The Graph Pattern Matching Problem through Parameterized Matching

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(Dedication)

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Abstract

We propose a new approach to solve graph isomorphism using parameterized matching. Parameterized matching is a string matching problem where two strings parameterized-match if there exists a bijective function, on the symbols of the alphabet, that maps one of the strings into the other. Given that parameterized matching is defined for linear structures, we define the concept of graph linearization to represent the topology of a graph as a walk on it. Then, our approach to determine whether two graphs are isomorphic consists of determining whether there exists a walk in one of the graphs that parameterized-matches a linearization of the other graph.

Our solution has two main steps: linearization and matching. We develop an efficient linearization algorithm, that generates short linearizations with an approximation guarantee, and develop a graph matching algorithm. We show that this solution also works for subgraph isomorphism, which is the problem of determining whether an input graph $H$ is isomorphic to a subgraph of another input graph $G$. We evaluate our approach experimentally on graphs of different types and sizes, and compare to the performance of VF2, which is a prominent algorithm for graph isomorphism. Our empirical measurements show that graph linearization finds a matching graph faster than VF2 in many cases, especially in Miyazaki-constructed graphs which are known to be one of the hardest cases for graph isomorphism algorithms.

We extend this approach to query attributed graphs. An attributed graph is a graph data structure, in which nodes and edges may have identifiers, types and other attributes.Attributed graphs are used in many application domains, for example to model social networks in which nodes represent people, photos, and postings and edges represent friendship, person-tagged-in-photo and mentioned-in-post relationships. Queries are used to extract information from such graphs. Several graph queries are expressed as graph pattern matching, which is the problem of finding all instances of pattern match query $P$ in a larger attributed graph $G$. A pattern match query may specify both a graph structure and predicates on the attributes of the graph elements. Clearly, this problem is associated to subgraph isomorphism.

Furthermore, we define a more general class of graph queries called generalized pattern queries on attributed multigraphs. The goal of this class is to find paths and subgraphs that satisfy query reachability and predicates. The query language is expressive: It allows (i) using regular expression operators (e.g., Kleene star and union); (ii) specifying structural predicates on graph nodes and edges; and (iii) using attribute predicates on nodes and edges. Pattern match queries, reachability queries, their combination, and even more queries can be expressed through generalized pattern queries. We use our approach to solve this new type of queries.

The proposed technique has two phases. First, the query is linearized, i.e., represented as a graph walk that covers all nodes and edges. There are several linearizations for a given query; we derive
heuristics to produce a good linearization that is short and places selective predicates early in the linearization. Second, we search for a bijective function that maps each element of the query to an element of the attributed multigraph that satisfies the reachability requirements and the predicates. Specifically, we develop an algorithm that matches the linearization by traversing the attributed graph in a manner similar to a breadth first traversal constrained by the linearization. We evaluate our solution experimentally using a real graph (the DBLP citation network) to assess its practicality and efficiency. Our results show that our techniques and optimizations are effective in querying attributed graphs, offering several factors of reduction in query response time when graph statistics are utilized.

**Keywords:** parameterized matching, graph theory, graph algorithms, graph matching, pattern matching, graph isomorphism, subgraph isomorphism, attributed graphs, graph queries, social networks.

**Resumen**

En esta tesis se propone un nuevo enfoque de solución para resolver el problema de isomorfismo de grafos usando búsqueda parametrizada. La búsqueda parametrizada es un problema de búsqueda de cadenas de texto en el cual dos cadenas coinciden si existe una biyección que mapee los símbolos de una cadena en los símbolos de la otra. Dado que la búsqueda parametrizada está definida para estructuras lineales, se define el concepto de linearización de grafos para representar la topología de un grafo como un camino sobre este. Entonces, la solución para determinar si dos grafos son isomorfos consiste en determinar si existe un camino en uno de los grafos que haga coincidencia parametrizada con la linearización del otro grafo.

La solución propuesta tiene dos pasos: linearización y búsqueda. Se presenta un algoritmo eficiente que genera linearizaciones aproximadamente óptimas en longitud, y un algoritmo de búsqueda. Se demuestra que esta solución también resuelve el problema de isomorfismo de subgrafos, en el cual se determina si un grafo $H$ es isomorfo a un subgrafo de otro grafo $G$. Se evaluó experimentalmente la solución con grafos de diferentes tipos y tamaños. Se comparó su desempeño con el de VF2, el cual es un algoritmo competitivo de isomorfismo de grafos. Los resultados experimentales muestran que la solución propuesta es más eficiente que VF2 en varios casos, en especial en grafos Miyazaki, los cuales se caracterizan por constituir uno de los casos más difíciles para los algoritmos de isomorfismo de grafos.

Este enfoque de solución se extiende para resolver consultas sobre grafos semánticos. Un grafo semántico es un grafo en el cual los nodes y arcos pueden tener identificadores, tipos y otros
atributos. Estos grafos tienen aplicaciones importantes en diversas áreas, como por ejemplo para modelar redes sociales en las que los nodos representan personas, fotos y publicaciones y los arcos representan relaciones de amistad, etiquetado y mención. Se usan consultas para extraer información de estos grafos. Varias de estas consultas se expresan como búsqueda de patrones, la cual consiste en encontrar las coincidencias del grafo patrón $P$ en un grafo semántico $G$. El grafo patrón especifica tanto la estructura de las coincidencias a encontrar, como los predicados sobre los atributos que deben cumplir los nodos y los arcos de las mismas. Claramente, este problema está asociado al isomorfismo de subgrafos.

Además, se define un tipo de consultas más general sobre grafos semánticos. Estos nuevos patrones se denominan grafos patrón generalizados. El objetivo de estos es encontrar caminos y subgrafos que satisfagan ciertos requisitos semánticos, de estructura y de alcance. Estos patrones son expresivos, pues permiten (i) usar operadores de expresiones regulares (e.g., la estrella de Kleene y la unión); (ii) especificar predicados estructurales en los nodos y arcos; y (iii) evaluar predicados sobre los atributos de los nodos y arcos. Los patrones grafo tradicionales, las consultas de alcance, la combinación de estos y más se pueden representar a través de grafos patrón generalizados. Se usa el enfoque de solución propuesto para resolver los grafos patrón generalizados.

La solución tiene dos fases. Primero, el patrón es linearizado, i.e., representado como un camino que incluye todos sus nodos y arcos. Hay muchas linearizaciones para un patrón dado; se proponen heurísticas para producir linearizaciones cortas que ubican los predicados selectivos al comienzo. Segundo, se busca una función biyectiva que mapee cada nodo en el patrón a un nodo en el grafo semántico que satisfaga los requisitos de alcance y los predicados. Específicamente, se propone un algoritmo de búsqueda que recorre el grafo semántico siguiendo una búsqueda en amplitud restringida por la linearización. La solución se evaluó experimentalmente usando un grafo semántico real (la red de citas DBLP) para evaluar su practicidad y eficiencia. Los resultados experimentales muestran que las técnicas y optimizaciones propuestas son efectivas en consultar grafos semánticos, ofreciendo un alto factor de reducción en el tiempo de ejecución cuando se utilizan las estadísticas del grafo semántico.

**Palabras clave:** búsqueda parametrizada, teoría de grafos, algoritmos de grafos, búsquedas en grafos, búsqueda de patrones, isomorfismo de grafos, isomorfismo de subgrafos, grafos semánticos, redes sociales.
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1. Introduction

Graphs are interesting data structures due to their expressive power that allows them to represent real-world phenomena in diverse areas. Graphs’ expressive power lies in their ability to represent different kinds of concepts and the relationships among such concepts. Nowadays, graph-based models are found in different domains where both concepts and relationships contain rich and diverse information expressed by types and attributes. In some applications, graphs contain millions of concepts and it is required to support queries efficiently. However, the growing size of graph databases and the richness and the variety of their data have made it difficult to resolve this problem in reasonable time. Given that existing solutions are not able to deal with the current demands, it is quite pertinent to keep exploring new solutions from different approaches in order to efficiently query large graphs. In this thesis, we apply a string matching technique called parameterized matching to evaluate the isomorphism and containment relations between graphs with or without attributes.

In the rest of this chapter, we examine these problems and their relationships more closely. Specifically, in Section 1.1, we consider some of the most fundamental concepts and application areas of graphs. The different variants of the graph matching problem are introduced in Section 1.2. Then, in Section 1.3, some applications of querying graphs are discussed. The basic concepts of parameterized matching are presented in Section 1.4. Finally, in Section 1.5 our contributions are outlined.

1.1. Graphs: Concepts and Applications

A graph $G = (V, E)$ consists of a set $V$ of nodes (or vertices), $n = |V|$, and a set $E$ of edges, $m = |E|$, where the edges are ordered pairs of the nodes that represent links between them, i.e., $E \subseteq V \times V$. Let $E_G = V \cup E$ denote the set of graph elements of $G$, i.e., the set of nodes and edges in $G$. In Figure 1-1, three examples of graphs are presented. For more generality, in this thesis we consider multigraphs. A multigraph is a graph where multiple edges between two distinct nodes and self loops are permitted. We distinguish the edges that have the same end nodes by the notation of the edge; for example, $e = (u, v)$ and $e' = (u, v)$.

Multigraphs are useful for representing sets of entities of different kinds and their relationships. Some of the domains where multigraphs can be found include the Semantic Web [131], the worldwide web [3], communication networking [20], social networking [114], interactive gaming [91],
Figure 1-1.: Examples of graphs. Graphs (a) and (b) are isomorphic. There exists a subgraph in graph (c) that is isomorphic to graph (a) and (b).

geographic information systems [98], pattern recognition [126], pattern analysis [146], computer vision [65], artificial intelligence [118], information retrieval [119], knowledge discovery [100], data mining [153], electronics [138], computer aided design [86], chemoinformatics [67] and bioinformatics [92]. For instance, multigraphs are used to provide structural descriptions of images by decomposing them into different components that are modelled through nodes and the relationships of such components are modelled through edges [44]. Some of the types of images that have been described in this way are handwritten characters, ideograms and symbols [45]. On the other hand, in bioinformatics, several types of information can be represented through multigraphs: a protein structure, considering the set of residues as the nodes and their spacial proximity as the edges, or a protein interaction network where the nodes represent the proteins and the edges represent the physical interactions [78].

However, multigraphs are not only useful to represent real-world phenomena with nodes of the same type connected by edges of the same type. In many applications, it is necessary to use multigraphs with attributes, called attributed multigraphs, which are multigraphs where both the set of nodes and the set of edges are sets of entities with different types and characteristics. For example, if the domain is music, we may have an information network where node types are singer, song and album, and edge types are performs and containedIn to connect singer with song and song with album, respectively. The attributes of singer may be name, birthday and website while the attributes of album may be name, length and year. Moreover, there is an ontology associated to an attributed multigraph that establishes the possible concepts, the types of relationships permitted between two types of concepts and the restrictions on the attributes of both concepts and relationships [69]. For instance, the ontology associated to the multigraph of our running example would forbid a relationship of type performs between concepts of types album and song.

Another important area where attributed multigraphs are used is social networks. For instance, in Figure 1-2, we show an example of a social network where nodes represent people and photos.
while edges establish friendship and person-tagged-in-photo relationships.

Figure 1-2.: Example of a social network represented as an attributed multigraph.

Considering the great amount of information that is represented through multigraphs nowadays, in these and in several other areas, the problem of querying multigraphs has significantly gained importance in recent years. In the next section, we introduce the main concepts of graph matching.

1.2. Graph Matching Problems

There are different problems associated to matching multigraphs. We first consider the general problems tackled in theoretical computer science and then the problems of practical interest for attributed multigraphs. The most basic problem is graph isomorphism which consists of determining whether two multigraphs have the same structure, i.e., there exists a bijection that associates the nodes/edges of the two multigraphs such that the adjacency relation is preserved. More formally, graph isomorphism can be defined as follows.
1.2 Graph Matching Problems

Problem 1 (Graph Isomorphism). Let $G_1 = (V_1, E_1)$ and $G_2 = (V_2, E_2)$ be two multigraphs such that $n = |V_1| = |V_2|$ and $m = |E_1| = |E_2|$. The Graph Isomorphism problem determines whether there exists a bijective mapping function $f : E_{G_1} \rightarrow E_{G_2}$, such that

$$\forall u, v \in V_1, e = (u, v) \in E_1 \iff f(u), f(v) \in V_2 \land f(e) = (f(u), f(v)) \in E_2$$

For example, the multigraph in Figure 1-1(a) is isomorphic to the multigraph in Figure 1-1(b) under the bijection $f : (A, B, C, D, E) \rightarrow (H, F, I, G, J)$. Other example of graph isomorphism is presented in Figure 1-3(a,b); furthermore there are two possible mapping functions that define the isomorphism (see Figure 1-3(c)). A closely related problem is subgraph isomorphism.

