them through a relationship with the respective version number, and capture changes by having different state nodes [26]. In the data model, the name of the status nodes shall end with *Details*, and the version relationships shall be named as **CONTAINS_DETAILS**. For example, the allele node is connected to the respective version by the **CONTAINS_DETAILS** relationship between the *Allele* and the *Allele Details* nodes. If the allele is modified, then a new *Allele Details* node is connected to the *Allele* node, while all profiles that were related to that allele version are still valid.

Phylogenetic data are composed of taxonomic units, *loci*, and alleles. Taxonomic units consist of several *loci*, thus this is represented by a relationship named **CONTAINS** between taxonomic units and *loci* nodes. For
instance, to represent the taxonomic unit *Streptococcus pneumoniae* which contains several *loci*, such as *aroE* and others, the *Streptococcus pneumoniae* node would have a CONTAINS relationship to *aroE* node. *Loci* may hold specific locations for a set of alleles, as shown in Figure 2.1. Therefore, to represent this relationship in the data model, a relationship named CONTAINS is used between the nodes of each locus and the associated nodes of the alleles. For instance, to express the Figure 2.1 in the data model, the locus *aroE* would have relationships CONTAINS to two allele nodes. In Figure 3.3 a simplified example of this case is presented.

The typing methods execution relies on and generates different types of data [1, 54], such as typing schemas, allelic profiles, isolates and ancillary data, as explained before.

Typing schemas can use several *loci* to characterize different allelic profiles, and in the data model this is expressed by the relationships HAS between the details of a schema node and the respective *loci* nodes. For instance, to describe the MLST schema used in Figure 2.3, the details of the schema node should contain relationships HAS to the *loci* *nusA*, *rpoB*, *eno*, *gltB*, *lepA*, *nuoL*, and *aroE*. Figure 3.4 presents an example dataset that follows this MLST schema. However, only the relationships to the *aroE* locus are shown to keep the figure simple, but in the complete scenario it would have a relationship to each of the mentioned locus.
Allelic profiles belong to a specific dataset as they are a result of applying a typing methodology. Thus, this is described by the relationship \texttt{CONTAINS} between dataset and profile nodes. These profiles follow the same schema, hence they should be related to the typing method used. To impose this concern, the \texttt{HAS} relationship is used between the dataset and schema nodes, which means that all profiles from that dataset follow the related schema. For example, the details node of a dataset containing the profiles shown in Figure 2.3 would have a relationship named \texttt{HAS} to a \texttt{MLST} schema node. However, having the dataset connected to the schema, only allows to perceive what \textit{loci} were used in the typing operation.

Therefore, to know what is the allele that characterizes a profile for each locus used in the schema, the details node of a profile must be connected to the respective allele nodes. Hence, this is represented in the data model by using a relationship called \texttt{HAS} between the profile details and the alleles nodes. For instance, in Figure 2.3 the profile with \texttt{ST} value 1 is represented by the allele with identifier 1 of locus \texttt{nusA}, consequently that profile should have a detail with a relationship called \texttt{HAS} to that allele. This example is also presented in Figure 3.4.

Isolates may have related ancillary data, as it is represented in Figure 2.2. Thus, in the data model this is expressed as a relationship named \texttt{HAS} between the detail of an isolate and ancillary data nodes. Since an isolate may be associated to a profile, there is also a relation between the two, which is called \texttt{HAS}. For example, to represent the associations between profiles, isolates, and ancillary data, shown in Figure 2.2 the detail of the isolate with name U2523 would have a relationship called \texttt{HAS} to the profile with id 1. The detail of this isolate would also have several relationships \texttt{HAS} to each ancillary data associated to it. Figure 3.4 also represents a simplified variation of this example.

Graphs and trees are generated by performing analysis over the datasets generated by the typing process.
These analyses are based on the execution of inference and visualization algorithms.

The inference algorithms rely on the genetic distances between profiles. These distances are calculated by computing a distance matrix, as shown in Figure 2.5. Based on these distances, the algorithm is then executed and relationships DISTANCES are created between the different profiles composing the graph. For example, to describe the result of an inference algorithm shown in Figure 2.6, the several profiles that were used in the execution of the algorithm would have relationships DISTANCES between them, quantified by the respective weights. This example is also presented in Figure 3.5 in a simplified way. This strategy allows to consider multilayer networks since the same nodes shall be used to represent different graphs.

Visualization algorithms execute over the graphs resulting from the inference algorithms, and create visualization coordinates for each node of the graphs. Hence, a relationship HAS between a profile and coordinate nodes exists to represent the coordinate of some profile, for a given inference and visualization algorithm. Figure 3.5 also represents an example of the coordinates for the nodes of a given graph.
Users and projects are concepts introduced to handle the authorization concern. The datasets of the profiles may belong only to some users, hence this access restriction should be expressed in the data model. Thus, a user may participate in several projects, and a project may contain a dataset of profiles and several algorithm executions. By restricting the access to a project, the ancillary details, and the results of the algorithms are also restricted. The data model embodies these concerns by using the relationships HAS between project and user nodes, and CONTAINS between project and dataset nodes.
3.4.2 Client Server

The client-server style is used for analysing the modifiability, reusability, scalability and availability of the solution, by factoring out common services, to better understand how the several units of computation interact between themselves. The client is a component that invokes services of a server component, and the server is a component that provides services to the client components. The client-server view for this framework is presented in Figure 3.6.

![Client-Server Architectural View](image)

**Figure 3.6: Client-Server architectural view.**

The Phylodb Server component provides a web application programming interface (API), to perform several operations over the data stored in the database, namely access data, load datasets, execute algorithms, and obtain results. It should be possible to scale horizontally to handle more operations by adding more instances of the Phylodb Server component. The communication between the users and the Phylodb Server component should be done over the Hypertext Transfer Protocol (HTTP) protocol.

The Phylodb Server interacts with the database component in two different ways. That is, it can normally query data, but it can also queue executions of the algorithms that are deployed in the database and reside in the Algorithms Plugin component. These algorithms also read the needed inputs and write the computed results. The communication between the database and the Phylodb Server component shall be done over the Bolt protocol [34].

The Authorization Server component manages the user information, and provides operations to perform the authentication of a user. It needs to receive the authentication requests over the HTTP protocol.

3.4.3 API

The API structure should be based on three layers, namely Controllers, Services and Repositories, as represented in Figure 3.7. This figure represents a layered view that puts together layers, which are
groupings of modules that offer a cohesive set of services, in a unidirectional allowed-to-use relation with each other.

![Layered architectural view](image-url)

**Figure 3.7: Layered architectural view.**

When a request is received by the API, it is passed through the **Controllers** layer. This layer contains the controllers that parse the received input, execute the respective service, and retrieve the response containing the respective status code and the formatted content. The **Services** layer contains the services that perform the business logic and use the needed repositories. The **Repositories** layer holds the repositories that shall provide operations to interact with the database. Apart from these layers, there is a validation logic that verifies the request authenticity and the user permissions before the request is processed by them.

Another approach for decomposing the structure of the API is to separate it by modules. This approach can be visualized by relying on a decomposition view. This view is used for decomposing a system into units of implementation, which allows to describe the organization of the code as modules, to better understand how the system responsibilities are partitioned across them, and to reason about the location of changes. The modules that shall compose the structure of the API are the **phylogeny**, **typing**, **analysis**, **security**, **io**, **error**, and **utils**. These modules are represented in Figure 3.8.

The **phylogeny**, **typing**, and **analysis** modules contain the main operations to handle the different types of data. The **phylogeny** module should combine the logic of managing taxonomic units, loci and alleles. The **typing** module shall contain the operations to administer datasets, profiles, isolates, and schemas. The **analysis** intends to contain the operations to manage the graphs and coordinates, resulting from the execution of inference and visualization algorithms. Each of these modules follows the layered structure previously explained, that is, each module contains the respective controller, service and repository.

The **security**, **io**, **error**, and **utils** modules represent the remaining concerns, which are important to the overall functioning of the API. The **security** module aims to hold the logic of managing users and projects, and contains the authentication and authorization concerns. The **io** module shall aggregate the
input parsing and output formatting logic. The error module should provide operations to perform error handling. Finally, the purpose of the utils module shall be to provide common operations which do not belong to a specific module and can be used by several modules.

Since the structure is decomposed in several modules, it is also possible to understand how they interact with each other by analysing an uses view. This view shows how modules depend on each other and which modules can be affected by changes in other modules. This view is represented in Figure 3.9.

It shows that the modules phylogeny, typing, and analysis should rely on the security module, which allows them to define the roles and permissions of the users that can perform their respective operations. They shall also depend on the io module to parse and format their respective inputs and outputs. And they intend to use the common operations provided by the utils module. The security module shall depend on io and utils modules for the same reasons. Finally, the io module and utils modules shall not rely on other modules. It can be concluded that if changes are made in the phylogeny, typing, or analysis modules, they do not influence any other module. However, if a change is conducted in any of the other modules then the phylogeny, typing, analysis, and security modules might need to be reviewed.