Problem 2 (Subgraph Isomorphism). Let $G_1$ and $G_2$ be two multigraphs. The Subgraph Isomorphism problem consists of determining whether there exists a subgraph in $G_2$ isomorphic to $G_1$. For example, if we remove the node $P$ and its adjacent edges from the multigraph in Figure 1-1(c), we obtain a multigraph that is isomorphic to the multigraphs presented in Figures 1-1(a) and (b). A naive solution for these problems could search for all the possible mappings; however, its search space is exponential.

A lot of research about both graph and subgraph isomorphism has been carried out. Interestingly, even though subgraph isomorphism has been proven to be NP-Complete, the exact complexity of graph isomorphism has not been determined yet [44]. Due to the similarity of the problems, most of the existing solutions solve both of them. In particular, Ullmann’s algorithm [148] is the traditional solution. Notwithstanding, a more recent algorithm, called VF2, experimentally outperformed...
Ullmann’s algorithm for many cases [44]. On the other hand, the NAUTY algorithm [102] is another traditional solution for only graph isomorphism. These algorithms have exponential worst-case performance since isomorphism is a hard problem. Except for some easy cases, solving isomorphism generally takes much longer time if there is no match; in such case, all the possible mappings are progressively searched until shown not to lead to an isomorphism. Several heuristics, however, are employed to find likely mappings quickly. A good algorithm for determining isomorphism should quickly find isomorphic multigraphs in many cases. In this thesis, we propose a new approach to solve both graph and subgraph isomorphism that makes use of some heuristics to detect isomorphism at an early stage of the search.

On the other hand, some models of interesting queries on attributed multigraphs have been proposed. Particularly, reachability queries consist of determining whether two nodes in the multigraph are somehow connected through an unrestricted path. For example, considering the multigraph of Figure 1-2, we can say that Photo 3 is reachable from Alice as one of her friends is tagged in such photo. Besides large-scale social networking, reachability queries have important applications in several other areas. For instance, on biological multigraphs, it is relevant to find genes whose expressions are influenced by a given molecule [149]. Moreover, reachability queries are also useful to query XML databases and domain ontologies [84].

In recent years, special types of constrains, like the permitted edge types on the connecting path, have been included in reachability queries [83]. For example, considering the attributed multigraph of Figure 1-2, we might want to know whether Dave is connected to a female using friend edges, i.e., whether there is a female in his network. The output is true as Alice can be reached through paths coming from either Bob or Chris. Later, reachability queries were extended to support regular expressions that establish the edges types on the connecting path [59]. However, such regular expressions have limited expressive power as they do not support the Kleene operator nor predicates on intermediate nodes. Then, a model that supports these features was developed [127, 128].

Other type of queries on attributed multigraphs is pattern match queries. Each pattern match query is a query multigraph that searches for matches in an attributed multigraph such that: (i) the adjacency relation of the matches is the same as the one of the query; and (ii) each node/edge in the query specifies a predicate to be satisfied by its corresponding node/edge in the match. More formally, the problem of finding all the matches can be defined as follows.

**Problem 3** (Pattern Match Query Problem). Let $P$ be a pattern match query and $G$ be an attributed multigraph. The Pattern Match Query problem consists of finding the set of subgraphs of $G$ that are isomorphic to $P$, and whose graph elements satisfy the predicates on the corresponding graph elements in $P$.

This problem is clearly associated to subgraph isomorphism; thus, it can be solved with straight-
forward adaptations of subgraph isomorphism algorithms that include predicate evaluations. For example, in Figure 1-4(a), we show a pattern match query that aims to find a pair of friends, where one of them is a female, that are tagged in a photo. The output of this query on the attributed multigraph of Figure 1-2 is presented in Figure 1-4(b) and (c).

![Diagram of a pattern match query](image)

**Figure 1-4.** Example of a pattern match query for the attributed multigraph presented in Figure 1-2. (a) Pattern match query. (b, c) Output reported.

Later, a new type of query on attributed multigraphs was introduced [59]. These queries, called *pattern queries*, constitute a combination of reachability and pattern match queries. Specifically, the queries are multigraphs where each node is associated to a predicate and each edge, along with its end nodes, establishes a reachability query. The output is associated to the set of nodes in the attributed multigraph that correspondingly satisfy both the predicates and the reachability queries in the query graph. However the output expressed corresponds to the set of global matches for each edge, i.e., reachability query in the query graph [59]. Then, the relative relationships between the nodes presented in the output is not easy to interpret. Moreover, these queries do not support the Kleene operator nor predicates on intermediate nodes [59].
In this thesis, we introduce *generalized pattern queries* as a new type of queries that evaluates attribute predicates, structural requirements and reachability requirements. These queries, besides allowing edge-to-path mappings, also support predicates on intermediate nodes/edges and operators like union and the Kleene star in the reachability requirements. Moreover, the output produced is easy to interpret: it consists of the set of all the solution instances where each instance is an ordered set of nodes that correspondingly satisfy the nodes in the query; thus, the relative relationships between the output nodes from each solution instance and the query is straightforward to determine. Then, the graph matching problems we consider in this thesis are graph isomorphism, subgraph isomorphism, solving pattern match queries and solving generalized pattern queries. In the next section, we discuss some applications of these problems.

### 1.3. Applications of Graph Matching

The graph matching problems have applications in different domains [78, 159]. Some examples are shown below:

- To find all heterocyclic chemical compounds that contain a given aromatic ring and a side chain. In this context, chemical compounds are modelled as graphs where the nodes represent atoms and the edges represent bonds.

- To find all protein structures that contain an $\alpha$-$\beta$-barrel motif that is specified as a cycle of $\beta$ strands embraced by another cycle of $\alpha$ helices [27].

- To determine whether a protein complex query from one species is functionally conserved in another species. The protein complex can be represented as a graph where the nodes are proteins labelled by Gene Ontology.

- To find all the instances from a Resource Description Framework (RDF) graph where two departments of a company share the same shipping company. The nodes are of type *department* and *company* and the edges of type *shipping*.

- To locate the occurrences of a suspicious bug that arises as a distortion in the control flow within a large software system that can be represented as a large static or dynamic call graph [55].

- To find all the co-authors from a bibliographic information network, such as DBLP, in a specified set of conference proceedings.

Furthermore, there are many other applications of graph matching in different areas including: 2D and 3D image analysis [135, 96, 150, 144], image database [80, 121], video analysis [133, 71, 125],
1.4 Parameterized Matching

In this section, we introduce string matching as we propose to use it to solve the graph matching problems. String matching is definitely one of the foremost and most basic and useful computational primitives [8]. The input to the string pattern matching problem consists of two strings: the pattern \( P = P_1...m \) and the text \( T = T_1...n \). The output should list all the occurrences of the pattern in the text, i.e., all the positions \( i \) in \( T \) such that \( P_j = T_{i+j-1} \) for all \( 1 \leq j \leq m \). Note that the symbols in the strings are chosen from some set which is called an alphabet. An alphabet could be any collection of symbols and it is normally drawn from a set of pre-existing characters which is habitually designated as the common ASCII code set. Over the years, several variants of this problem have been proposed in order to support a wider range of applications. For instance, in the early nineties, a string matching variant called parameterized matching was proposed as an aid to detect duplicate code in large software systems.

Duplication in code occurs when there are some sections of code that are exactly equal (such as literals and reserved keywords) and some other sections of code that are the same, except for a systematic change of parameters (such as identifiers or constants). Then, the code can be seen as a string of tokens, where each token belongs to either of the following alphabets: (i) an alphabet \( \Sigma_C \) of constant symbols for the tokens of code sections that remain exactly the same; and (ii) an alphabet \( \Sigma_P \) of parameter symbols for the tokens of code sections that could have the mentioned systematic change. Then, the parameterized matching problem can be defined as follows.

Definition 1 (Parameterized-Match). Let \( \Sigma_C \) be the constant symbol alphabet and \( \Sigma_P \) be the parameter symbol alphabet, where \( \Sigma_C \) and \( \Sigma_P \) are disjoint. Two length-\( \ell \) strings \( X = X_1...\ell \) and \( Y = Y_1...\ell \), defined over \( (\Sigma_C \cup \Sigma_P)^* \), are said to be a parameterized-match, or a p–match, if there exists a bijective function \( g : \Sigma_C \cup \Sigma_P \mapsto \Sigma_C \cup \Sigma_P \) such that \( g(Y_i) = X_i, \ 1 \leq i \leq \ell \) so that \( g \) is identity for the the symbols from \( \Sigma_C \).

In other words, \( X \) and \( Y \) are a p-match if one string can be transformed into the other by renaming its parameters through a bijective function \( g : \Sigma_C \cup \Sigma_P \mapsto \Sigma_C \cup \Sigma_P \), such that \( g \) is identity for the constant symbols. Note that, \( g \) can be chosen from \( |\Sigma_P|! \) different possible mapping functions. As an example, Figure 1-5 shows two equal-length strings \( X = xyxybx \) and \( Y = zbxzbxz \) defined over \( \Sigma_C \cup \Sigma_P \), where \( \Sigma_C = \{b\} \) and \( \Sigma_P = \{x,y,z\} \). In Figure 1-5(a), the 6 possible bijective functions from the symbols in \( Y \) to the symbols in \( X \) are shown. We conclude that \( X \) and \( Y \)

\footnote{American Standard Code for Information Interchange.}
parameterized-match because there is a function, specifically $r$, such that $r(Y_i) = X_i$ for every $1 \leq i \leq 7$, and the only constant symbol, $b$, has an identity mapping, i.e., $r(b) = b$.

![Table and Diagram](image)

**Figure 1-5.** Example of a parameterized-match between the strings $X = xbyyxby$ and $Y = zbxxzbz$ both defined over $\Sigma_C \cup \Sigma_P$, where $\Sigma_C = \{b\}$ is the constant alphabet and $\Sigma_P = \{x, y, z\}$ is the parameter alphabet. (a) All the 6 possible bijective functions from the symbols in $Y$ to the symbols in $X$ such that the constant symbol has an identity mapping. (b) Successful attempt to transform $Y$ into $X$ through $r$.

Furthermore, two equal-length strings $X$ and $Y$ that parameterized-match have the same structure. Let us suppose that $i$ and $j$ are the only occurrences of the symbol $\alpha$ in $Y$. Then, the existence of a bijective function $g$ that maps the symbols in $Y$ to the symbols in $X$ implies that $g(\alpha) = X_i = X_j = \beta$ and that $\beta$ has no other occurrences in $X$. As this applies for all the distinct symbols $\alpha$ in $Y$, we can conclude that the following facts hold: (i) $X$ and $Y$ have the same number of distinct symbols; (ii) the first occurrence of each distinct symbol $\alpha$ in $Y$ takes place in the same position of the first occurrence of the symbol $g(\alpha)$ in $X$; and (iii) the relative distances among the different occurrences of each $\alpha$ in $Y$ are the same relative distances among the occurrences of $g(\alpha)$ in $X$. Therefore, two strings that parameterized-match have the same structure, i.e., they are the same except for a systematic change of the symbols. We illustrate these properties for our running example in Figure 1-6. For instance, notice that the first occurrence of $x$ in $Y$ is at position 3 and its second occurrence is one position away; the occurrences of $y$ in $X$ take place at corresponding positions.

In 1993, Brenda Baker [15] was the first researcher to have addressed this problem, and many others [5, 9, 16, 17, 18, 41, 76, 68, 90, 53, 124] since have followed Baker’s work. She did, indeed, open up a wide field of extensive research that soon was generalized to other fields. Over the years,
1.5 Our Contributions

In this thesis, we propose a new application for parameterized matching: the solution of the graph matching problems. Our initial motivation for this is based on the fact that in parameterized matching we must determine whether two strings that have the same structure, i.e., the relative distances among the occurrences of the distinct symbols in the strings are equal (see Figure 1-6). Similarly, in graph matching, we must determine if two graphs have the same structure, i.e., if the graphs are isomorphic. Furthermore, in parameterized matching (and graph isomorphism) the match depends upon the existence of a bijection from the symbols (nodes) in one string (graph) to the symbols (nodes) in the other string (graph). Therefore, this thesis defines a new model for solving graph matching based on parameterized matching. Specifically, given two graphs $G_1$ and $G_2$, we represent $G_1$ in a linear manner which we call graph linearization. Then, we evaluate if this representation parameterized-matches a walk on $G_2$.

This document is comprised by two parts. In Part I, we propose a new solution for both graph and subgraph isomorphism. Then, in Part II, we adapt this solution to answer different types of queries on attributed graphs. In particular, we solve pattern match queries and generalized pattern queries.
More specifically, in Part I, we present our approach to solve isomorphism in multigraphs through parameterized matching as follows:

- We define graph linearization, as a linear representation of graphs, and formally show how it can be used to solve graph and subgraph isomorphism. We develop the *Graph Linearization Algorithm* – GLA, an asymptotically length-optimal algorithm that efficiently linearizes a graph through greedy heuristics (Chapter 3).

- We propose a matching algorithm, called PMG, that solves graph isomorphism. Given two multigraphs $G_1$ and $G_2$, PMG calculates a linearization of $G_1$ and determines whether there exists a walk on $G_2$ that parameterized-matches such linearization. If so, the graphs are isomorphic (Chapter 4).

- We adapt the PMG algorithm to solve subgraph isomorphism; this adaptation is called PMG-SI (Section 4.6).

Then, in Part II, we redefine our approach to efficiently solve queries of interest on attributed graphs as follows:

- We define the data model and the query model. Especially, we define generalized pattern queries, a new type of queries with high expressive power that supports structural predicates, attribute predicates and reachability evaluation (Chapter 5).

- We extend the concept of graph linearization to represent generalized pattern queries. Moreover, we show how the linearization of a generalized pattern query can be used to find its solution on an attributed graph. We present the *Enhanced–Graph Linearization Algorithm* – E-GLA, an algorithm that exploits the attributed graph statistics to generate linearizations of the query that will incur in low matching cost (Chapter 6).

- We develop a solution for generalized pattern queries on attributed graphs also based on query linearization through E-GLA (Chapter 7).

Some contents of this thesis have already been presented in [105, 110, 104, 106, 103, 109, 93]. Other articles also developed during the PhD program, that are related to this thesis, are [22, 108, 107, 111, 112, 23, 21, 46].
2. Related Work

In this thesis we propose a new solution for the graph matching problems based on parameterized matching. Thus, in this chapter we present a literature review of these topics. In particular, Section 2.1 describes the most efficient algorithms for graph isomorphism and subgraph isomorphism. Then, Section 2.2 considers graph matching in attributed graphs. Specifically, we cover different types of queries on attributed graphs, including reachability queries, pattern match queries and pattern queries. Finally, Section 2.3 includes the basic problems, solutions and extensions of parameterized matching.

2.1. Solutions for Graph and Subgraph Isomorphism

It has been proven that subgraph isomorphism is a NP-Complete problem [70]; however, the exact complexity of graph isomorphism is still an open question [70, 44]. Because the two problems are very related, most of the solutions for one of them works for the other. In this section, we describe the most efficient algorithms: Ullmann’s algorithm (Section 2.1.1) and VF2 (Section 2.1.2).

2.1.1. Ullmann’s Algorithm

One of the first algorithms for solving both problems was proposed by Ullmann back in 1976 [148]. His solution is similar to Corneil and Gotlieb’s algorithm for graph isomorphism [47] but differs from it in that graphs are not processed separately. Ullmann’s solution consists of backtracking in a search tree using a mechanism to prune the search space called refinement procedure. For generality, we describe the algorithm for subgraph isomorphism, but it also solves graph isomorphism. First, we show how the possible mappings are enumerated and then how to apply the refinement procedure.

Ullmann’s algorithm determines whether graph $G_1 = (V_1, E_1)$, where $n_1 = |V_1|$ and $m_1 = |E_1|$, is isomorphic to a subgraph in $G_2 = (V_2, E_2)$, where $n = |V_2|$ and $m = |E_2|$. Let us denote the adjacency matrices of $G_1$ and $G_2$ as $A$ and $B$, respectively. The possible mappings are stored in a binary matrix $M$ of $n_1$ rows and $n$ columns. In particular, $M[i][j]$, for $1 \leq i \leq n_1$ and $1 \leq j \leq n$, is equal to 1 if node $v_i \in V_1$ can be mapped to node $v_j \in V_2$; otherwise, $M[i][j] = 0$. An important property of this matrix is that it must contain exactly one 1 in each row and at most one 1 in each column. This is to ensure the injective mapping from the nodes of $G_1$ (i.e., the rows) to a subset of...
the nodes of $G_2$ (i.e., the columns).

The idea of the algorithm is exploring all the possible mappings by permuting the rows and columns of $B$ and comparing adjacency with $A$. This can be done by multiplying $B$ with the possible mappings $M$. Specifically, for a given $M$, the multiplication $MB$ moves row $j$ to row $i$, for all $M[i][j] = 1$. Thus, $(MB)^T$ moves column $j$ to column $i$. Similarly, $C = M(B)^T$ moves column $j$ to column $i$ and row $j$ to row $i$ for every $M[i][j] = 1$. Then, in order to evaluate if a given mapping $M$ is an isomorphism, we check whether the adjacency relation defined in $A$ is contained in $C$. In other words, if $M$ is an isomorphism, the following condition must be satisfied:

$$\forall_{i,j} (A[i][j] = 1) \Rightarrow (C[i][j] = 1) \quad (2-1)$$

Notice that, in case of graph isomorphism, $\Rightarrow$ is replaced by $\iff$ as the adjacency matrices must be equal.

The possible mappings are explored as follows. An initial matrix $M^0$ of $n_1$ rows and $n$ columns is constructed by setting $M^0[i][j] = 1$ if mapping $v_i \in V_1$ to node $v_j \in V_2$ is possible. Specifically, it is possible to do such mapping if the degree of $v_j$ is greater or equal to the degree of $v_i$ in the case of subgraph isomorphism; in the case of graph isomorphism, the degrees must be equal. Otherwise, we set $M^0[i][j] = 0$. Then, $M^0$ becomes the root of the search space of the permutation matrices. In particular, a Depth-First Search (DFS) approach is used: at level $k$ of the DFS search tree, the matrix $M^k$ only keeps one of the 1’s at the row $k$ of $M^{k-1}$; the others are reassigned to 0. Thus, at level $n_1$, the mapping for all nodes in $V_1$ should have been found. Therefore, for each matrix $M^k$, we evaluate condition 2-1 to determine if it is an isomorphism. Note that if, at any point, a given $M^k$ has a row with no 1’s, it is not necessary to explore its successors as it will not lead to an isomorphism.

Now Ullmann’s refinement procedure, which is the key idea of his algorithm to prune the search space, is introduced. If, at any point, a node $v_j \in V_2$ is among the possible mappings of a node $v_i \in V_1$, then every neighbour of $v_i$ must have at least one possible mapping among the neighbours of $v_j$. If this condition does not hold, we can safely remove $v_j$ from the possible mappings of $v_i$ as we know that forthcoming mappings cannot be established in this direction. Recall that, in terms of the mapping matrix $M$, the possible mappings for $v_i$ are the nodes $v_j$ for which $M[i][j] = 1$. Then, for each $M[i][j] = 1$ in matrix $M$, the refinement procedure must be evaluated as follows:

$$\forall_x (A[i][x] = 1) \Rightarrow \exists_y (M[x][y] \cdot B[y][j] = 1) \quad (2-2)$$

If this condition does not hold, then we set $M[i][j] = 0$. This is procedure is performed for every $M[i][j] = 1$. However, changing a 1 to 0 can make condition 2-2 is no longer satisfied for other 1’s in $M$. Then this process must be repeated over and over again until there is an iteration where no 1 in the matrix is changed to 0. The refinement procedure is applied for each matrix $M^k$ in the DFS
search tree, including $M^0$ [148]. The space complexity of Ullmann’s algorithm is $\Theta(n^3)$; its time complexity in the best case is $\Theta(n^3)$ while its worst-case time complexity is $\Theta(n! n^2)$ [44].

Other refinement procedures to reduce the search space have also been considered. Specifically, Haralick and Elliot proposed forward-checking and looking-ahead [74], and Kim and Kak used discrete relaxation [88]. Another approach that has been taken consists of a reduction to the maximal clique detection problem [56, 115]. Besides, Blake proposed a solution based on a partition according to lattice theory to reduce computational complexity [24].

However, Ullmann’s algorithm is one of the most used solution because of its good performance. Ullmann’s algorithm was compared against other graph matching solutions and it turned out to be the one with best matching time [113]. Furthermore, this algorithm also permits the comparison of semantic information during the process. Notwithstanding, for large graphs, the time required by Ullmann’s algorithm is still too high to be tractable. During the last three decades, there have been many attempts for solving the graph matching problem on large graphs. Most of them achieve low time complexity by imposing restrictions on the topology of the graphs. Some of the most important contributions in this direction are polynomial algorithms for trees, planar graphs and bounded valence graphs [95]. On the other hand, some algorithms based on continuous optimization methods like neural networks, simulated annealing [79], genetic algorithms [28] and probabilistic relaxation [39] have been proposed. They find solutions in a reasonable time without imposing constraints on the topology of the graph; the drawback is that their solutions are approximate.

### 2.1.2. The VF2 Algorithm

More recently, new mechanisms to query large graphs have been devised. Specifically, Cordella et al. proposed a deterministic algorithm, called VF2, that does not impose any restrictions on the topology of the graph and supports attributed graphs [44]. This algorithm achieves a reduced computational complexity by using a set of feasibility rules during the matching process. Furthermore, sophisticated data structures are used to reduce space complexity. Experimental results prove that VF2 is a competitive graph matching algorithm.

In particular, VF2 determines if graphs $G_1 = (V_1, E_1)$ and $G_2 = (V_2, E_2)$ are isomorphic by constructing a mapping set $M \subset V_1 \times V_2$, where each pair $(v_1, v_2)$ represents the mapping of a node $v_1$ in $G_1$ to a node $v_2$ in $G_2$. Then, if $G_1$ and $G_2$ are isomorphic, the mapping set $M$ is a bijection that preserves the adjacency relation of the graphs. The process to find $M$ can be described using a State Space Representation (SSR) [117] where each state is a partial mapping. Specifically, a partial mapping $M(s)$ leads to a new state $M(s')$ by inserting a new pair $(v_1, v_2) \in V_1 \times V_2$ that maintains consistency, i.e., that does not preclude the possibility of obtaining a complete mapping. A partial mapping is consistent if the subgraphs that its nodes constitute in $G_1$ and $G_2$ are isomorphic.
Let $M_1(s)$ and $M_2(s)$ be the set of nodes of $V_1$ and $V_2$ in the mapping $M(s)$, respectively. Also, let $T_1^{in}(s)$ be the set of nodes in $V_1$ that are not in $M_1(s)$ but are the origin of edges ending into nodes in $V_1$. Likewise, let $T_1^{out}(s)$ be the set of nodes in $V_1$ that are not in $M_1(s)$ but are the destination of edges starting from nodes in $V_1$. Furthermore, let $R_1(s)$ denote the nodes in $V_1$ that are neither in $M_1(s)$, $T_1^{in}(s)$ or $T_1^{out}(s)$. Sets $T_2^{in}(s)$, $T_2^{out}(s)$ and $R_2(s)$ are defined in a similar way. Then, consistency maintenance is verified by five feasibility rules:

- For each predecessor $v'_1$ of $v_1$ in $M_1(s)$, there must be a predecessor $v'_2$ of $v_2$ in $M_2(s)$ such that $(v'_1, v'_2) \in M(s)$, and vice versa.

- For each successor $v'_1$ of $v_1$ in $M_1(s)$, there must be a successor $v'_2$ of $v_2$ in $M_2(s)$ such that $(v'_1, v'_2) \in M(s)$, and vice versa.

- The number of successors of $v_1$ that are in $T_1^{in}(s)$ must be equal to the number of successors of $v_2$ that are in $T_2^{in}(s)$. Similarly, the number of predecessors of $v_1$ that are in $T_1^{in}(s)$ must be equal to the number of predecessors of $v_2$ that are in $T_2^{in}(s)$.

- The number of successors of $v_1$ that are in $T_1^{out}(s)$ must be equal to the number of successors of $v_2$ that are in $T_2^{out}(s)$. Similarly, the number of predecessors of $v_1$ that are in $T_1^{out}(s)$ must be equal to the number of predecessors of $v_2$ that are in $T_2^{out}(s)$.

- The number of successors of $v_1$ that are in $R_1(s)$ must be equal to the number of successors of $v_2$ that are in $R_2(s)$. Likewise, the number of predecessors of $v_1$ that are in $R_1(s)$ must be equal to the number of predecessors of $v_2$ that are in $R_2(s)$.

The last three rules should be adapted for subgraph isomorphism. In particular, the equal constraints must be replaced for inequalities. In case of isomorphism in attributed graphs, semantic feasibility is also evaluated. Specifically, when a new pair $(v_1, v_2) \in V_1 \times V_2$ is considered to extend the partial mapping $M(s)$, it is verified that $v_1$ and $v_2$ are semantically compatible. Furthermore, for each edge $(v_1, v'_1)$ where $v'_1 \in M_1(s)$, there must be a node $v'_2 \in M_2(s)$ such that $(v_2, v'_2)$ is semantically compatible with $(v_1, v'_1)$. Similarly, for each edge $(v'_1, v_1)$ where $v'_1 \in M_1(s)$, there must be a node $v'_2 \in M_2(s)$ such that $(v'_2, v_2)$ is semantically compatible with $(v'_1, v_1)$.