### 3.4.4 Plugin

The structure of the plugin can also be based on three layers, namely Procedures, Services and Repositories as demonstrated in the layered view represented by Figure 3.10. This view implies that the modules within the Procedures layer are allowed to use the modules belonging to the Services layer, and so on, but the opposite is not allowed. The Procedures layer intends to hold the definitions of the operations that allow to execute the supported algorithms, hence a call to an algorithm is directed to them. Their implementation shall parse the received input and execute the respective service provided by the Services layer. The Services layer shall read the input data for the algorithm from the database, compute the respective algorithm and store the obtained result back to the database. The reading and writing of data is accomplished by using the methods provided by the Repositories layer.

Such structure is constituted by several modules and it can be further decomposed into the inference,
Figure 3.9: Uses architectural view.

Figure 3.10: Layered architectural view.
visualization, and utils modules. These are shown in the decomposition view presented in Figure 3.11. The inference module should contain the inference algorithms and hold the operations to retrieve and store data, related to an inference, from the database. The visualization module is similar, in the way that it aims to support the visualization algorithms and hold the operations to retrieve and store data, related to a visualization, from the database. Finally, the utils module shall provide common operations which do not belong to a specific module and can be used by different modules. Each module, except for the utils module, follow the layered structure previously explained, that is, each module contains the respective procedure, service and repository.

![Diagram showing the decomposition view](image)

Figure 3.11: Decomposition architectural view.

The interaction between these modules can be understood by analysing the uses view represented by Figure 3.9. In this view, it can be recognized that the inference and visualization modules rely only on themselves and on the common operations provided by the utils module. Hence, a change can be conducted without the need to review the other module, since they do not interact with each other. Additionally, the utils module shall not rely on any other module.

### 3.5 Technology

The technologies that shall support the framework are the Spring Framework [14], Google Identity Provider [18], and Neo4j. These technologies will be further explained ahead, and it will be specified what component of the framework they shall support.

#### 3.5.1 Spring Framework

The Spring Framework is an application framework and inversion of control container for the Java platform. The appealing features of Spring are that it allows to develop applications of different types and provides appealing aspect-oriented programming (AOP) features to deal with cross-cutting concerns [39]. Furthermore, a library which connects the Spring system to the Neo4j database already exists. This library benefit from the AOP framework of Spring to make the access to the database simpler. For instance, instead
of managing the transactions manually, spring offers the annotation `@Transactional`, which specifies that the method annotated with it must be run in a transactional environment. Finally, the extensions for the query language of Neo4j and most of the already implemented phylogenetic inference algorithms are implemented in the Java language. Considering these facts, the Spring Framework shall support the application server component of the framework.

### 3.5.2 Google Identity Provider

An identity provider is a system that vouches for the identity of a user. The identity provider role is to authenticate a user and provide an authentication token to the service provider. The OAuth 2.0 [API](http://example.com), from Google, which conform to the OpenID Connect specification [38], will be used to authenticate the users. Hence, the authorization server of the framework will rely on Google Identity Provider, which shall authenticate the client applications by any of the existing flows [19]. Once the authentication process is completed, an identity token should be granted to the client application. It must then be sent within the request to the application server, so it can authenticate the request.

An example of the authentication process is shown in Figure 3.13, where there are three main components, namely the client application, the application server and the authorization server. It is assumed that the client application already obtained the identity token, thus step 1 starts by sending a request to the application server to perform a given operation. Once the application server receives it, the step 2 begins by sending a request to the authorization server to validate the token. Afterwards, in step 3 the authorization server validates the token and sends a response to the application server. Finally, in step 4, the application proceeds with the operation, and sends a response to the client.
3.5.3 Neo4j

Considering the graph database system comparisons made previously, it can be recognized that Neo4j offers the most interesting set of capabilities and features, and also has an active community. The most interesting features being the fact that it uses Cypher as its query language and is able to extend the query language semantics by implementing custom procedures. As well as the fact that it allows scheduling query executions over the database, and provides a library of graph algorithms, utility functions, and trigger mechanisms. It also allows to perform data sharding.

**Cypher** An effort was started to standardize Cypher as the query language for graph processing. It allows to store and retrieve data from the graph database, and the syntax of Cypher provides a visual and logical way to match patterns of nodes and relationships in the graph. It is a declarative, Structured Query Language (SQL)-inspired language for describing visual patterns in graphs and allows to state what to select, insert, update, or delete from the graphs without a description of exactly how to do it. Through Cypher, it is possible to construct expressive and efficient queries to handle the needed create, read, update, and delete functionalities.

**Apoc** The library of graph algorithms, which Neo4j yields [37], is the Awesome Procedures on Cypher (APOC). It provides algorithms with different focus such as, community detection, centrality, path finding, similarity, and link prediction. These algorithms are presented respectively in Table 1, 2, 3, 4, and 5 in the appendices. Besides algorithms, the library contains a collection of functions and procedures, that cover collective operations such as sorting, graph operations, text searches, conversions, geospatial operations, data integration, and reporting.

**Triggers** The APOC library provides some background operations, such as trigger mechanisms. These trigger mechanisms allow to register Cypher statements that are executed when data in database are changed. For instance, it is possible to run a trigger before or after events such as when data are created, updated, or deleted. The trigger mechanism is an appealing feature to achieve a kind of incremental computation when executing the algorithms.

![Authentication process using an identity provider.](image)

Figure 3.13: Authentication process using an identity provider.
Scheduler  The APOC library also provides other type of background operations, which are the background jobs. They allow procedures to run in the background or asynchronously, and execute procedures periodically. The background jobs mechanism is based in a Java scheduler, which relies on a queue, to post the tasks to execute, and on a thread pool, where the threads will execute the tasks that are on the queue. This feature is appealing since some algorithms may take several minutes to finish executing.

Custom Procedures  A user defined procedure or custom procedure, is a mechanism that enables to extend the Neo4j semantics by writing custom code, which can be invoked directly from Cypher. Procedures can take arguments, perform operations on the database, and return results. These procedures are implemented through Java plugin projects, that after compiled, must be placed in the Neo4j database instance, where it is needed to run the procedure.

Sharding  Data sharding allows to divide data into horizontal partitions that are organized into various instances or servers, most often to spread load across multiple access points. These shards can be accessed individually or aggregated to see all of the data, when required. This is particularly interesting when the quantity of data is becoming large enough that it makes sense to divide the data into smaller graphs to run on smaller sized hardware and be accessed by necessary parties.

3.6  Discussion

This chapter defines the proposed solution, which outlines the development of a framework that conforms to the phylogenetic analysis. This framework shall allow to manage the data resulting from the phylogenetic analysis in a graph database, execute inference and visualization algorithms, retrieve their results, and load datasets of different formats. Hence, the framework shall be composed of a web API, a graph database with a plugin to support the algorithms and an authorization server. There are different types of architectural views to describe such components, namely data model, client server, layered, decomposition and uses. Each of these architectural views is complemented with the respective explanation.

The components of the framework shall rely on the Spring Framework, the Google Identity Provider, and Neo4j technologies. The Spring Framework shall support the web API since it has some appealing features, such as providing aspect oriented programming features to deal with cross-cutting concerns. The Google Identity Provider shall handle the authentication of the framework clients. It allows to perform authentication conforming to the OpenID Connect specification. Finally, Neo4j will support the database, as it offers the most interesting set of capabilities and features, and also has an active community. Some of these features include the custom procedures, the APOC library and the scheduler mechanism.
Chapter 4

Implementation

This chapter describes and reasons about the implementation details of the framework presented in Chapter 3. It starts by describing what are the components of the solution. Then, it explains the details of the API component, such as the interface, the parsing and formatting of files, the error handling, and the security concerns. Finally, it describes details of the plugin component, such as the algorithms. Besides these details, it explains how the respective architecture for each component was materialized, and presents several exemplifications of the different concerns that are described throughout this chapter.

The solution was implemented considering an agile methodology and is publicly available at https://github.com/Brunovsk/phyloDB along with its issues, milestones, and documentation. For testing purposes, the solution is hosted in a server as a Docker image.

4.1 Structure

This implementation relies on three components, namely the API, the plugin, and the database. The API receives requests and processes them into queries that are executed over the database. The plugin holds the inference and visualization algorithms and allows the API to execute them over the data stored in the database. The database contains the phylogenetic data, which is stored as a set of nodes and edges. A database schema is not implied by Neo4j, hence the database schema is defined by the queries performed through the operations provided by the API.

4.2 API

The API provided by the framework was implemented considering the representational state transfer (REST) architecture, which is based on statelessness. That is, the server should not store any state about the client session on the server-side. Each request from the client to the server must contain all the information necessary to interpret the request. Therefore, the session state must be kept entirely on the client. This architecture has many advantages, such as the possibility to scale the API to several instances by deploying it to multiple servers, since any server can handle any request, as they do not possess any session related dependencies.