From a global perspective, the algorithm works as follows. For a given state $M(s)$, a recursive procedure called MATCHVF2(), is performed. In this procedure, all the pairs $(v_1, v_2) \in V_1 \times V_2$ that can be considered for inclusion in $M(s)$ are inserted into the set $P(s)$. For each pair $p \in P(s)$ that satisfies the feasibility and semantic rules, a new state $M(s')$ is generated by inserting $(v_1, v_2)$ into $M(s)$. Then, the procedure MATCHVF2() is recursively called for each $M(s')$. The process ends when a complete mapping is found or when all the feasible mappings are explored. Note that VF2 explores the search space in a DFS fashion starting from an empty mapping set.
In order to reduce space requirements, and then make it scalable to large graphs, VF2 makes use of the following data structures:

- **Vectors** $\text{core}_1$ and $\text{core}_2$, whose length is $n$ and $m$, respectively. Specifically, $\text{core}_1[i]$ contains the index of the current mapping of node $v_i \in M_1(s)$ into $G_2$; in case $v_i$ is not in the mapping yet, then $\text{core}_1[i] = \text{null}$. Vector $\text{core}_2$ is defined in the same manner to establish the mappings of $M_2(s)$ into $G_1$.

- **Vectors** $\text{in}_1$ and $\text{in}_2$, whose length is $n$ and $m$, respectively. In particular, $\text{in}_1[i]$ is non-$\text{null}$ if $v_1$ is either in $M_1(s)$ or $T_1^{in}(s)$; the value stored in $\text{in}_1[i]$ is the depth in the SSR tree of the state in which the node entered the corresponding set. Vector $\text{in}_2$ is defined in the same manner with respect to $G_2$.

- **Vectors** $\text{out}_1$ and $\text{out}_2$, whose length is $n$ and $m$, respectively. They are defined in a similar way as $\text{in}_1$ and $\text{in}_2$. For instance, $\text{out}_1[i]$ contains the depth in the SSR tree in which $v_i$ entered either $M_1(s)$ or $T_1^{out}(s)$. In case $v_i$ has been inserted in neither of these sets, $\text{out}_1[i]$ is equal to $\text{null}$.

These vectors allow to perform different operations of the algorithm efficiently. For instance, membership queries can be evaluated in constant time. Furthermore, VF2 does not require to store a copy of these vectors at each state of the search. Just one instance of each vector must be stored because if an element is non-$\text{null}$ in a given state, it will have the same value in the descending states from $s$. This, along with the traversal order of DFS, allows to restore the previous values of the vectors when the algorithm backtracks. Then, the space complexity of VF2 is $\Theta(n)$, i.e., the size of the vectors. In the best case, the algorithm finds a complete mapping on the first attempt, i.e., just one length-$n$ path in the SSR is explored. Then, the best-case time complexity of VF2 is $\Theta(n^2)$. On the other hand, the worst-case time complexity of VF2 is $\Theta(n!n)$ given that the algorithm visits $n!$ states in the worst case, i.e., the case where the complete mapping is the last one explored or when there is no complete mapping at all.

The VF2 algorithm was experimentally compared against Ullmann’s algorithm and NAUTY to evaluate its performance on graph isomorphism. In 56% of the cases, the best results were achieved by VF2 while, in 44% of the cases, NAUTY was the most efficient. Moreover, the results indicate that NAUTY is more convenient for randomly connected graphs, while VF2 is more efficient for graphs with a more regular structure, especially for large graphs. It is important to remark that, in most applications, graphs follows certain regularity. This makes VF2 a good candidate to consider in practical environments. Furthermore, VF2 was compared against a straightforward adaptation of Ullmann’s algorithm that supports attributes to evaluate subgraph isomorphism. Results show that VF2 performs significantly better. While the matching time rapidly grows with the size of the subgraph for Ullmann’s algorithm, it remains almost independent of such size for VF2 [44].
2.2. Queries on Attributed Graphs

Several techniques have been proposed in the literature for querying graphs. However, the existing graph querying methods mainly focus on querying the topological structure of the graphs \[122, 156, 159\] and very few of them have considered the use of attributed graphs \[127, 143\]. In practice, it is more common that the querying requirements for the applications of graph databases (e.g., social networks or bibliographical networks) would involve querying the graph data (attributes on nodes/edges) in addition to the graph topology.

Answering queries that involve predicates on the attributes of the graphs (nodes or edges) is more challenging as it requires extra memory consumption for building indices over the graph attributes in addition to the structural indices in order to accelerate the query evaluation process. Furthermore, it makes the query evaluation and optimization process more complex (e.g., evaluation and join orders). Existing graph optimization techniques focus on either building index structures \[156, 159\] or on developing estimation modules for certain graph queries \[120, 158\]. These techniques are complementary to this work. They can be used to accelerate query processing in our approach, which exploits both attribute histograms and the topology of the graph to produce efficient linearizations that yield lower matching time. At the same time, these index structures cannot be used to answer all queries as they may require graph exploration. Two types of query which are widely used in many applications are reachability queries \[40, 151, 84\], and pattern match queries \[159, 161\]. In this section, we review the related work on reachability queries (Section 2.2.1), pattern match queries (Section 2.2.2), and a recently defined type of queries called pattern queries (Section 2.2.3).

2.2.1. Reachability Queries

Given a directed attributed graph \(G = (V, E)\) and two nodes \(u, v \in V\), a reachability query determines whether there exists a directed path in \(G\) from \(u\) to \(v\). Graph reachability is closely related to the concept of transitive closure. Specifically, the transitive closure of \(G\), denoted as \(TC(G)\), is defined as the set of the pairs of nodes \((u, v)\) such that \(v\) is reachable from \(u\). Then, in order to determine reachability from \(u\) to \(v\), it suffices to evaluate the membership of \((u, v)\) in \(TC(G)\). However, the transitive closure of large and dense graphs can be very large.

A reachability query on a directed graph \(G\) can be evaluated by using an associated directed acyclic graph (DAG) \(G' = (V', E')\) of \(G\). Such graph can be obtained by finding the strongly connected components of \(G\), which takes \(O(|V| + |E|)\). Each node in \(V'\) represents one of the components. Each edge \((u, v) \in E\), such that \(u\) and \(v\) belong to different components, is associated to an edge in \(E'\) between the nodes in \(V'\) that represent the corresponding components of \(u\) and \(v\). Then, given the nodes \(u, v \in V\), we can say that \(v\) is reachable from \(u\) if they are in the same component or if the component of \(v\) is reachable from the component of \(u\). In the next algorithms, we assume that
the directed graph has been transformed into a DAG.

There are two main approaches to evaluate a reachability query on a graph \( G = (V, E) \): (i) performing a DFS/BFS traversal; and (ii) pre-computing and maintaining the transitive closure of the graph. The former has high time complexity (i.e., \( O(|V| + |E|) \)). The latter allows to answer reachability queries in constant time but incurs high space requirements (i.e., \( O(|V|^2) \)). Existing solutions attempt to obtain a convenient trade-off between time and space requirements where the query time complexity varies between \( O(1) \) and \( O(|V| + |E|) \) (the complexity of approaches (i) and (ii)). This trade-off is achieved by constructing different types of indices that reduce the space storage of the transitive closure [157]. Table 2-1 summarizes the complexity of constructing indices under different approaches, their size and the corresponding query response time [84].

Table 2-1.: Complexity of solutions for reachability queries on a DAG \( G = (V, E) \) [84]. Notation: \( t \) is the number of non-tree edges in the spanning-tree-based solutions.

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>Query Time</th>
<th>Index Size</th>
<th>Index Construction</th>
</tr>
</thead>
<tbody>
<tr>
<td>DFS/BFS</td>
<td>( O(</td>
<td>V</td>
<td>+</td>
</tr>
<tr>
<td>Transitive Closure [136]</td>
<td>( O(1) )</td>
<td>( O(</td>
<td>V</td>
</tr>
<tr>
<td>Tree Cover [1]</td>
<td>( O(\log</td>
<td>V</td>
<td>) )</td>
</tr>
<tr>
<td>Labelling + SSPI [35]</td>
<td>( O(</td>
<td>E</td>
<td>-</td>
</tr>
<tr>
<td>GRIPP [145]</td>
<td>( O(</td>
<td>E</td>
<td>-</td>
</tr>
<tr>
<td>Dual-Labelling [151]</td>
<td>( O(1) )</td>
<td>( O(</td>
<td>V</td>
</tr>
<tr>
<td>Chain Cover [36]</td>
<td>( O(\log k) )</td>
<td>( O(k</td>
<td>V</td>
</tr>
<tr>
<td>Path-Tree Cover [85]</td>
<td>( O(\log^2 k') )</td>
<td>( O(k'</td>
<td>V</td>
</tr>
<tr>
<td>2–Hop Cover [40]</td>
<td>( O(\sqrt{</td>
<td>E</td>
<td>}) )</td>
</tr>
<tr>
<td>3–Hop Cover [84]</td>
<td>( O(\log</td>
<td>V</td>
<td>+ k) )</td>
</tr>
</tbody>
</table>

In the following, we describe selected solutions of Table 2-1 as presented in a recent survey [157]. In the late eighties, an algorithm that computes the transitive closure without any compression in \( O(|V| \times |E|) \) time was developed by Simon [136]. In particular, this algorithm is an improvement on the solution proposed by Goralčíková and Koubek [72] a decade earlier.

Also, in the late eighties, Agrawal et al. propose the use of a spanning tree of the DAG to compress its transitive closure; such spanning tree is called the tree cover [1]. In this approach, the edges of the graph are classified based on their membership in the tree: if an edge appears in the tree, it is called a tree edge; otherwise, it is called a non-tree edge. First, let us consider the case where the tree cover is comprised of only tree edges, i.e. the graph is a tree. Each node \( u \) is associated to an interval-based label \([i, j]\), where \( i \) is the index and \( j \) is the postorder number of the node. The
postorder number of \( u \) is its relative position in the postorder traversal of the tree. The index of \( u \) is the lowest postorder number among its descendants or, in case it is a leaf, its own postorder number. Using these intervals, reachability can be checked with the following lemma: a node \( v \), with postorder number \( k \), is reachable from \( u \) iff \( i \leq k < j \), i.e. the interval of \( v \) is contained in the interval of \( u \) [1].

This index is applied to general graphs as follows. A spanning tree of the graph is computed. In case the graph contains more than one connected component, a virtual root node can be added. Then, each node is associated to a set of intervals rather than a single interval. The set of each node initially contains only the interval \([i, j]\) with its index and postorder number, respectively, just like in the case of trees. Then, the nodes are sorted in reverse topological order. For each node \( u \), all its outgoing edges are considered. Specifically, for each edge \((u, v)\), all the intervals associated to the node \( v \) are inserted into the set of intervals of node \( u \). If an interval is subsumed by another interval in the set, it is discarded. It can be concluded that a node \( u' \) can reach node \( v' \) if the postorder number of \( v' \) is contained in one of the intervals of node \( u' \). An algorithm that computes an optimal spanning tree, in the sense that it minimizes the required storage of the transitive closure, was also presented [1].

More recently, different variants of this approach, based on a spanning tree, have been developed. For instance, Chen et al. proposed a set of stack-based algorithms in 2005 [35]. In their approach, a label is assigned to each node in the tree; we denote the label of node \( u \) as \( \text{label}(u) \). A predicate \( P(\text{label}(u), \text{label}(v)) \) is used to determine graph reachability from node \( u \) to node \( v \) in the DAG. Specifically, if \( P(\text{label}(u), \text{label}(v)) = \text{true} \), then \( v \) is reachable from \( u \). However, because not all the edges appear in the tree, \( P(\text{label}(u), \text{label}(v)) = \text{false} \) does not imply that \( v \) is not reachable from \( u \). In order to tackle these cases, another data structure called \textit{Surrogate & Surplus Predecessor Index} (SSPI) is used [157]. Another solution that makes use of a spanning tree of the DAG and labelling was devised by Trißl and Leser [145]. This approach, called \textit{GRaph Indexing based on Pre- and Postorder numbering} (GRIPP), extends the pre- and postorder numbering scheme to support graphs. The search phase is performed by means of a hop technique and a set of pruning strategies. Notice that the complexity of this solution is the same as the one of Chen et al.

Also following the spanning-tree based approach, Wang et al. proposed a solution suitable for large sparse graph where graph reachability needs to be evaluated [151]. This approach also constructs a spanning tree where each node \( u \) is associated to an interval-based label \([\text{start}, \text{end}]\), where \( \text{start} \) and \( \text{end} - 1 \) are the preorder and postorder number of \( u \) in the spanning tree. This labelling scheme is called \textit{Dual-I}. Furthermore, the non-tree edges are stored in a \textit{link table}; such table is desired to be small. The selection of a convenient spanning tree and strategies to avoid superfluous non-tree edges have been studied [151]. This solution achieves constant query time while reduces the size of the index to \( O(|V| + t^2) \), where \( t \) is the number of non-tree edges. Notice that this is efficient only if \( t \) is much lower than \( |V| \), which does not occur in many real graphs [84].
Most of the algorithms that efficiently solve shortest distance queries [40, 32, 37, 154], and reachability queries for graphs do not support predicates on the connecting paths [40, 32, 151, 37, 154]. Because of the need for supporting semantic restrictions in the queries, without sacrificing the response time, recent work has been developed in this direction. For instance, a special type of reachability query, called label-constraint reachability query, that only accepts edge labels from a given set was proposed [83]. More recently, a revised definition of reachability query, where a regular expression of edge labels is used to specify the connecting path in the query, was proposed [59]; however such regular expression has limited expressiveness: neither the Kleene operator nor predicates on intermediate nodes are supported.

### 2.2.2. Pattern Match Queries

Pattern match queries have been defined and solved in terms of subgraph isomorphism due to its appropriateness for practical applications; some of the most relevant solutions following this approach are presented in [29, 35, 143, 38, 160, 159, 161], as outlined in the surveys [69, 132]. In order to make the pattern match queries more flexible and support more applications, the edges of the query have been allowed to map to paths in the graph [58, 61, 161]. The main challenge of the techniques based on subgraph isomorphism is scalability on the graph size given that the problem is NP-Complete. To address this issue, a new version of pattern match query was proposed [59].

On the other hand, given that graphs can be considered as databases, the database community introduced a formal language called GraphQL to query semantic graphs [78]. Moreover, different techniques are applied on GraphQL to support large graphs: use of neighborhood subgraphs, joint reduction of the search space and optimization of the search order. Experimental results on a biological database of thousands of nodes showed that GraphQL outperforms an SQL-based implementation [78]. Later, Zhao and Han proposed an indexing mechanism called SPath that leverages decomposed shortest paths around node neighborhoods as basic indexing units [159]. This mechanism achieves effectiveness in pruning the search space and scalability in index construction and deployment. Two experimental tests were performed: one on the same biological database used for testing GraphQL, and another one on a synthetic database that contains one million nodes; in both tests SPath significantly outperformed GraphQL [159].

### 2.2.3. Pattern Queries

To address this issue, a new version of pattern match query was proposed [59]. This variant is based on graph bounded simulation rather than subgraph isomorphism. Graph bounded simulation [60] is an extension of graph simulation for pattern match queries where bounds on the number of permitted hops are imposed. This restriction allows to solve pattern match queries in polynomial time, even when we allow that edges in the query map to paths in the graph. The corresponding
output presents in batch the global matches for each edge (reachability query) in the query.

However, like in the reachability queries also studied in [59], pattern match queries using the Kleene operator or containing predicates on intermediate nodes cannot be solved with such approach. In this paper, we introduce generalized pattern queries as a new type of queries that evaluates both attribute predicates and structural requirements. These queries, besides allowing edge-to-path mappings, also support predicates on intermediate nodes/edges and operators like union and the Kleene star in the reachability requirements. Moreover, the output produced is easy to interpret: it consists of the set of all the solution instances where each instance is an ordered set of nodes that correspondingly satisfy the nodes in the query; thus, the relative relations among the output nodes from each solution instance and the query is straightforward to determine.

On the other hand, graph query languages, based on either regular expressions [42, 51, 73], SQL-like languages [10, 123, 134], or procedural languages [78], to solve graph queries have been proposed. Such languages have limited expressive power and lack the support of declarative query interfaces. For instance, comparing with well-known query languages such as SPARQL and SQL, we support queries that cannot be expressed by either of them. (1) The SPARQL query language expresses pattern match queries over RDF data, and cannot express general reachability queries. In contrast, we target both reachability queries and pattern match queries. A recent specification of SPARQL allows a limited form of reachability with the triple pattern (subject, verb+, object). (2) SQL cannot express paths of arbitrary length, unless it is extended with recursion [69] to support closure operators. Notwithstanding, we use an idea that is similar to the query optimizer in a database system: the most restrictive conditions are often pushed down to the evaluation tree of a query plan; thus, the selective predicates are evaluated early in the execution.

2.3. Parameterized Matching

As an aid in software maintenance, parameterized matching was first defined by Brenda Baker to detect duplicate code in large software systems [15]. Later, the study on this problem was further extended due to its practical applications in different areas. In this section, we cover some of this research. Specifically, in Section 2.3.1, the formal definition of parameterized matching and its variants are given. Some parameterized matching algorithms are reviewed in Section 2.3.2 and some extensions are presented in Section 2.3.3. Finally, some of the most important applications of this pattern matching variant are shown in Section 2.3.4.

2.3.1. Definition of the Basic Problems

Let $\Sigma_C$ be the constant symbol alphabet and $\Sigma_P$ be the parameter symbol alphabet. We assume that $\Sigma_C$ and $\Sigma_P$ are disjoint from each other and the set of nonnegative integers. A parameterized string or a $p$–string is defined as a string of symbols in $(\Sigma_C \cup \Sigma_P)^*$. Furthermore, two length-$m$
2.3 Parameterized Matching

Parameterized Matching is the problem of finding all the parameterized–matches of a pattern in a text. More formally, let us consider two p–strings: the pattern \( P = P_{1...m} \) and the text \( T = T_{1...n} \), both defined over \( \Sigma_C \cup \Sigma_P \). Also, let \( T^i \) denote the length–\( m \) text window starting at position \( i \) of \( T \), i.e., \( T^i = T_{i...i+m-1} \). Then, pattern \( P \) is said to parameterized-match \( T^i \) iff there exists a bijective mapping function \( g_i \) such that \( g_i(P_j) = T_{i+j-1} \), \( 1 \leq j \leq m \), so that \( g_i \) is identity for the symbols from \( \Sigma_C \). Notice that if there exists a function \( g_i(P_j) = T_{i+j-1} \), \( 1 \leq j \leq m \), the inverse of \( g_i \) also exists given that \( g_i \) is bijective. So we can equivalently say that \( P \) parameterized-matches \( T^i \) if there exists a bijective mapping function \( g_i'(T_{i+j-1}) = P_j \) so that \( g_i' \) is identity for the symbols from \( \Sigma_C \). Note that, at each position \( i \) of \( T \), a different \( g_i \) can be considered to determine the existence of a parameterized-match between the pattern and the text window starting at position \( i \). The output of the problem is the set of indices \( i \), \( 1 \leq i \leq n-m+1 \), such that \( P \) parameterized-matches \( T^i \). This problem is also referred as Parameterized Fixed Pattern Matching (PFPM) [81].

Some other problems related to parameterized matching have been defined to be able to support more applications. One of them is finding the maximal \( p \)–matches over a threshold length of a \( p \)–string text, defined as follows. Let \( T = T_{1...n} \) be a \( p \)–string and \( T_{i...i+k} \) and \( T_{j...j+k} \) two \( p \)–substrings of it that \( p \)–match. This \( p \)–match is said to be left–extensible if \( T_{i-1...i+k} \) and \( T_{j-1...j+k} \) are a \( p \)–match and is right–extensible if \( T_{i...i+k+1} \) and \( T_{j...j+k+1} \) are a \( p \)–match, where \( 1 \leq i \leq i + k \leq n \), \( 1 \leq j \leq j + k \leq n \) and \( i \neq j \). If a \( p \)–match is neither left–extensible or right–extensible, it is said to be a maximal \( p \)–match. Maximal \( p \)–matches are not an equivalence relation, because they are not transitive, so the output of the maximal \( p \)–matches problem must list pairs of \( p \)–substrings rather than an equivalence class. Thus, the output of the maximal \( p \)–matches of a \( p \)–string text \( T = T_{1...n} \) over a threshold length \( t \) problem must report the set of all pairs of \( p \)–substrings of \( T \) that \( p \)–match and whose length is at least \( t \).

On the other hand, the searching of multiple patterns has been extended to parameterized matching [81]. For a given fixed set \( D \) of \( p \)–string patterns over \( \Sigma_C \cup \Sigma_P \), the Parameterized Multiple Pattern Matching (PMPM) problem consists of preprocessing \( D \) as an aid to later determine the \( p \)–matches (for all of the patterns in \( D \)) in a query text \( T \). A dynamic variant of this problem, called Parameterized Dynamic Dictionary Matching (PDDM), has also been considered [81]. In this problem, a dictionary \( D \) of \( p \)–string patterns is preprocessed and maintained with available operations of inserting/deleting patterns into/from \( D \) and searching a query text \( T \) for \( p \)–matches for the patterns currently in \( D \).
2.3.2. Solutions

The maximal parameterized matching over a threshold length problem was the first parameterized matching problem to ever be considered, even before some of the basic definitions of parameterized matching were formalized. Baker tackled this problem motivated by the observation that there was a considerable amount of duplicate code in large software systems [13]. Therefore, she presented a program, called DUP, as an aid to find all the duplicate sections of code with a minimum length, specified by the user, in a large software system. DUP simplifies the problem to an exact matching problem replacing all the parameters by a determined symbol and then looks for the p–matches among the exact matches found. The algorithm is based on recursions over the suffix tree of the text.

A suffix tree of a string $X$ is a compacted trie\(^1\) defined on the set of the suffixes of $X\$$, where $\$$ is a unique end marker so that no suffix is a prefix of another suffix (cf. [101, 147, 155]). A compacted trie is a tree data structure defined on a set of strings (in this case, the set of the suffixes of a string) such that: (i) every inner node has at least two children; (ii) every edge is labelled with a substring of one of the strings in the given set; and (iii) the concatenation of the labels on the path from the root to each leaf is a distinct string in the set [41]. The key property of compacted tries is that, for every pair of leaves, the string formed by concatenating the labels from the root to their lowest common ancestor is the longest common prefix of the strings in the set associated with these leaves. For each node $v$ of the trie, the pathstring of $v$ is the concatenation of the edge labels on the path from the root to $v$ and the length of the pathstring of $v$ is called the pathlength of $v$.

For constructing the suffix tree for the text, Baker suggests McCreight’s Algorithm [101]. This algorithm builds a suffix tree for a string $X = X_1...m$ in $m$ stages each one of which corresponds to the insertion (from left to right) of a suffix $X_i...m$, $1 \leq i \leq m$, of $X$. The insertion of the $i$-th suffix is made so that the first part of the path coincides with the path of the longest common prefix of $X_i...m$ and $X_j...m$ for some $j < i$ (a previously inserted suffix). One of the key points to make this algorithm efficient is the use of suffix links. If an internal node has pathstring $aX$, where $a$ is a symbol and $X$ is a string, its suffix link points to an internal node with pathstring $X$ (which is guaranteed to exist due to the Common Prefix Property and the Distinct Right Context Property of strings). Suffix links are also useful for pattern matching in space proportional to the size of the pattern [34].

Experiments with real data proved that DUP is highly useful in software maintenance but also showed that the algorithm is inefficient given that just a few of the found exact matches correspond to p–matches. For this reason, the same author proposed a more elaborate theory [15, 18] aiming to find better solutions and support a wider range of applications. This theory includes the definition of the parameterized pattern matching problem.

\(^1\)a.k.a. Multiway Patricia Trie.
Some core aspects of parameterized matching are discussed, as follows. For the case of the string comparison problem, a naive way to determine whether two length\( m \) \( p \)-strings \( X = X_1 \ldots m \) and \( Y = Y_1 \ldots m \) are a \( p \)-match was proposed [18]. It consists of the following steps. Traverse both \( X \) and \( Y \) from left to right while constructing a table that establishes the mapping function that allows to transform one of the \( p \)-strings into the other one. Continue with this procedure until a mismatch is found. A mismatch between two corresponding symbols occurs in any of the following three cases: (i) one symbol is a parameter (from \( \Sigma_P \)) and the other is a non-parameter (from \( \Sigma_C \)); (ii) both symbols are non-parameters and they are different; and (iii) both symbols are parameters but any of them has previously been assigned to a different parameter in the mapping table. If no mismatch occurs, then \( X \) and \( Y \) are a \( p \)-match. The time complexity is \( O(m) \) and the space complexity is \( O(|\Sigma_P|) \). Nevertheless, this approach is not proper for the pattern matching problem.

A procedure called \( \text{prev} \) was defined to yield more efficient solutions for parameterized matching [18]. Given the constant alphabet \( \Sigma_C \), the parameter alphabet \( \Sigma_P \) and a length\( m \) \( p \)-string \( X = X_1 \ldots m \), \( \text{prev}(X) \) is a string in \( (\Sigma_C \cup \mathbb{N})^* \) where every constant symbol in \( X \) remains the same in \( \text{prev}(X) \) but the parameter symbols are replaced by nonnegative integers: the leftmost occurrence of a determined parameter is represented by a 0 and the other occurrences are represented by the difference in position compared to the previous occurrence of this parameter. The numbers that represent difference in position are called parameter pointers. The time complexity of the computation of \( \text{prev} \) is \( O(m) \) and the space complexity is \( O(|\Sigma_P|) \) by means of a table containing the last occurrence position of each parameter. Notice that \( \text{prev}(X) \) is calculated in such a way that it does not matter what the parameters of \( X \) are; what is really relevant is the relative distance among the different occurrences of the same parameter (represented by the parameter pointers) which provides valuable information about the structure of the \( p \)-string. Thus, two \( p \)-strings \( X \) and \( Y \) are a \( p \)-match, iff \( \text{prev}(X) = \text{prev}(Y) \).