This API relies on the Spring Framework, which allows to use the inversion of control (IoC) features that it provides. This principle has several advantages, such as allowing to have a greater modularity and
facilitating the change between different implementations. The IoC pattern used in the API is the dependency injection, which supports setting the dependencies of the objects. That is, the act of injecting objects into other objects is done by an assembler rather than by the objects themselves.

the Component annotation and its implementations were used to achieve the dependency injection. This annotation is used to denote classes as components, which then Spring uses to auto detect them for dependency injection. Besides the Component annotation Spring offers more specific annotations, such as the Controller annotation, that is used to define controllers, the Service annotation, which allows to depict classes that hold business logic, and the Repository annotation, that is utilized to describe data access classes. These annotations are useful in the implementation of the framework, since they follow the same paradigm as the proposed architecture, which is specifically presented by Figure 3.7.

4.2.1 Interface

The interaction between the API and users is accomplished through HTTP requests. Thus, the HTTP protocol allows the API to retrieve different types of responses depending on several factors, such as the parameters used in the request, the type and result of the operation that is being executed.

The API relies on models to parse the input which comes within the body of the requests. These models are defined by the InputModel interface. It defines the toDomainEntity method, which intends to parse the JavaScript Object Notation (JSON) object contained in the body of the request. That is, the input models are used to parse the input data from the requests into domain entities. For instance, the ProjectInputModel, which extends from InputModel and implements the respective method to retrieve a Project, would be used to parse the data contained in the --data-raw section of the following request. This request is an example of a curl request that is used to create a project.

```
curl --location --request POST 'http://localhost:8080/projects?provider=google'
   --header 'Content-Type: application/json'
   --header 'Authorization: Bearer {Access Token}'
   --data-raw '{
       "name": "Example Project",
       "visibility": "private",
       "description": "Example project",
       "users": [{"email": "example1@gmail.com", "provider": "google"}, ...]
   }
```

Likewise, the API also relies on a different type of models to format the domain entities into output objects. These models are defined by the OutputModel interface. It outlines the toResponseEntity method, that aims to format a domain entity into an output object, which is then retrieved in a response to the user. For example, the ProjectOutputModel extends from OutputModel and implements the respective method to retrieve a response containing the 200 OK status code and an object with some of the properties of the respective project. Thus, the database schema is not exposed since these output models contain only the data that is meant to be retrieved.

The status code of the responses, which contains these output models, are defined conforming to the HTTP protocol. That is, the status codes depend on the type and result of the respective operations. For example, the 200 OK status code is used for operations that shall retrieve representations of domain entities, the 201 CREATED status code is used for operations that create a domain entity and retrieve the generated
identifier, and the 204 NO CONTENT status code is used for operations which do not retrieve anything. While the 400 BAD REQUEST status code is used to indicate that the request is not valid due to some input, and the 401 UNAUTHORIZED is utilized for indicating that the request is not valid due to lack of authentication or permissions. Another example is the 500 INTERNAL SERVER ERROR which is used to signal that an internal error occurred while processing a request.

The mime type for the responses containing these output models is always the application/json, since these output models only allow to format objects or lists of objects. However, this API also supports the parsing and formatting of more complex data types, namely files, that can have several formats. Thus, the file parsing and formatting will be further detailed ahead. However, the mime type used to retrieve a list of domain entities formatted in a file format is the text/plain.

4.2.2 Formatters

Imports and exports of large quantities of data are based on files. There are different formats of files depending on the type of data that they hold. Thus, the interface Formatter defines methods to support the imports and exports operations, namely the parse and format methods. The parse method iterates over the lines of a file and calls another method to translate them into domain entities. The latter must be implemented by the inheriting classes. The format method iterates over a list of domain entities to transform them into a String representation formatted in a specific file format.

There are many file formats used in the phylogenetic analysis. These formats are used to represent alleles, profiles, isolates, and inferences. The alleles can be formatted in the FASTA format. The profiles can be represented in the MLST, MLVA or SNP formats. The MLST and MLVA formats are very similar, that is, the only difference is the number of columns used to represent the profiles. The isolates follow a tabular format. And the inferences can be represented in Newick or Nexus.

A class that extends from the Formatter interface exists for each of these formats. Each of these classes implements the inherited methods according to the format that they represent. For instance, the AlleleFormatter, which extends from Formatter, implements the method used by the parse method to allow the parsing of alleles from a FASTA file, and the format to represent alleles in a FASTA formatted String. This structure of classes is presented in Figure 4.1.

These formatters can rely on certain settings, that are defined in the application.properties configuration file, namely the missing symbols to be considered in some formats and the maximum length of the lines to be formatted in the FASTA format. These settings are stored in the configuration file, so they can be modified without having to recompile the project.

This structure of formatter classes can be easily extended. For instance, to support a new file format it is only needed to create a class that extends from the Formatter interface and implement the method used by the parse method, and implement the format method, according to the format that it shall represent.

4.2.3 Errors

Errors can occur when processing requests. The cause of these errors can either be a user mistake or an internal processing issue. When errors occur, an error response is retrieved to the user with the respective status code. These responses are built based on error models. The ErrorOutputModel receives the type of error that occurred and produces the respective error response. The type of errors is defined by the enum Problem. For example, an ErrorOutputModel is built using the Problem.UNAUTHORIZED problem to retrieve
an error response which signals that the request did not have a valid authentication token.

User errors can happen due to lack of authentication or permissions, and wrong inputs. However, internal errors may only occur when an unexpected exception is thrown. When internal errors and some of the user errors occur, they are handled by a ControllerAdvice component. This component allows to declare exception handling methods using the ExceptionHandler annotation. These methods are globally applied to all controllers and allow to receive exceptions as arguments. That is, this component acts as a try catch which encloses all the controllers, where each method catches the exceptions corresponding with the ones declared in their arguments. For example, a method that receives an HttpMediaTypeNotAcceptableException exception as argument is executed when an exception of the same type occurs.

Besides retrieving error responses when internal errors occur, log entries are also created. These logs are useful to understand the internal errors that occur. The org.slf4j.Logger logger is used to perform these logs. It is configured by defining the respective settings in the configuration file. For instance, this file should contain a property that holds the name of the file to where the logs should be written to.

### 4.2.4 Security

The security concerns, namely authenticity and permissions should be validated before a request is processed by the controllers. Thus, a security pipeline which validates these concerns was built based on the Spring interceptor components \[49\]. These components allow to implement the preHandle method, that is executed before a request is passed to the controllers.

The security pipeline is composed by the AuthenticationInterceptor and AuthorizationInterceptor
interceptors. These components implement the `SecurityInterceptor` interface. This interface defines the method `handle`, which shall contain the authenticity and user permissions validations. However, to ensure that these validations are executed before a request is passed to the controllers, the `SecurityInterceptor` must implement the Spring `HandlerInterceptor` interface and define that the `handle` method is executed within the `preHandle` method. This structure is represented in Figure 4.2.

Figure 4.2: Security generalization architectural view.

The `AuthenticationInterceptor` is the first component of the security pipeline to be executed. It supports the bearer token authentication [32]. This type of authentication is based on tokens that are acquired after an authentication process with an identity provider. Thus, the requests must include the authorization header with the value `Bearer` followed by a space and the authentication token. Currently, the authentication relies only on the Google Identity provider, hence the `GoogleInterceptor` component extends from the `AuthenticationInterceptor`. It implements the `instrospect` method to define how to validate the token with Google. In case of the token being invalid, an error response with 401 `UNAUTHORIZED` status code is retrieved. Otherwise, the request is allowed to continue to the next security component.

The `AuthorizationInterceptor` is the following component of the security pipeline. It relies on roles and annotations. Each user has a role, and each endpoint of the controller is annotated with the minimum role, which the user must hold, and with the type of operation that it represents. Hence, when this component processes a request it gathers that information and verifies if the request can proceed to the controllers. In case the needed conditions are not met, an error response with 401 `UNAUTHORIZED` status code is retrieved. Otherwise, the request is allowed to proceed to the controllers.

The security pipeline can be easily extended to support more authentication providers. To do so, it is only needed to create a new component that extends from `AuthenticationInterceptor` and implement the `instrospect` method with the specific settings to validate the token.
4.2.5 Controllers

The controllers yield the endpoints of the API. Thus, several instances of controllers compose the Controllers layer. Each of these controllers are annotated with the Controller annotation and defines different methods, however they all extend from the Controller base class which contains utility methods that can be used by any of the extending controllers. For instance, it holds the getAllFile method that allows to retrieve a list of domain entities from a given service and map it to a specific file format. This structure is represented in Figure 4.3.

![Controller generalization architectural view.](image)

These controllers rely on input and output models to receive and retrieve data as JSON objects, and rely on formatters to import and export large quantities of data in different file formats. Moreover, they depend on services to perform the logic associated to each endpoint. Hence, the controllers rely on the dependency injection mechanism to inject them. In our case, they are injected through the constructor of the controllers.

A few configurations that are defined in the configuration file are used by the controllers, namely the maximum limits of the lists of domain entities that are retrieved, depending on the type of data. That is, there are different limits for the JSON and the file formatted String responses.

4.2.6 Services

The services provide methods to perform the business logic. Thus, several instances of them compose the Services layer. They yield operations to manage persistent or transient domain entities. The majority of the domain entities exists independently of the time, however jobs only exist while the respective algorithm execution is not processed. Therefore, the JobService defines operations to manage jobs, while the EntityService define operations to manage persistent entities. Although they provide operations for different types of entities, they are annotated with the Service annotation and extend from the Service base class.
The `EntityService` can be further decomposed depending on the need to consider versioning. That is, services which shall manage domain entities that need versioning must extend from `VersionedEntityService`, while services which do not need versioning must extend from `UnversionedEntityService`. This difference translates in the yielding of different methods. For example, the `VersionedEntityService` provides the `find` method that allows to retrieve the specified entity based on a `Key` and a `long`, which are the identifier and the version of the entity. However, the `UnversionedEntityService` provides a similar `find` method that has the same functionality but does not consider the version. That is, it retrieves the specified entity based only on the `Key` that is received as argument.

The `VersionedEntityService` can be further extended to support batch operations. Thus, `BatchService` extends from `VersionedEntityService` to provide different operations that process multiple entities. For example, it yields the `saveAll` method that intends to save a list of entities that is received as arguments. This structure is represented in Figure 4.4.

In the implementation of each of these services, it was considered that the methods must be executed inside a transaction to maintain the consistency of the data stored in the database. Hence, the `Transactional` annotation was used to achieve such behaviour. This annotation is used to combine more than one write operation on a database as a single atomic operation. When a method annotated with it is called, all or none of the writes on the database are executed.

These services normally rely on repositories that perform the interactions with the database. Hence, the
services utilize a set of repositories which are also injected by the Spring mechanism of dependency injection.

The service implementation classes can either extend from BatchService, VersionedEntityService, UnversionedEntityService or Service depending on their need. For example, the ProfileService must implement the operations inherited from BatchService, VersionedEntityService, and EntityService, because it considers a persistent domain entity that needs versioning and batch operations. This structure can be easily extended to support more services. To do so, it is only needed to create the new service and extend it from one of the mentioned services depending on the needs.

4.2.7 Repositories

The repositories provide methods to perform the data access. Hence, several instances of them compose the Repositories layer. The structure of repositories is very similar to the services one, in the sense that it also yields operations to manage persistent or transient domain entities. Therefore, the JobRepository defines operations to manage jobs, while the EntityRepository define operations to manage persistent entities. Although they provide operations for different types of entities, they are annotated with the Repository annotation and extend from the Repository base class which allows to perform queries to the database. Likewise, the EntityRepository can also be further decomposed into VersionedEntityRepository and UnversionedEntityRepository, depending on the need to consider versioning.

Moreover, the VersionedEntityRepository can be further extended to support batch operations. Thus, BatchRepository extends from VersionedEntityRepository to define methods that process lists of entities. For example, it yields the `saveAll` method that intends to save the list of entities that is received as arguments. This structure is represented in Figure 4.5.

Several details were considered in the implementation of the methods provided by each repository, such as the use of pagination, object graph mapper (OGM) and parameterized queries.

An implementation consideration was to not use the usual functionalities of an OGM. The use of these functionalities may add some overhead [56]. Hence, it was decided that each query should be implemented from scratch to increase the performance of the data access operations. That is, the implementation of each repository method contains the respective Cypher query that it should perform, for example the `findAllEntities` method implementation contains the query that allows to retrieve the resumed information of a subset of entities.

Moreover, the use of pagination in these queries was adopted because this framework is intended to handle great quantities of data. This approach allows to control the quantity of data that is dealt with in the methods that retrieve many domain entities. Thus, memory issues are less likely to happen since a maximum number of records that can be retrieved by the queries is defined. To achieve this behaviour, the respective methods must receive the page and the limit of records to retrieve. For instance, the `findAllEntities` method provided by the VersionedEntityRepository receives the page and limit values as arguments, which allows to retrieve the respective set of domain entities according to them.

Also, each of those queries is parameterized. By using parameterized queries the performance can be increased because Neo4j can cache the query plans and reuse them in the following executions, which increases the following query speed. And, it also allows to protect from injection attacks, since parameters are never allowed to be interpreted as part of the query and have no means of escaping out of being anything other than a value of some sort [56]. This can be achieved by executing a query that contains specific placeholders for each parameter and pass the values of the parameters, in the correct order, as arguments.

Another implementation characteristic is the use of indexes [35]. They can increase the performance of
some of the read queries. A database index is a redundant copy of some of the data in the database for the purpose of doing searches of related data more efficiently. They are useful when there are large quantities of data stored in the database, because it will allow the queries to rely on the index that points to a specific node instead of searching in the respective group of nodes for it. These indexes are created through a script that must be executed before the API is started.

Finally, the UNWIND feature of Neo4j was used to implement the batch operations. This feature is optimal to deal with up to fifty thousand records per operation, and allows to process a list of rows into individual rows that contain the information for each of the updates. Furthermore, queries using this feature can also be parameterized, and are constant, which allows Neo4j to use them for caching. Neo4j struggles to handle queries that contains many lines, therefore by using this feature small enough queries are used.

4.3 Plugin

The plugin of the database is based on the feature of user-defined procedures from Neo4j. A user-defined procedure is a mechanism that allows to extend Neo4j by writing custom code, which can be invoked directly from Cypher. These procedures can take arguments, perform operations on the database, and return results. Moreover, some resources can be injected into them from the database, which is similar to the dependency injection mechanism from Spring.
This plugin intends to extend Neo4j to support inference and visualization algorithms, which shall be available as procedures. The inference algorithms procedures are executed over the profiles of a dataset, while visualization algorithms procedures are executed over the results of the inference algorithms. These procedures can be invoked directly from Cypher like any other standard procedure. For example, the execution of the goeBURST algorithm over a dataset with identifier `dataset_id`, belonging to a project with identifier `project_id`, only considering a maximum distance of three, and storing the result with the identifier `inference_id`, can be represented as:

```cypher
CALL algorithms.inference.goeburst(project_id, dataset_id, 3, inference_id)
```

However, these procedures are meant to be executed only by the API since certain validations are performed by it before executing them. For example, the existence of an inference is verified before a visualization algorithm execution is scheduled by the API.

This plugin is based on the three layers previously presented in Figure 3.10. Each of these layers was materialised into a structure of classes, that allows to easily extend themselves. These structures are presented in Figure 4.6.

![Figure 4.6: Plugin generalization architectural views.](image)

### 4.3.1 Algorithms

Algorithms receive inputs, perform the respective computation, and produce results. Hence, inference algorithms receive distance matrices and produce graphs, while visualization algorithms receive the graphs produced by inference algorithms and generate coordinates for each of the nodes of the graph.

Thus, the interface `Algorithm` defines a method `compute` that is extended by the `InferenceAlgorithm` and `VisualizationAlgorithm` interfaces to specify the respective arguments and results. For example, `InferenceAlgorithm` defines that the `compute` method must receive a `Matrix` and retrieve an `Inference`.

The goeBURST algorithm is the only inference algorithm supported, and the Radial Static Layout algorithm is the only visualization algorithm supported. Therefore, a class that implements the `compute` method exists for each of these algorithms. For instance, the class `GoeBurst` extends from `InferenceAlgorithm`
to implement the `compute` method, which receives a `Matrix` and retrieves an `Inference`, with the logic of the `goeBURST` algorithm. This structure of classes is represented in Figure 4.7. Implementations of the `goeBURST` and Radial Static Layout algorithms already exist, thus our implementations are based on them [47, 30].

![Figure 4.7: Algorithms generalization architectural view.](image)

This structure of algorithms classes can be easily extended to support new inference or visualization algorithms. It is only needed to create a class that extends from the respective base class to implement the `compute` method. For instance, to support a new inference algorithm, it is needed to create a class that extends from `InferenceAlgorithm` and implements the `compute` method with the respective algorithm logic.

### 4.3.2 Procedures

The procedures yield the interface of the plugin. Thus, different instances of procedures compose the `Procedures` layer. The methods declared within these procedure classes, that are annotated with the `Procedure` annotation, are allowed to be executed as a standard procedure. This annotation allows to define the designation that shall be used by the users to execute these procedures. For instance, the `InferenceProcedure` defines a method named `goeburst`, however the annotation holds the designation `algorithms.inference.goeburst`. Hence, the users need to use that designation to execute the `goeburst` method.

Each of these procedures defines different methods, however they all extend from the `Procedure` base class. However, the base class only holds the `GraphDatabaseService` object, which is injected by the database. This object allows to interact with the database and can only be injected at the procedures level, thus, it must be passed as an argument to the services, and then to the repositories, since the procedures rely on services to perform the respective logic. This structure is represented in Figure 4.6. These procedures are
similar to the controllers presented before, in the sense that they intend to yield the endpoints of the plugin.

4.3.3 Services

The services provide methods to perform the business logic. Thus, different instances of them compose the Services layer. They yield methods to execute algorithms. That is, the methods defined by each service are responsible for gathering the algorithms input data from the database, computing the algorithms, and storing the results back to the database. For instance, InferenceService provides the goeburst method that obtains the distance matrix for the specified dataset, executes the goeBURST algorithm, and then stores the resulting graph in the database.

Each of these services defines different methods, however they all extend from the Service base class, as shown in Figure 4.6. The base class holds the GraphDatabaseService object that is passed by the Procedure. Since each of those methods relies on the repositories to interact with the database, this object must also be passed to them.