**Example.** For the example presented in Figure 1-5, where \( \Sigma_C = \{b\} \), \( \Sigma_P = \{x, y, z\} \), \( X = xbyyxbx \) and \( Y = zbxzbz \), we find that \( \text{prev}(X) = 0b014b2 = \text{prev}(Y) \) and therefore \( X \) and \( Y \) are a \( p \)-match (see Figure 2-1).

The \( \text{prev} \) of any substring of a \( p \)-string \( X \) can be calculated from \( \text{prev}(X) \) given that any symbol of the substring is the same as in \( \text{prev}(X) \) except when it is a parameter pointer that points to a position before \( i \); in such case, it will correspond to the first occurrence of the parameter in the substring so it must replaced by a 0. On the other hand, the parameterized pattern matching problem could be defined, in terms of \( \text{prev} \), in the following manner: Given the pattern \( P = P_1 \ldots m \) and the text \( T = T_1 \ldots n \), both defined over \( \Sigma_C \cup \Sigma_P \), \( P \) is said to parameterized-match \( T^i \) iff \( \text{prev}(P) = \text{prev}(T^i) \) (recall that \( T^i = T_{i \ldots i+m-1} \)). Thus, using \( \text{prev} \) is a convenient approach for the parameterized pattern matching case, given that any \( \text{prev}(T^i) \) can be calculated as follows.
Figure 2-1.: Determination of a p–match between $X = xbyyx$ and $Y = zbxxz$ through the prev procedure, where $\Sigma_C = \{b\}, \Sigma_P = \{x, y\}$.

\[
prev(T^i)_j = \begin{cases} 
0 & \text{if } prev(T)_{i+j-1} > j - 1 \\
prev(T)_{i+j-1} & \text{otherwise}
\end{cases}
, \quad \text{for } 1 \leq j \leq m.
\]

In this sense, with the use of prev, parameterized matching can be seen as a standard exact matching problem without losing information about the chains of parameters. Reminiscing about the use of suffix trees for exact matches in DUP, Baker defined a new data structure called parameterized suffix tree to aid in directly searching for parameterized–matches [18]. Parameterized suffix trees are a generalization of suffix trees for strings.

To generalize suffix trees to parameterized suffix trees, it is necessary to review the definition of p–suffix [18]. The $i$–th p–suffix of a p–string $X = X_1...m$ is defined as $psuffix(X, i) = prev(X_{i..m})$.

So we can calculate each p–suffix, just like the prev of any substring of $X$, by copying the corresponding symbols of prev($X$) except when they are parameter pointers that point to a symbol outside the substring (in which case they are replaced by 0). Then, p–suffix trees are defined as follows. If $X$ is a p–string that ends with a unique end marker $\$ in $\Sigma_C$, a parameterized suffix tree, also called p–suffix tree, for $X$ is a compacted trie (multiway Patricia trie) that stores the p–suffixes of $X$ [18]. Following, we give an example, as it appears in [18], of the p–suffixes that the p–suffix tree of a given string must store.

**Example.** Let $\Sigma_C = \{b, \$\}$ be the constant alphabet, $\Sigma_P = \{x, y\}$ be the parameter alphabet and $X = xbyyx$ be a p–string. Then, prev($X$) = 06014b2 so the p–suffix tree of $X$ must encode 06014b2$, b010b2$, 010b2$, 00b2$, 0b2$, b0$, 0$ and $\$ (see Fig. 2-2).
2.3 Parameterized Matching

Figure 2-2.: A p–suffix tree for $X = xbyyxbx\$ where $\Sigma_C = \{b, \$\}$ and $\Sigma_P = \{x, y\}$.

An algorithm to construct p–suffix trees, called LAZY, was proposed [18]. It is based on McCreight’s algorithm for constructing suffix trees [101]. Nevertheless, in this case, a suffix link for a node with pathstring $aX$ cannot point to a node with pathstring $X$ because that node may not exist. This is because the Distinct Right Context Property does not hold for p–strings. Therefore, suffix links were redefined in such a way that, for a node with pathstring $aX$, the suffix link points to the node whose pathstring is the longest prefix of $X$ among all the nodes in the tree. This algorithm is linear in the p–string length in both time and space for fixed alphabets. For variable alphabets, the time complexity is $O(n(\Sigma_P|\log(\Sigma_C| + |\Sigma_P|)))$.

Later, Baker proposed a new algorithm to build p–suffix trees, called EAGER, where the suffix links for a node with pathstring $aX$ point to the node whose pathstring is the shortest of all those for which $X$ is a prefix [15]. This idea is more convenient for the structure of p–suffix trees. Even though for fixed alphabets the time and space complexity remain linear, for variable alphabets the time complexity is $O(n(|\Sigma_P| + \log(|\Sigma_C| + |\Sigma_P|)))$. Nevertheless, for both LAZY and EAGER, the time complexity of the variable alphabet case can be reduced to $O(n \log n)$ by using auxiliary data structures like concatenable queues [2] and Sleator-Tarjan dynamic trees [137]. However, the use of these structures makes the algorithms not practical.

Other authors have worked on developing faster algorithms for constructing p–suffix trees. Kosaraju proposed an algorithm whose time complexity is $O(n \log(|\Sigma_P| + |\Sigma_C|))$ [90]. Furthermore, a randomized algorithm to construct suffix trees for cases where there are missing suffix links, such as p–suffix trees and suffix trees for two–dimensional arrays, was proposed [41]. It was the first algorithm whose time complexity is $O(n)$ even for variable alphabets. It is based on adding
a back–propagation component to McCreight’s Algorithm and using a high probability hashing scheme for large degrees.

Two solutions for the parameterized matching problem that use p–suffix trees were developed [15]. Given the pattern p–string $P = P_1...m$ and the text p–string $T = T_1...n$, one of the algorithms consists of following the path determined by the symbols of $\text{prev}(P)$ on the p–suffix tree of $T$ to find out if $\text{prev}(P)$ is identical to a length–$m$ substring of $T$. For fixed alphabets, to determine all the positions in $T$ where there is a p–match with $P$ takes $O(m + \text{occ})$ time and $O(n)$ space, where occ is the number of p–matches. For variable alphabets, the time complexity is $O(m \log(|\Sigma_C| + |\Sigma_P|) + \text{occ})$. The other algorithm consists of searching in a p–suffix tree for $P$ through an adaptation of the corresponding algorithm for strings [33]. Its space complexity is $O(m)$ and its time complexity is $O(n)$ for fixed alphabets; for variable alphabets, its time complexity is $O(n |\Sigma_P| + \log(|\Sigma_C| + |\Sigma_P|))$. Nevertheless, it could also be improved to $O(n \log(|\Sigma_C| + |\Sigma_P|))$ by using some auxiliary data structures for computing lowest common ancestors [75, 130].

On the other hand, an algorithm, called PDUP, for finding the maximal p–matches over a threshold length of a text $T = T_1...n$ was devised [18]. PDUP is similar to DUP, but constructs a p–suffix tree of the text instead of a suffix tree. This algorithm generalizes to p–strings the algorithm for finding maximal p–matches over a threshold length in a string [14]. In this generalization, it is necessary to augment the p–suffix tree with lists that store valuable data that makes possible to determine whether there is left–extensibility in the p–matching substrings. The time complexity of PDUP is $O(n + \text{occ})$, where occ is the number of maximal p–matches found, even for variable alphabets.

Soon after Baker proposed the parameterized matching theory and its first algorithms, other researchers started to work on this topic. For instance, Amir et al. analysed Baker’s theory and defined a related model called Mapped Matching which is a special case of parameterized matching where all symbols are in the parameter alphabet $\Sigma_P$ [5]. Through this model, an algorithm that extends the KMP algorithm [89] to parameterized matching and runs in $O(n \log \min(m, |\Sigma_P|))$ time was proposed [5]. This was the first parameterized matching algorithm independent from the size of the constant alphabet $\Sigma_C$. Furthermore, it was proven that the $\log \min(m, |\Sigma_P|)$ factor is inherent to any algorithm for parameterized matching in the comparison model and, consequently, that the provided algorithm is optimal. This demonstration was achieved through a reduction from the element distinctness problem to parameterized matching.

This new research may have motivated Baker to look for parameterized matching solutions based on classical exact string matching algorithms [16]. Given that the BOYER–MOORE algorithm [26] is one of the most efficient, she attempted to generalize it to p–strings but found its worst case performance was poor. Therefore she turned to one of its variants, TURBOBM [49]. Her non-trivial generalization of TURBOBM to p–strings, called PTURBOBM, runs in $O(n \log \min(m, |\Sigma_P|))$ time and $O(n)$ space; the preprocessing time is $O(m \log \min(m, p))$. Its time complexity is the sa-
me as the generalization of KMP complexity so it is optimal [5]. Nevertheless some experiments show that PTURBOBM works better for long patterns over different alphabet sizes. Anyhow, for variable alphabets, both of these algorithms are notably better than then p–suffix tree based parameterized matching algorithms.

Other important contributions were made by Idury and Schäffer who proposed some variants of the basic problem (see Section 2.3.1) and solutions for all of them [81]. For the Parameterized Multiple Pattern Matching Problem, they proposed an algorithm that uses a modified Aho–Corasick automaton and runs in $O(n \log(|\Sigma_C| + |\Sigma_P|) + occ)$ time, where $occ$ is the number of occurrences of all the patterns. As for the Parameterized Dynamic Dictionary Problem, they devised an automaton algorithm that supports different operations with the following time complexity: (i) $O((n + occ)(\log(|\Sigma_C| + |\Sigma_P|) + \log d))$ for searching the p–string patterns of the dictionary in a p–string text $T = T_1...n$; (ii) $O(m(\log(|\Sigma_C| + |\Sigma_P|) + \log^2 d))$ for inserting a new pattern $P = P_1...m$ into the dictionary; and (iii) $O(m(\log(|\Sigma_C| + |\Sigma_P|) + \log d))$ for deleting a pattern $P = P_1...m$ from the dictionary, where $d$ is the total size of all the patterns.

More recently, Fredriksson and Mozgovoy proposed two new algorithms for both the single and multiple parameterized matching problems [68]. Both of them make use of Baker’s lemma to compute the $\text{prev}$ of a text substring through the $\text{prev}$ of the container p–string [18]. One of them is a bit–parallelism based algorithm called P–SHIFT–OR. It is a generalization of the SHIFT–OR algorithm [12] to p–strings and runs in $O(n \lceil m/w \rceil)$ worst case time and $O(n)$ average time. This algorithm can be extended to solve the multiple parameterized matching problem.

Fredriksson and Mozgovoy also devised an algorithm called Parameterized Backward Trie Matching (PBTM) [68] based on the Backward DAWG Matching (BDM) algorithm [25, 49]. First, the set of p–suffixes of the reverse of the pattern are stored in a trie. Then, this trie is used to fastly determine whether there is a p–match in the current text window; otherwise, the trie is used to calculate the length of the shift to consider the next text window where a p–match could be found. The average time complexity of PBTM is $O(n \log(m)/m)$. This process could also make use of a suffix array [99] instead of a trie, in which case the algorithm is called Parameterized Backward Array Matching (PBAM). PBTM and PBAM are also extensible for the multiple parameterized matching problem. It is remarkable that these algorithms are the first parameterized matching algorithms for which an average time complexity analysis has been made. They have optimal average case running for both single and multiple patterns, as confirmed by experimental results.

The diagram in Figure 2-3 shows the algorithms for solving the different parameterized matching problems presented in this section organized by the nature of their approaches.
2.3.3. Extensions

Parameterized Matching has been studied in many directions. For instance, an investigation about the periodicity of parameterized strings was done [9]. They attempted to generalize to p–strings two of the periodicity lemmas of strings: the Lyndon and Schitzenberger lemma (referred as Weak Version) [97], and the Fine and Wilf lemma [63]. They found out that only the Weak Version holds for p–strings only when the two mappings inducing the periodicity commute. These results and some other studies about the repetitions in p–strings showed considerable differences between p–strings and ordinary strings. Nevertheless, binary p–strings behave in a very similar way as ordinary strings with respect to periodicity and repetitions.

On the other hand, parameterized matching was extended to the two dimensional case by considering matrices of symbols instead of p–strings. Two–dimensional parameterized matching consists of finding all the p–matches of a pattern of size $m \times m$ in a text of size $n \times n$. An algorithm for the problem that runs in $O(n^2 + m^{2.5} \text{polylog } m)$ time was proposed [76]. Other solutions include a
2.3 Parameterized Matching

\(O(n^2 \log^2 m)\) deterministic algorithm and a \(O(n^2 \log n)\) randomized algorithm that reports all the p–matches [4]. Nevertheless, it may report a mismatch as match with probability of \(1/n^k\), where \(k\) is a given constant.