These services rely on repositories to perform the interactions with the database. For example, the goeburst method from the InferenceService uses the InferenceRepository to obtain the distance matrix that will be used by the algorithm.

4.3.4 Repositories

The Repositories layer is composed by different repositories that provide methods to interact with the database. The objectives of these repositories are to retrieve the input data for the algorithms from the database, and to write their results back to the database. To accomplish these objectives, they rely on the GraphDatabaseService object, which allows them to perform operations over the database.

Also, based on those objectives, the Repository interface is provided, which defines the read and write methods. These methods are meant to be implemented to retrieve and store a certain type of data respectively. Thus, the InferenceRepository and VisualizationRepository implement those methods to read and write the respective types of data. For example, the InferenceRepository implements the read operation to retrieve a Matrix and implements the write operation to store an Inference. This structure is also presented in Figure 4.6.

The implementation of these methods is based on the Java Core API of Neo4j. There were other alternatives to implement the data access, such as using Cypher and using the Traversal framework. However, Cypher takes more time to execute and the Traversal framework became deprecated.

The structure of repositories classes can be easily extended to support new repositories. It is only needed to create a new class which extends from the base class, specify which type of data shall be retrieved by the read operation and received by the write operation, and implement those methods based on the Java Core API.

4.4 Discussion

This chapter describes and reasons about the implementation details of the API and plugin components that compose the framework. The API section includes implementation details, such as the compliance to the REST architecture, the use of dependency injection, and how the authentication and authorization processes are accomplished. The plugin section describes details such as, how it relied on the user-defined procedures...
mechanism of Neo4j, and how the algorithms were implemented. Besides these details, each section explains how the respective architecture for each component was materialized and presents several exemplifications of the different concerns that were described.
Chapter 5

Experimental Evaluation

This chapter explains the types of tests, the operations, the datasets, and the system settings used to evaluate the framework. Afterwards, it provides an analysis of the results obtained from these tests.

There are three types of tests that were used to analyse the framework, namely functional, performance, and load testing. The functional tests aim to validate the functionality of the framework, that is, verify if each operation provided by the framework works as expected. Thus, unit tests were designed for almost every functionality, achieving a code coverage of more than 90%. The performance tests objective is to check how some operations behave in terms of time and memory usage, under a particular workload. This type of tests allows to uncover what can still be improved in future work. The load tests purpose is to test how the framework behaves when handling several clients at a given moment. The framework might perform well for one client during performance testing, but it might degrade when many clients try to use it simultaneously during load testing due to lack of system resources. This type of tests also allows to uncover what can still be improved in future work.

The performance and load tests were executed over a set of read and write operations, which were picked to be analysed in terms of time and memory. These operations are composed by the Save Profiles, Run Inference, Run Visualization, Save Profile, Get Profiles, Get Resumed Profiles, Get Profile, Get Inference, and, Get Visualization. In these tests, the Save Profiles saves a file containing an increasing number of profiles. The Run Inference executes the goeBURST algorithm over an increasing number of profiles that are stored in the database. The Run Visualization executes the Radial Static Layout algorithm over the result of the goeBURST algorithm execution with an increasing number of profiles and edges. It must be taken into account that these two operations not only compute the algorithm, but also include the work of gathering the data and storing the results. Note that, in the case of the inference algorithms, the calculation of the distance matrix is also included in this process. Then, the Save Profile saves a single profile in the database that holds an increasing number of profiles. The Get Profiles and Get Resumed Profiles retrieve the respective information about an increasing number of profiles. The Get Profile retrieves only a single profile independently of the quantity of profiles stored in the database. Finally, the Get Inference and Get Visualization retrieves the results of the algorithms executed over an increasing number of profiles and edges.

The tests relied on the *Streptococcus pneumoniae* [40 MLST] dataset, which was specifically chosen because it is part of several published studies and also because it is publicly available, which will facilitate the interpretation of the results. This dataset contains a profile length of 7 and a total of around 16000 profiles.
This experimental evaluation was performed on a machine with an Intel Core i7 2.40 GHz quad core processor and 8 GB of memory, where 2 GB were allocated for the database and another 4 GB were allocated for the API.

### 5.1 Performance Testing

This section analyses the average running time of the operations, displaying the results in a table and a plot, as well as the average allocated memory by these operations, displayed in another table and plot. These tests were developed relying on a microbenchmarking tool for Java, namely Java Microbenchmark Harness, and using 5 warmups and 10 iterations for each operation being tested.

The average running time that each operation took to complete, over an increasing number of profiles, is represented in Table 5.1 in milliseconds. From these values, it can be observed that the **Save Profiles** write operation, takes around 6000 milliseconds to process 500 profiles, while the **Get Profiles** read operation takes much less time (68 milliseconds). The presented results confirm that the graph database operations that operate over a fixed amount of data are not affected by the increasing volume of data. This is confirmed by analysing the results for the **Get Profile** operation, which shows that the time of execution is constant and independent of the increasing number of profiles. Additionally, it can be concluded that the presented execution times for the algorithms comply with their time complexity, which is quadratic for the goeBURST algorithm [16], and linearithmic for the Radial Static Layout algorithm [3] since the children nodes are sorted.

The presented results also reveal that relying in a graph database to handle this type of data allows to have a good performance in read and single write operations.

| Operation          | Profiles | 500       | 1000      | 2000      | 5000      | 10000     | 15000     |
|--------------------|----------|-----------|-----------|-----------|-----------|-----------|-----------|
| Save Profiles      |          | 6294.861  | 12916.707 | 25434.663 | 64377.122 | 133941.541| 212235.800|
| Run Inference      |          | 316.292   | 1043.079  | 4199.889  | 25543.573 | 104885.648| 244421.421|
| Run Visualization  |          | 193.064   | 560.269   | 2055.067  | 12784.802 | 51034.175 | 119885.683|
| Save Profile       |          | 54.371    | 58.351    | 57.265    | 50.065    | 61.265    | 55.035    |
| Get Profiles       |          | 68.595    | 89.940    | 200.015   | 550.311   | 1549.135  | 2355.374  |
| Get Resumed Profiles|        | 6.392     | 10.320    | 19.018    | 47.900    | 96.402    | 150.546   |
| Get Profile        |          | 1.915     | 1.520     | 2.108     | 1.582     | 1.866     | 1.364     |
| Get Inference      |          | 5.954     | 10.777    | 20.097    | 48.638    | 100.737   | 152.563   |
| Get Visualization  |          | 7.625     | 13.712    | 27.242    | 81.630    | 138.806   | 214.127   |

Table 5.1: Performance testing time results in milliseconds.

This conclusion is also backed by the plot represented in Figure 5.1 which allows us to see that when the number of profiles used by a read operation is increased, it has little effect in the time taken to execute the operation. However, the time used by a batch writes operation increases almost or even linearly proportional to the number of profiles, when the latter is increased. From this plot, it can be seen that the line that corresponds to the **Save Profiles** write operation grows linearly proportional to the number of profiles as it grows, while the line which corresponds to the **Get Profiles** operation is almost a flat line, and the line which corresponds to the **Get Profile** operation is flat. Additionally, the lines corresponds to the algorithms are more curve than the others, that also supports that the algorithms follow their expected time complexity.
The average memory allocated that each operation consumed until completion, over an increasing number of profiles, is represented in Table 5.2 in megabytes. From these values, it can observed that the **Save Profiles** write operation allocates around 40 megabytes to process 500 profiles, while the **Get Profiles** read operation allocates much less memory space (9 megabytes). The presented results reveal that the framework allocates the memory linearly proportional to the amount of data that it is handling, and that the read and single write operations allocate much less memory than batch writes operations. However, it must be noticed that this analysis only considers the amount of memory allocated in the API. This is relevant because the algorithms are executed within the database, which causes them to use the database memory instead of the API memory. Hence, the allocated memory for the algorithms executions are minimal.

![Figure 5.1: Performance testing time plot.](image-url)

| Operation         | Profiles | 500     | 1000    | 2000    | 5000    | 10000   | 15000   |
|-------------------|----------|---------|---------|---------|---------|---------|---------|
| Save Profiles     | 40.283   | 80.519  | 160.746 | 402.086 | 803.059 | 1203.376|
| Run Inference     | 0.018    | 0.019   | 0.020   | 0.028   | 0.053   | 0.088   |
| Run Visualization | 0.018    | 0.019   | 0.020   | 0.023   | 0.032   | 0.047   |
| Save Profile      | 0.147    | 0.147   | 0.147   | 0.147   | 0.147   | 0.147   |
| Get Profiles      | 9.107    | 18.170  | 36.308  | 90.620  | 181.275 | 271.995 |
| Get Resumed Profiles | 1.940   | 3.867   | 7.732   | 19.243  | 38.571  | 57.781  |
| Get Profile       | 0.045    | 0.0426  | 0.0426  | 0.0426  | 0.0426  | 0.0426  |
| Get Inference     | 0.820    | 1.713   | 3.548   | 9.042   | 18.195  | 27.291  |
| Get Visualization | 0.569    | 1.148   | 2.340   | 5.889   | 11.826  | 17.719  |

Table 5.2: Performance testing memory results in megabytes.
This conclusion is also backed by the plot represented in Figure 5.2, which allows us to see that when the number of profiles used by any operation is increased, the amount of memory allocated grows almost linearly proportional to the number of profiles. For instance, the line that corresponds to Save Profiles write operation grows linearly proportional to the number of profiles as it increases.