Other topic that arose as a matter of interest was the calculation of similarity between two p–strings. In particular, Baker defined the parameterized edit distance or p–edit distance of two p–strings as the cost of a minimal edit script, called p–edit script, that transforms one p–string into the other [19]. The valid operations are insertions, deletions and parameterized replacements (the replacement of a substring with a p–string that p–matches it). Moreover, Baker proposed an algorithm [19] for calculating the p–edit distance \(D\) of two prev–encoded p–strings, \(X = X_{1...m}\) and \(Y = Y_{1...n}\), by generalizing Myers’s algorithm for finding the LCS of two strings [116]. The algorithm runs in \(O(D(n + m))\) time and \(O(n + m)\) space. However, the complexities can be improved by using p–suffix trees [18] and the lowest common ancestor [75, 130]. Furthermore, a divide-and-conquer based algorithm for reporting the minimal p–edit script was proposed [19]. It also runs in \(O(D(n + m))\) and \(O(n + m)\) space. Finally, it is shown that these techniques can be extended to solve the approximate parameterized problem under the p–edit distance [19], defined as follows. For a given p–string pattern \(P = P_{1...m}\), a p–string text \(T = T_{1...n}\) and an integer \(k\), the goal is to report all the positions \(1 \leq i \leq n\) such that \(T^i\) are within the p–edit distance \(k\) of \(P\). This can be done in \(O((k + \log|\Sigma_C| + \log|\Sigma_P|)(n + m))\) time and \(O(n + m)\) space.

There have been some works about approximate parameterized problem under hamming distance. In particular, the \(\pi\)–match between two p–strings \(X = X_{1...m}\) and \(Y = Y_{1...m}\) was defined as the number of matches between \(\pi(Y_i)\) and \(X_i\), for \(1 \leq i \leq m\) [7]. For two equal–length p–strings, the approximate parameterized matching problem, also called parameterized matching with mismatches, consists of finding a \(\pi\) of maximal \(\pi\)–match. Given a p–string pattern \(P = P_{1...m}\) and a p–string text \(T = T_{1...n}\), the approximate parameterized searching problem under hamming distance consists of computing the approximate parameterized matching between \(P\) and every length–m p–substring of \(T\). It is not necessary to choose the same \(\pi\) for every text window, as in standard parameterized matching. Furthermore, a linear algorithm to solve this problem, for the case where both \(P\) and \(T\) are run–length encoded and one of them is a binary p–string, was devised [7].

Further studies about parameterized matching and hamming distance have been developed [77, 76]. Specifically, a related problem, called parameterized matching with a threshold of \(k\) mismatches, was proposed. Its goal is finding all the p–matches of a pattern \(P = P_{1...m}\) in a text \(T = T_{1...n}\) with at most \(k\) mismatches. For two equal–length p–strings \(X = X_{1...m}\) and \(Y = Y_{1...m}\), they proposed a \(O(m + k^{1.5})\) time algorithm and a \(O(m^{1.5})\) time algorithm for the cases when \(k\) is considered and when it is not considered, respectively. These solutions are based on maximum matching algorithms; furthermore, it was demonstrated that the maximum matching problem is reducible to the approximate parameterized matching problem. For a p–string pattern \(P = P_{1...m}\), a p–string text \(T = T_{1...n}\) and a given \(k\), a \(O(nk^{1.5} + mk \log m)\) time algorithm for the parameterized matching
with $k$ mismatches problem was also proposed. It is shown that this could be extended to the two dimensional case in $O(n^2 mk^{1.5} + m^2 k \log m)$ time.

Another approximate version of parameterized matching is based on $\delta$– and $\gamma$– distances. Specifically, we defined $\delta\gamma$–approximate parameterized matching [93, 103]. Given two equal-length integer strings $X = X_1...m$ and $Y = Y_1...n$, string $X$ is said to $\delta\gamma$–parameterized match string $Y$ if $X$ can be transformed in a string $X'$, via a bijection $\pi$ (i.e., $X_i' = \pi(X_i)$ for $1 \leq i \leq m$), such that $X' \delta\gamma$–matches $Y$. Constants $\delta$ and $\gamma$ are bounds for the local and global errors, respectively, on the difference between the corresponding symbols of the strings. A $O(nm)$ algorithm to report the $\delta\gamma$–parameterized matches of a pattern $P = P_1...m$ in a text $T = T_1...n$ was proposed [93, 103].

This variant is defined as a combination of two string matching paradigms: parameterized matching and $\delta\gamma$–matching. The latter, is very effective in searching for all similar but not necessarily identical occurrences of a given pattern. This problem has been well-studied (cf. [30, 50, 48]) due to its applications in bioinformatics [107, 111] and music information retrieval [30].

The parameterized matching problem under the LCS distance problem has also been considered. The longest common parameterized subsequence (LCPS) for two $p$–strings $X = X_1...m$ and $Y = Y_1...n$ was defined as the pair of sequences $I$ and $J$ of maximum length, such that $I$ is a subsequence of the $p$–string $X$, $J$ is a subsequence of the $p$–string $Y$, and $I$ and $J$ are a $p$–match [87]. It is important to remark that it is not required that the symbols in $I$ and $J$ are consecutive in $X$ and $Y$. The LCPS could be useful as a similarity measure between code sections; nevertheless, this problem has been proven to be NP–hard. Then, an approximate algorithm was proposed [87]. On the other hand, in [82] some algorithms for computing the longest parameterized common subsequences are presented; nevertheless it is important to mention that their definition of parameterization is considerably different from the one developed by Baker and tackled in this thesis.

Another parameterized paradigm, called parameterized pattern queries, that does not correspond to Baker’s initial definition, was proposed [53]. However, this model is indeed closely related to the theory developed by Baker. They use a set of symbols and a set of variables that correspond to Baker’s constant alphabet and parameter alphabet. They also defined a concept of valuation that could be associated with the mapping bijection and the $p$–match definition. The parameterized pattern queries paradigm was conceived as an extension of traditional pattern expressions to enhance the querying and clustering operations over sequence databases. Thus, the definition of a set of predicates on the variables (constraints) is also permitted under this new model. Furthermore, a KMP–based algorithm for this problem is also proposed. Experimental results showed that it notably decreases the query evaluation time compared to a naive approach.

In order to support more applications, parameterized matching was generalized to function matching by allowing the mapping function to be of any type, and not just bijections as in parameterized matching [4]. In other words, many symbols of the pattern can be mapped to the same text.
2.3 Parameterized Matching

A deterministic solution for the function matching problem, that runs in \(O(n|\Sigma_P| \log m)\) time, was devised [4]. Furthermore, they proposed a Monte Carlo algorithm that runs in \(O(n \log m)\) time with failure probability of \(1/n^k\), where \(k\) is a given constant. Function matching was also extended for the two–dimensional case and a randomized algorithm that runs in \(O(kn^2 \log n)\) time was proposed [4]. This algorithm has a \(1/n^k\) probability of reporting a false positive.

We derived an approximate version of function matching to permit certain degree of error. In particular, we proposed \(\delta \gamma\)--approximate function matching [106]. Given two integer strings, \(X = X_1...m\) and \(Y = Y_1...m\), and two given constants, \(\delta, \gamma \in \mathbb{N}\), we say that there is a match from \(X\) to \(Y\) if \(X\) can be transformed into a string \(X'\), by means of a function \(f\), such that \(X'\) is \(\delta\)--equal and \(\gamma\)--equal to \(Y\). Two equal-length integer strings are \(\delta\)--equal if the maximum difference between their corresponding symbols is at most \(\delta\); they are \(\gamma\)--equal if the sum of such differences is at most \(\gamma\). A \(O(nm)\) algorithm to find the \(\delta \gamma\)--function matches of a pattern \(P = P_1...m\) in a text \(T = T_1...n\) was proposed [106].

To support even a much wider range of applications, function matching was extended to the generalized function matching with don’t cares problem [6]. In this problem, the image of the mapping function can be any substring in \((\Sigma_C \cup \Sigma_P)^*\) and not just a single symbol as in function matching. Furthermore, an extra symbol \(\phi\), called the don’t care character, can be present in the strings. A \(\phi\) in the text matches any pattern symbol; a \(\phi\) in the pattern matches any text substring. This problem represents many pattern searching types but, as a consequence, it is much more complex. It was shown that the alphabet sizes appear in the exponent of the naive solution’s complexity which leads to a considerable difference between the cases of finite and infinite alphabets. A polynomial algorithm for the finite alphabet case was presented; for the case of infinite alphabets, it was demonstrated that the problem is \(NP–hard\) [6]. This is the first problem, so far, for which there is a polynomial solution for the finite alphabet case and there is not one for the infinite alphabet case.

2.3.4. Applications

Parameterized matching was initially defined as a tool for software maintenance [13]. This was motivated by the observation that programmers introduce duplicate code into large software systems when they are adding new features or fixing bugs possibly generated for not having considered special cases in the initial programming. Instead of adapting working sections of code, the programmers prefer to copy and slightly modify new instances of those sections in order to avoid making major revisions and introducing new bugs. They do it specially when the working sections were written by another programmer. With time, the amount of duplicate code is highly increased and the code gets larger, more complex and more difficult to maintain. For instance, when a new issue in a determined part of the program is fixed, it will not be automatically fixed in the other copies of that section of code and sometimes they may be hard to find.
The definition of parameterized matching assumes that some sections of code are copied and modified through text editors such that the corresponding copies are mostly the same, except for a systematic change of the variables and procedures’ names. Then, the code is considered as a sequence of tokens (variables, constants, operands, reserved keywords and procedure names) where the constant alphabet $\Sigma_C$ is comprised by the operands and the reserved keywords while the parameter alphabet $\Sigma_P$ is comprised by the variables, constants and procedures’ names.

**Example.** In Fig. 2-4 two edited code excerpts from the X Window [129] source code are presented. These two fragments are a p–match given that they are identical except for a correspondence between $pfi$ and $pfh$, $lbearing$ and $left$, and $rbearing$ and $right$. Notice that the p–matching sections are like expansions of the same macro with different parameters.

```c
    copy_number(&pmin, &pmax, pfi->min_bounds.lbearing,
   pfi->max_bounds.lbearing);
  *pmin++ = *pmax++ = ' ',
    copy_number(&pmin, &pmax, pfi->min_bounds.rbearing,
   pfi->max_bounds.rbearing);
  *pmin++ = *pmax++ = ' ',
    copy_number(&pmin, &pmax, pfh->min_bounds.left,
   pfh->max_bounds.left);
  *pmin++ = *pmax++ = ' ',
    copy_number(&pmin, &pmax, pfh->min_bounds.right,
   pfh->max_bounds.right);
  *pmin++ = *pmax++ = ' ',
```

**Figure 2-4.** Two sections of code that parameterized–match. Taken from [15].

In that sense, if it is required to look for all the copies of a determined section of code, the problem can be solved through a **parameterized fixed matching** algorithm by considering that section of code as the pattern and the code where copies are searched as the text. If the goal is finding all the copies of many sections of code (multiple patterns), then a **parameterized multiple matching** algorithm would be useful. If there exists no pattern, but the goal is finding all the pairs of duplicate code that have at least a determined length specified by the user, then the problem can be solved with a **maximal p–matches over a threshold length** algorithm, like DUP or PDUP.

Experiments with DUP on a large subsystem of over a million lines of code showed that 22% of the lines were involved in parameterized matching. This is a great amount of duplicate code, given that a proportional percentage of the code could be shrunk by using better programming techniques like procedures and functions. A reduction of this magnitude would make the code much more simple and easier to maintain. In general, all the parameterized matching problems and the approximate parameterized matching problems (under the p–edit and hamming distance) produce important results that facilitate the analysis of the code and provide useful information to simplify
2.3 Parameterized Matching

it and shrink it. This is the reason why software maintenance is still one of the main areas where parameterized matching is most useful at.

Other area of application of parameterized matching is image processing [77, 4]. Searching for color images on the web is an interesting problem [11, 139]. The Human–Computer Interaction Lab at the University of Maryland tackled the problem of searching for an icon in the screen. If the colors are fixed, the problem can be solved with an exact two-dimensional pattern matching algorithm. Nevertheless, sometimes the pattern image appears in other ranges of colors within the text, which makes impossible for exact–matching algorithms to find these occurrences. In this kind of cases it is proper to use two dimensional parameterized matching algorithms. However, images often have some errors resulting from distortion and loss of resolution, so such occurrences of a pattern image could not be reported by parameterized matching algorithms either (due to the absence of perfect bijections). But occurrences with these errors can indeed be found by taking either a function matching approach [4, 106] or an approximate parameterized matching approach under the hamming, p–edit, or $\delta_\gamma$ distance [19, 76, 77, 93, 103].