![Figure 5.2: Performance testing memory plot.](image)

5.2 Load Testing

This section analyses the average running times of the operations that are presented in a table and a plot. These tests are executed over a database that contains around 500 profiles, and the Save Profile operation also involves 500 profiles. These tests were developed relying on a load testing tool, namely Apache JMeter, and using 10 iterations for each operation being tested.

The average time, in milliseconds, that each operation took to complete, over an increasing number of clients, is represented in Table 5.3. From these values it can be observed that the Save Profiles write operation, takes around 12000 milliseconds to process 10 different requests at the same moment, while the Get Profiles read operation takes much less time (101 milliseconds). The presented results also reveal that if the number of clients in a given moment increases, then it is still possible to maintain a good performance for the read operations. Lastly, these results also show that if there is nearly 250 clients performing the Save Profiles operation in a given moment, then the framework cannot handle all of those requests. However, this is not very problematic since the most common use cases for this framework for saving data rely on the use of the Save Profile operation, which performs efficiently.

This conclusion is also backed by the plot represented in Figure 5.3, which allows us to see that when the number of clients performing read operations is increased, it has a small effect in the time taken to
Table 5.3: Load testing, simulating several parallel users, time results in milliseconds.

| Operation                  | 1   | 10  | 50  | 100 | 250 | 500 | 1000 |
|----------------------------|-----|-----|-----|-----|-----|-----|------|
| Save Profiles              | 3005| 12348| 62511| 112772| -   | -   | -    |
| Save Profile               | 65  | 176 | 1305| 3118| 9164| 18259| 39030|
| Get resumed Profiles       | 6   | 6   | 6   | 5   | 7   | 320 | 1055 |
| Get Profiles               | 97  | 101 | 975 | 1469| 4019| 8154 | 17095|
| Get Profile                | 2   | 2   | 2   | 2   | 2   | 2   | 319  |
| Get Inference              | 33  | 21  | 26  | 102 | 493 | 1338 | 3267 |
| Get Visualization          | 30  | 16  | 14  | 15  | 312 | 915  | 1701 |

execute. However, the time line of a batch writes operation grows almost vertically and grows almost linearly proportional, in the single write operations, to the number of clients when it is increased.

Figure 5.3: Load testing time plot, simulating several parallel users.

5.3 Discussion

In this chapter it was discussed the implications of our solution in terms of time and memory requirements in two different types of analyses, namely performance and load testing. The most important results found are related with the read operations computational cost, when comparing to write operations, since the first represents a notable difference over the latter. Overall, we consider our implementation efficient in terms of read and single write operations, and recognize that the response time of the batch writes operations can still
be improved. However, the most common use cases of the framework will not be affected, since the saving of a single profile or a relatively small quantity of profiles are more typical than saving large quantities of profiles. The latter may happen in the early uses of the framework, which shall be in an offline environment and by a specific user.

Nonetheless, a possible improvement can be accomplished by using an alternative architecture for these operations, for example these operations could be handled with the same strategy applied for the algorithms, that is, the use of a queue mechanism. Using a queue mechanism would give us the advantages of providing a faster response time, as the writes are processed in the background, and of not having the constraint of only being able to process, in parallel, less than 250 clients. This alternative is proposed and explained in Chapter 6.
Chapter 6

Final Remarks

Epidemics have become an issue of increasing importance due to the growing exchanges of people and merchandise between countries. Hence, phylogenetic analyses are continuously generating huge volumes of typing and ancillary data. And there is no doubt about the importance of such data, and phylogenetic studies, for the surveillance of infectious diseases and the understanding of pathogen population genetics and evolution. The traditional way of performing phylogenetic analysis is not feasible anymore as a result of the amount of data generated. This analysis often produces graphs and trees, which have many relationships. Although graph oriented databases can be of much help in this setting, as far as it is known there is no solution relying on these technologies to address large scale phylogenetic analysis challenges.

6.1 Conclusions

This document starts by analysing what is the current problem with phylogenetic analysis when huge amounts of data are considered. Then, although graph oriented databases can be useful in this setting, we conclude that there is no solution based on these technologies.

Afterwards, it provides an explanation of what are graph databases, their use cases, their most important capabilities, and comparisons between them and a relational database. Then, we conclude that Neo4j offers the most interesting set of capabilities and features, beyond the general capabilities of graph databases, and also has an active community. This set of interesting capabilities and features are composed by the APOC library, the trigger and scheduler mechanism, and the user-defined procedures.

The following topic which the document addresses is the explanation of the phylogenetic analysis process. This process starts with a scientific workflow that takes biological samples as input to obtain the genetic sequences in the form of (short) reads through a NGS process. Then, alignment tools and/or assemblers are executed to align/assembly the genomes, which are further processed for identifying alleles at specific loci of interest. These genomes are then typed with a given typing methodology based on their loci, leading to several allelic profiles, and which are complemented by ancillary data. These allelic profiles can then be used as input for inferring evolutionary patterns, in the form of trees or graphs, for instance based on distance matrix methods. These patterns provide evolutionary hypotheses for the organisms being analysed and are usually presented to the analyst through different visualizations.

Then, the document provides an overview, based on the phylogenetic analysis process, of the main use cases and requirements for phylogenetic analysis. And it proposes a framework which complies with such
use cases and considers all of the requirements. This framework is composed of a Spring Framework API, a Neo4j database and a plugin for the latter, which relies on the custom procedures capabilities of Neo4j. It allows to authenticate with Google, manage all types of data that were analysed throughout the document, import and export dataset files of different formats, and execute inference and visualization algorithms. The framework shall contribute with some advantages for the phylogenetic analysis such as, by storing the results of algorithms will avoid having to compute them again, and using multilayer networks will make the comparison between them more efficient and scalable. Then, the architecture of this framework is presented with the use of different architectural views, and implementation characteristics are presented and described.

Finally, this document presents and explains how the experimental evaluation of the implemented framework was conducted, and also describes the respective results. From these results, we can conclude that the most important facts found are related with the read operations computational cost, when comparing to write operations, since the former represents a notable difference over the latter. We can also conclude that by using a graph database the operations executed over a fixed amount of data are not affected by an increasing volume of data. Furthermore, we observed that the execution times for the algorithms comply with their time complexity, which is quadratic for the goeBURST algorithm, and linearithmic for the Radial Static Layout algorithm. Overall, we consider our implementation efficient in terms of read and single write operations. However, with the presented results and their analysis, we understand that the framework can still be improved.

6.2 Future Work

There are several possible continuations of the work done throughout this thesis. One could be the extension of our solution to provide more algorithms, and make use of parallelization to improve their performance. The already provided algorithms by Neo4j make use of parallelization, hence we could improve our algorithms execution time by parallelizing their computations. Other potential development could be on how to use the background triggers functionality to achieve the dynamic computation of inference algorithms. Another possibility could be a study of how to perform the batch writes processing based on a queue mechanism. Or even to study how to use the Neo4j functionality of integrating with big data processing engines, such as Apache Spark. With this changing the batch writes operations of the API would have a faster response time. Finally, the API interface could be improved to be based on hypermedia. This would allow the API to be explorable and decouple the client applications from it.
References

[1] João Almeida, João Tiple, Mário Ramirez, José Melo-Cristino, Cátia Vaz, Alexandre P. Francisco, and João A. Carriço. An ontology and a rest api for sequence based microbial typing data. In Ana T. Freitas and Arcadi Navarro, editors, Bioinformatics for Personalized Medicine, pages 21–28, Berlin, Heidelberg, 2012. Springer Berlin Heidelberg.

[2] Aurelius. Titan: Distributed graph database., 2015. Last accessed 28 December 2020. URL: http://titandb.io

[3] Christian Bachmaier, Ulrik Brandes, and Falk Schreiber. Biological networks. 2014.

[4] J A Carriço, A J Sabat, A W Friedrich, M Ramirez, and Collective on behalf of the ESCMID Study Group for Epidemiological Markers (ESGEM). Bioinformatics in bacterial molecular epidemiology and public health: databases, tools and the next-generation sequencing revolution. Eurosurveillance, 18(4), 2013. URL: https://www.eurosurveillance.org/content/10.2807/esr.18.04.20382-en doi:https://doi.org/10.2807/esr.18.04.20382-en

[5] Paul Clements, David Garlan, Len Bass, Judith Stafford, Robert Nord, James Ivers, and Reed Little. Documenting software architectures: views and beyond. Pearson Education, 2002.

[6] Paul Clements, David Garlan, Len Bass, Judith Stafford, Robert Nord, James Ivers, and Reed Little. A tour of some module style. In Documenting software architectures: views and beyond [5], pages 109–120.

[7] Neo4j Contrib. Neovis.js. Last accessed 28 December 2020. URL: https://github.com/neo4j-contrib/neovis.js

[8] Nicholas J Croucher, Simon R Harris, Christophe Fraser, Michael A Quail, John Burton, Mark van der Linden, Lesley McGee, Anne von Gottberg, Jae Hoon Song, Kwan Soo Ko, et al. Rapid pneumococcal evolution in response to clinical interventions. science, 331(6016):430–434, 2011.

[9] Dgraph. Dgraph: A distributed, fast graph database., 2016. Last accessed 28 December 2020. URL: https://dgraph.io/

[10] Nick Dimiduk and Amandeep Khurana. HBase in Actions. O’Reilly Media, Inc., 2012.

[11] Edward J Feil, Bao C Li, David M Aanensen, William P Hanage, and Brian G Spratt. eburst: inferring patterns of evolutionary descent among clusters of related bacterial genotypes from multilocus sequence typing data. Journal of bacteriology, 186(5):1518–1530, 2004.

[12] FlowCraft. Flowcraft, 2016. Last accessed 28 December 2020. URL: https://github.com/assemblerflow/flowcraft
[13] The Linux Foundation. Janusgraph: Distributed graph database., 2017. Last accessed 28 December 2020. URL: http://janusgraph.org/

[14] Spring Framework. Spring framework. Available on: https://spring.io/. Access in, 3, 2018.

[15] Nadime Francis, Alastair Green, Paolo Guagliardo, Leonid Libkin, Tobias Lindaaker, Victor Marsault, Stefan Plantzikow, Mats Rydberg, Petra Selmer, and Andrés Taylor. Cypher: An evolving query language for property graphs. In Proceedings of the 2018 International Conference on Management of Data, pages 1433–1445. ACM, 2018.

[16] Alexandre P Francisco, Miguel Bugalho, Mário Ramírez, and João A Carriço. Global optimal ebust analysis of multilocus typing data using a graphic matroid approach. BMC bioinformatics, 10(1):152, 2009.

[17] Alexandre P Francisco, Cátia Vaz, Pedro T Monteiro, José Melo-Cristino, Mário Ramírez, and Joao A Carriço. Phyloviz: phylogenetic inference and data visualization for sequence based typing methods. BMC bioinformatics, 13(1):87, 2012.

[18] Google. Openidconnect. Last accessed 28 December 2020. URL: https://developers.google.com/identity/protocols/OpenIDConnect.

[19] Dick Hardt. The oauth 2.0 authorization framework. 2012.

[20] Florian Holzschuher and René Peinl. Performance of graph query languages: Comparison of cypher, gremlin and native access in neo4j. In Proceedings of the Joint EDBT/ICDT 2013 Workshops, EDBT ’13, pages 195–204, New York, NY, USA, 2013. ACM. URL: http://doi.acm.org/10.1145/2457317.2457351 doi:10.1145/2457317.2457351.

[21] Docker Inc. Docker. Last accessed 28 December 2020. URL: https://www.docker.com/

[22] F. Inc. Allegrograph., 2004. Last accessed 28 December 2020. URL: https://franz.com/agraph/

[23] NHOGS Interactive. Popoto.js. Last accessed 28 December 2020. URL: http://www.popotojs.com/

[24] Keith A. Jolley and Martin CJ Maiden. Bigsdb: Scalable analysis of bacterial genome variation at the population level. BMC Bioinformatics, 11(1):595, 2010. doi:10.1186/1471-2105-11-595

[25] Avinash Lakshman and Prashant Malik. Cassandra: a decentralized structured storage system. ACM SIGOPS Operating Systems Review, 44(2):35–40, 2010.

[26] Ljubica Lazarevic. Keeping track of graph changes using temporal versioning. Last accessed 28 December 2020. URL: https://medium.com/neo4j/keeping-track-of-graph-changes-using-temporal-versioning-3b0f854536fa

[27] Jeremy Leipzig. A review of bioinformatic pipeline frameworks. Briefings in bioinformatics, 18(3):530–536, 2017.

[28] Bjørn-Arne Lindstedt. Multiple-locus variable number tandem repeats analysis for genetic fingerprinting of pathogenic bacteria. Electrophoresis, 26(13):2567–2582, 2005.
[29] Ann-Katrin Llarena, Bruno Filipe Ribeiro-Gonçalves, Diogo Nuno Silva, Jani Halkilahti, Miguel Paulo Machado, Michael Santos Da Silva, Anniina Jaakkonen, Joana Isidro, Crista Hämäläinen, Jasmin Joenperä, et al. Innuendo: A cross-sectoral platform for the integration of genomics in the surveillance of food-borne pathogens. *EFSA Supporting Publications*, 15(11):1498E, 2018.

[30] Leonardo Alexandre Luana Silva and Diogo Loureiro. Phyloviz-electron. Last accessed 28 December 2020. URL: [https://github.com/DrLDiogo/PHYLOViZ-Electron](https://github.com/DrLDiogo/PHYLOViZ-Electron).

[31] Martin CJ Maiden, Jane A Bygraves, Edward Feil, Giovanna Morelli, Joanne E Russell, Rachel Urwin, Qing Zhang, Jiaji Zhou, Kerstin Zurth, Dominique A Caugant, et al. Multilocus sequence typing: a portable approach to the identification of clones within populations of pathogenic microorganisms. *Proceedings of the National Academy of Sciences*, 95(6):3140–3145, 1998.

[32] Microsoft. The oauth 2.0 authorization framework: Bearer token usage. Last accessed 28 December 2020. URL: [https://tools.ietf.org/html/rfc6750](https://tools.ietf.org/html/rfc6750).

[33] Marta Nascimento, Adriano Sousa, Mário Ramirez, Alexandre P Francisco, João A Carriço, and Cátia Vaz. Phyloviz 2.0: providing scalable data integration and visualization for multiple phylogenetic inference methods. *Bioinformatics*, 33(1):128–129, 2016.

[34] Inc. Neo Technology. Bolt protocol. Last accessed 28 December 2020. URL: [https://boltprotocol.org/](https://boltprotocol.org/).

[35] Neo4j. Indexes for search performance. Last accessed 28 December 2020. URL: [https://neo4j.com/docs/cypher-manual/current/administration/indexes-for-search-performance/](https://neo4j.com/docs/cypher-manual/current/administration/indexes-for-search-performance/).

[36] Neo4j. Neo4j java driver 4.2 api. Last accessed 28 December 2020. URL: [https://neo4j.com/docs/api/java-driver/current/](https://neo4j.com/docs/api/java-driver/current/).

[37] Inc Neo4j. *The Neo4j Graph Algorithms User Guide v3.5*. Neo4j.

[38] OpenID. Openidconnect. Last accessed 28 December 2020. URL: [https://openid.net/connect/](https://openid.net/connect/).

[39] Inc. Pivotal Software. Spring framework. Last accessed 28 December 2020. URL: [https://spring.io/](https://spring.io/).

[40] PubMLST. Pubmlst. Last accessed 28 December 2020. URL: Available:[http://pubmlst.org/](http://pubmlst.org/).

[41] Roshan Punnoose, Adina Crainiceanu, and David Rapp. Rya: A scalable rdf triple store for the clouds. In *Proceedings of the 1st International Workshop on Cloud Intelligence*, Cloud-I ’12, pages 4:1–4:8, New York, NY, USA, 2012. ACM. URL: [http://doi.acm.org/10.1145/2347673.2347677](http://doi.acm.org/10.1145/2347673.2347677), doi:10.1145/2347673.2347677.

[42] Philip Rathle. Official release: 3 essentials of neo4j 3.0, from scale to productivity and deployment, 2016. Last accessed 28 December 2020. URL: [https://neo4j.com/blog/neo4j-3-0-massive-scale-developer-productivity/](https://neo4j.com/blog/neo4j-3-0-massive-scale-developer-productivity/).

[43] Ian Robinson, Jim Webber, and Emil Eifrem. Graph database internals. In *Graph Databases* [44], pages 149–170.

[44] Ian Robinson, Jim Webber, and Emil Eifrem. *Graph Databases*. O’Reilly Media, Inc., 2013.
[45] Ian Robinson, Jim Webber, and Emil Eifrem. Introduction. In *Graph Databases* [44], pages 1–10.

[46] Ian Robinson, Jim Webber, and Emil Eifrem. Options for storing connected data. In *Graph Databases* [44], pages 11–24.

[47] Luana Silva. phylolib. Last accessed 28 December 2020. URL: https://github.com/Luanab/phylolib

[48] BG Spratt. Multilocus sequence typing: molecular typing of bacterial pathogens in an era of rapid dna sequencing and the internet. *Current opinion in microbiology*, 2(3):312–316, June 1999. URL: https://doi.org/10.1016/S1369-5274(99)80054-X, doi:10.1016/s1369-5274(99)80054-x

[49] Spring. Spring framework documentation. Last accessed 28 December 2020. URL: https://docs.spring.io/spring-framework/docs/current/reference/html/web.html

[50] Andrew Bowman Neo4j Staff. Neo4j security. Last accessed 28 December 2020. URL: https://community.neo4j.com/t/neo4j-security/16044

[51] Michael Hunger Neo4j Staff. 5 tips & tricks for fast batched updates of graph structures with neo4j and cypher. Last accessed 28 December 2020. URL: https://medium.com/neo4j/5-tips-tricks-for-fast-batched-updates-of-graph-structures-with-neo4j-and-cypher-73c7f693c8cc

[52] Andreia Sofia Teixeira, Pedro T Monteiro, João A Carriço, Mário Ramirez, and Alexandre P Francisco. Not seeing the forest for the trees: size of the minimum spanning trees (msts) forest and branch significance in mst-based phylogenetic analysis. *Plos one*, 10(3):e0119315, 2015.

[53] Andreia Sofia Teixeira, Pedro T Monteiro, Joao A Carriço, Francisco C Santos, and Alexandre P Francisco. Using spark and graphx to parallelize large-scale simulations of bacterial populations over host contact networks. In *International Conference on Algorithms and Architectures for Parallel Processing*, pages 591–600. Springer, 2017.

[54] Cátia Vaz, Alexandre P. Francisco, Mickael Silva, Keith A. Jolley, James E. Bray, Hannes Pouseele, Joerg Rothganger, Mário Ramirez, and João A. Carriço. Typon: the microbial typing ontology. *Journal of Biomedical Semantics*, 5(1):43, 2014. doi:10.1186/2041-1480-5-43

[55] Cátia Vaz, Marta Nascimento, João A Carriço, Tatiana Rocher, and Alexandre P Francisco. Distance-based phylogenetic inference from typing data: a unifying view. *Briefings in Bioinformatics*, 07 2020. bbaa147. arXiv:https://academic.oup.com/bib/advance-article-pdf/doi/10.1093/bib/bbaa147/33551477/bbaa147.pdf, doi:10.1093/bib/bbaa147

[56] Christophe Willemsen. Cypher: Write fast and furious. Last accessed 28 December 2020. URL: https://neo4j.com/blog/cypher-write-fast-furious/

[57] Zhemin Zhou, Nabil-Fareed Alikhan, Martin J Sergeant, Nina Luhmann, Cátia Vaz, Alexandre P Francisco, João André Carriço, and Mark Achtman. Grapetree: visualization of core genomic relationships among 100,000 bacterial pathogens. *Genome research*, 28(9):1395–1404, 2018.
Appendices
Example Workflow

![Figure 1: Example of a workflow system.](image)

In this example of a workflow system shown in Figure 1, can be interpreted that, an input is provided to the system, which then applies the transformations T1, and T2 sequentially. With the result of the transformation T2, it is applied the transformations T3x, and T3y parallelly. Afterwards, each result produced by the latter transformations is merged, and the transformation T4 is applied producing the final output.

**Neo4j Algorithms**

| Algorithm                  | Objective                                                                                                                                 |
|----------------------------|------------------------------------------------------------------------------------------------------------------------------------------|
| Louvain                    | Is an algorithm for detecting communities in networks. It maximizes a modularity score for each community, where the modularity quantifies the quality of an assignment of nodes to communities by evaluating how much more densely connected the nodes within a community are, compared to how connected they would be in a random network. |
| Label Propagation          | Is a fast algorithm for finding communities in a graph. It detects these communities using network structure alone as its guide, and doesn’t require a pre-defined objective function or prior information about the communities. |
| Connected Components       | Finds sets of connected nodes in an undirected graph where each node is reachable from any other node in the same set. It differs from the Strongly Connected Components algorithm (SCC) because it only needs a path to exist between pairs of nodes in one direction, whereas SCC needs a path to exist in both directions. |
| Strongly Connected Components | Finds sets of connected nodes in a directed graph where each node is reachable in both directions from any other node in the same set. |
| Triangle Counting / Clustering Coefficient | Is used to determine the number of triangles passing through each node in the graph. A triangle is a set of three nodes, where each node has a relationship to all other nodes. |
| Balanced Triads            | Is used to evaluate structural balance of the graph. Balance theory differentiates between positive and negative relationships. Certain structures between individuals and objects are perceived as balanced whereas others are not. |

Table 1: Community Detection Algorithms.
| Algorithm       | Objective                                                                                                                                 |
|-----------------|-------------------------------------------------------------------------------------------------------------------------------------------|
| PageRank        | Measures the connectivity of nodes. It counts the number, of links to a page which determines an estimation of how important the page is. The underlying assumption is that pages of importance are more likely to receive a higher volume of links from other pages. |
| ArticleRank     | Is a variant of the PageRank Where ArticleRank differs to PageRank is that PageRank assumes that relationships from nodes that have a low out-degree are more important than relationships from nodes with a higher out-degree. ArticleRank weakens this assumption. |
| Betweenness Centrality | Is a way of detecting the amount of influence a node has over the flow of information in a graph. It calculates the shortest (weighted) path between every pair of nodes in a connected graph, using the breadth-first search algorithm. Each node receives a score. Nodes that most frequently lie on these shortest paths will have a higher betweenness centrality score. |
| Closeness Centrality | Is a way of detecting nodes that are able to spread information very efficiently through a graph. For each node, the Closeness Centrality algorithm calculates the sum of its distances to all other nodes, based on calculating the shortest paths between all pairs of nodes. The resulting sum is then inverted to determine the closeness centrality score for that node. |
| Harmonic Centrality | Is a variant of closeness centrality, that was invented to solve the problem the original formula had when dealing with unconnected graphs. Rather than summing the distances of a node to all other nodes, the harmonic centrality algorithm sums the inverse of those distances. |
| Eigenvector Centrality | Measures the connectivity of nodes. Relationships to high-scoring nodes contribute more to the score of a node than connections to low-scoring nodes. A high score means that a node is connected to other nodes that have high scores. |
| Degree Centrality | Measures the number of incoming and outgoing relationships from a node. It can help to find popular nodes in a graph, and can be used to find the popularity of individual nodes, but it is often used as part of a global analysis where we calculate the minimum degree, maximum degree, mean degree, and standard deviation across the whole graph. |

Table 2: Centrality Algorithms.
**Algorithm Objective**

| Algorithm                          | Objective                                                                                                                                                                                                 |
|------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Minimum Weight Spanning Tree       | Starts from a given node, and finds all its reachable nodes and the set of relationships that connect the nodes together with the minimum possible weight. Prim’s algorithm is one of the simplest and best-known minimum spanning tree algorithms. |
| Shortest Path                      | Calculates the shortest (weighted) path between a pair of nodes. In this category, Dijkstra’s algorithm is the most well known.                                                                             |
| Single Source Shortest Path        | Calculates the shortest (weighted) path from a node to all other nodes in the graph.                                                                                                                      |
| All Pairs Shortest Path            | Calculates the shortest (weighted) path between all pairs of nodes. This algorithm has optimisations that make it quicker than calling the Single Source Shortest Path algorithm for every pair of nodes in the graph |
| A*                                | Improves on the classic Dijkstra algorithm. It is based upon the observation that some searches are informed, and that by being informed we can make better choices over which paths to take through the graph. |
| Yen’s K-shortest paths             | Computes single-source K-shortest loopless paths for a graph with non-negative relationship weights.                                                                                                |
| Random Walk                        | Provides random paths in a graph. We start at one node, choose a neighbor to navigate to at random or based on a provided probability distribution, and then do the same from that node, keeping the resulting path in a list. |

Table 3: Path finding Algorithms.

| Algorithm                         | Objective                                                                                                                                                                                                 |
|-----------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Jaccard Similarity                | Measures similarities between sets. It is defined as the size of the intersection divided by the size of the union of two sets.                                                                           |
| Cosine Similarity                 | Is the cosine of the angle between two n-dimensional vectors in an n-dimensional space. It is the dot product of the two vectors divided by the product of the two vectors’ lengths (or magnitudes). |
| Pearson Similarity                | Is the covariance of the two n-dimensional vectors divided by the product of their standard deviations.                                                                                                  |
| Euclidean Distance                | Measures the straight line distance between two points in n-dimensional space.                                                                                                                           |
| Overlap Similarity                | Measures overlap between two sets. It is defined as the size of the intersection of two sets, divided by the size of the smaller of the two sets.                                                          |
| Approximate Nearest Neighbors     | Constructs a k-Nearest Neighbors Graph for a set of objects based on a provided similarity algorithm. The similarity of items is computed based on Jaccard Similarity, Cosine Similarity, Euclidean Distance, or Pearson Similarity. |

Table 4: Similarity Algorithms.
Algorithm | Objective
--- | ---
Adamic Adar | Is a measure used to compute the closeness of nodes, based on their shared neighbors.
Common Neighbors | Captures the idea that two strangers who have a friend in common are more likely to be introduced than those who don’t have any friends in common.
Preferential Attachment | Is a measure used to compute the closeness of nodes, based on their shared neighbors.
Resource Allocation | Is a measure used to compute the closeness of nodes, based on their shared neighbors.
Same Community | Is a way of determining whether two nodes belong to the same community.
Total Neighbors | Computes the closeness of nodes, based on the number of unique neighbors that they have. It is based on the idea that the more connected a node is, the more likely it is to receive new links.

Table 5: Link Prediction Algorithms.