On the other hand, parameterized matching has important applications in databases. For instance, in a database that contains urls of the pages visited by different users, parameterized pattern queries can be used to retrieve useful information for improving the ergonomy of the site and finding the best places for advertisement ads [53]. For example, given the symbol $a$ and the variable $x$ where both represent urls, the query of the parameterized pattern expression $axa$ would retrieve the set of urls that the users have visited before coming back to the previously visited page represented by $a$. In a similar fashion, this idea can be used in computational biology to retrieve all the amino acids substrings that follow a determined structure where the presence of determined amino acids at certain positions are a constraint. This is also applicable to databases of any type, where the analysis over the sequential occurrence of elements is a matter of interest.

In general, parameterized matching and its related problems are considerably useful in any area where patterns are defined in terms of structural correlation across the positions. This motivates us to extrapolate its use to the solution of graph matching.
Part I.

Graph Isomorphism through Parameterized Matching
3. Our Approach: Graph Linearization

Our approach to determine whether two graphs, \( G_1 = (V_1, E_1) \) and \( G_2 = (V_2, E_2) \), are isomorphic consists of two main steps: (i) linearizing \( G_1 \) into a walk \( p = p_1...\ell \); and (ii) exploring all the walks in \( G_2 \) to determine whether there is one that parameterized-matches \( p = p_1...\ell \). In this chapter, we define graph linearization and parameterized matching on graph walks (Section 3.1). Then, we discuss characteristics and algorithms for linearization (Section 3.2). Finally, we propose an efficient algorithm that produces asymptotically length-optimal linearizations (Section 3.3).

3.1. Definition of Graph Linearization

**Definition 2 (Graph Linearization).** Let \( G = (V, E) \) be a connected undirected multigraph. A walk \( p = p_1...\ell \) of nodes and edges is a linearization of \( G \) iff:

1. \( p_i \) is a node \( v \in V \) if \( i \) is odd, \( 1 \leq i \leq \ell \).
2. \( p_i \) is an edge \( e \in E \) if \( i \) is even, \( 1 \leq i \leq \ell \), such that \( e = (p_{i-1}, p_{i+1}) \).
3. Each node \( v \in V \) and each edge \( e \in E \) appears at least once in \( p \).

In other words, the linearization \( p \) of a connected undirected graph \( G = (V, E) \) is an alternating sequence of nodes \( v \in V \) and edges \( e \in E \) that starts and ends at a node. Each intermediate occurrence of a node in \( p \) must be preceded and followed by an adjacent edge. Furthermore, all nodes and edges in the graph must appear in \( p \) at least once.

Our motivation for defining graph linearization is representing the topology of a multigraph through a walk. Specifically, the linearization \( p \) of \( G \) is a walk that represents all its adjacency relation, which we use to solve the graph isomorphism problem by comparing walks instead of multigraphs. For this purpose, we define parameterized matching on walks as follows:

**Definition 3 (Parameterized Match on Graph Walks).** Let \( G_1 = (V_1, E_1) \) and \( G_2 = (V_2, E_2) \) be two connected undirected multigraphs. Also, let \( V'_1 \subseteq V_1 \) and \( E'_1 \subseteq E_1 \) be subsets of nodes and edges in \( G_1 \); similarly, \( V'_2 \subseteq V_2 \) and \( E'_2 \subseteq E_2 \) are subsets of nodes and edges in \( G_2 \). The walks \( p = p_1...k, \) in \( G_1, \) and \( q = q_1...k, \) in \( G_2, \) are said to parameterized-match if and only if there exists a bijective function \( f : (V'_1 \cup E'_1) \rightarrow (V'_2 \cup E'_2) \) such that \( q_i = f(p_i) \) for \( 1 \leq i \leq k \).
The core idea of using parameterized matching to solve the graph isomorphism problem is as follows. Let \( p \) be a linearization of \( G_1 \); hence, \( p \) represents the topology of \( G_1 \). Thus, if a walk \( q \) in \( G_2 \) parameterized-matches \( p \), then \( p \) and \( q \) have the same topology. Consequently, considering that \( q \) represents \( G_2 \), we conclude that \( G_1 \) and \( G_2 \) are isomorphic. More formally, we prove the following theorem that solves the graph isomorphism problem through parameterized walks.

**Theorem 1.** Let \( G_1 = (V_1, E_1) \) and \( G_2 = (V_2, E_2) \) be two connected undirected multigraphs such that \( n = |V_1| = |V_2| \) and \( m = |E_1| = |E_2| \); also, let \( p = p_{1\ell} \) be a linearization of \( G_1 \). Then, \( G_1 \) and \( G_2 \) are isomorphic if and only if there exists a walk \( q = q_{1\ell} \) in \( G_2 \) such that \( p = p_{1\ell} \) parameterized-matches \( q = q_{1\ell} \).

**Proof.** In order to prove the theorem, we need to show that (i) if \( G_1 \) and \( G_2 \) are isomorphic, then there exists a walk \( q = q_{1\ell} \) in \( G_2 \) that parameterized-matches \( p = p_{1\ell} \); and (ii) if there exists a walk \( q = q_{1\ell} \) in \( G_2 \) that parameterized-matches \( p_{1\ell} \), then \( G_1 \) and \( G_2 \) are isomorphic.

First we prove (i). According to Problem 1, if \( G_1 \) and \( G_2 \) are isomorphic, there exists a bijective function \( f : \mathcal{E}_{G_1} \rightarrow \mathcal{E}_{G_2} \) for which Equation 1-1 is evaluated as \textit{true}. Notice that \( p = p_{1\ell} \) represents all the adjacency relation of \( G_1 \), which is defined on the left side of the biconditional. Considering the format of \( p_{1\ell} \) (see Definition 2) and the existence of a bijective function \( f \) that satisfies the right side of the biconditional, we can conclude that \( q = f(p_1)f(p_2)\cdots f(p_{\ell}) \) is a walk in \( G_2 \). Furthermore, as \( f(p_i) = q_i \), walks \( p = p_{1\ell} \) and \( q = q_{1\ell} \) parameterized-match.

Now we prove (ii). Let \( q = q_{1\ell} \) be a walk in \( G_2 \) that parameterized-matches \( p = p_{1\ell} \). Then, there exists a bijective function \( f : \mathcal{E}_{G_1} \rightarrow \mathcal{E}_{G_2} \) such that \( q_i = f(p_i) \) for all \( 1 \leq i \leq \ell \). Recall that all the nodes in \( V_1 \) and all the edges in \( E_1 \) appear at least once in \( p \) (see Definition 2) and that all the adjacency relation of \( G_1 \) is represented in \( p = p_{1\ell} \). Therefore, the existence of a bijective function \( f \) such that \( q_i = f(p_i) \), for all \( 1 \leq i \leq \ell \), implies that Equation 1-1 is evaluated as \textit{true}; then, \( G_1 \) and \( G_2 \) are isomorphic. \( \square \)

**Corollary 1.** Let \( G_1 = (V_1, E_1) \) and \( G_2 = (V_2, E_2) \) be two connected undirected multigraphs where \( |V_1| \leq |V_2| \). Also, let \( p = p_{1\ell} \) be a linearization of \( G_1 \). Then, \( G_1 \) is isomorphic to a subgraph in \( G_2 \) if and only if there exists a walk \( q = q_{1\ell} \) in \( G_2 \) such that \( p = p_{1\ell} \) parameterized-matches \( q = q_{1\ell} \).

**Proof.** A walk \( q = q_{1\ell} \) on \( G_2 \) is a linearization of a subgraph \( G = (V, E) \) of \( G_2 \) where \( V \) and \( E \) are the sets of nodes and edges, respectively, included in \( q = q_{1\ell} \) (see Definition 2). Then, because of Theorem 1, \( G \), a subgraph of \( G_2 \), is isomorphic to \( G_1 \). \( \square \)
3.2. Characteristics and Algorithms for Graph Linearization

There may be many linearizations that represent the same graph. Many factors such as different starting nodes and different visiting orders can result in different linearizations. However, a compact representation is preferable. For solving graph isomorphism, the length of the linearization is an important measure on the matching time. This is because a shorter linearization often leads to a smaller cost at the matching stage. Next, we define *length-optimal linearization*.

**Definition 4 (Length-Optimal Linearization).** *The linearization \( p = p_1 \ldots \ell \) of a connected undirected multigraph is length-optimal if the length of \( p \) (i.e., \( \ell \)) is minimum.*

Finding the linearization of a graph is very similar to the Chinese Postman Problem (CPP). CPP finds a walk that visits all the edges (and all the nodes) in the multigraph at least once; the only difference is that graph linearization does not require the starting node to be the same final node. A \( O(n^3 + m^2) \) algorithm for the CPP was proposed \([54]\). We can adapt this algorithm to calculate a length-optimal linearization. However, for large multigraphs, it is desirable to have algorithms with lower time complexity even if they do not produce length-optimal linearizations. Then, we consider BFS-L and DFS-L which produce linearizations using the graph traversal algorithms, as follows:

- **BFS-L** traverses the multigraph in the breadth-first search manner with some differences. When it explores a node, it visits all of its unexplored adjacent edges regardless if the node they lead to has been visited before. After discovering such nodes, it goes back to each one of them to (i) check if it has unexplored adjacent edges; and (ii) explore such unexplored edges (if there is any). But it is necessary to add in \( p \) all the walk that connects the current node and the next node to be explored; this significantly increases the length of the linearization walk. The algorithm terminates when there are no discovered nodes to be explored.

- **DFS-L** traverses the multigraph in the depth-first search manner. It starts from any node and makes recursive calls to visit the adjacent nodes that are connected to it through unexplored edges. The base case of the recursive call is when it reaches a node whose edges are all explored; then, it returns directly. Every time a recursive execution is finished, the execution instance that called it must add the corresponding connecting edge and source node in \( p \) again; this is because they need to be visited again so that other neighbours can be explored. The procedure finishes when it goes back to the starting node and it does not have any unexplored edges.

Notwithstanding, the linearizations produced by these algorithms can be long. As an attractive trade-off between length-optimality and efficiency, we propose a greedy approximation algorithm with an approximation guarantee.
3.3. Graph Linearization Algorithm - GLA

This section presents the GLA or Graph Linearization Algorithm. First, we describe the key ideas of the algorithm in Section 3.3.1; then we go through the details in Section 3.3.2. Its correctness is proven in Section 3.3.3. In Section 3.3.4 we present an upper bound for the length of GLA linearizations; furthermore, we show with empirical examples that, in practice, the produced linearizations are close to optimal (see Section 3.3.5). Finally, in Section 3.3.6, we present the complexity analysis.

3.3.1. Key Ideas

One of the challenges of linearization algorithms is visiting all the edges with short linearization length. Given that, in order to visit an edge, it is necessary to visit first one of its end nodes, we use the number of unexplored adjacent edges that the nodes have to conduct the traversal in a convenient manner. Particularly, GLA does the traversal in a similar way as DFS-L. The base case of the traversal is when we reach a node that has no unexplored adjacent edges; in this case, we say that such node is covered. In GLA, the number of uncovered nodes is stored. Then, if all the nodes are covered when a base case takes place, we do not add the way back up to the root of the DFS tree in the linearization.

Furthermore, GLA takes into account the number of unexplored edges of the nodes, at all stages of the process, to produce shorter linearizations. Specifically, we develop three heuristics: (i) the traversal starts from the node with the lowest degree; (ii) the unexplored edges that lead to already explored nodes are visited before than the ones that lead to unexplored nodes; and (iii) the edges that lead to unexplored nodes are considered sorted, in ascending order, on the number of unexplored edges they have. Heuristics (i) and (iii) aim to put the nodes that are close to be covered in the top levels of the DFS tree. On the other hand, heuristic (ii) aims to cover the nodes in the highest levels of the DFS tree at an early stage.

The first insight is that we want to cover the nodes at the top of the DFS tree early. This is because when the base case of the recursion is reached, going back to a node in a higher level of the DFS tree makes the linearization longer than going back to lower levels; thus, we want to reduce the probability of needing to go back to a high level. The nodes with low degree are more likely to be covered early; thus our first heuristic chooses the node with the lowest degree as the starting node of the traversal (the root of the DFS tree). The next level of the DFS tree are the root’s adjacent nodes that we discover first. And we want to cover them rapidly as well. Therefore, heuristic (iii) prefers the edges $e = (u, v)$ that lead to unexplored nodes $v$, sorted on the number of unexplored edges of $v$. Then, the nodes at the top levels of the DFS tree will be the ones with the lowest degrees, as the ones with the highest degrees may be covered in a lower level of the tree. Furthermore, to increase the probability of early covering the nodes at the top of the tree, we use heuristic (ii).
3.3 Graph Linearization Algorithm - GLA

More specifically, heuristic (ii) works as follows. When a node \( u \) is processed, all its unexplored edges \( e \) that lead to already explored nodes \( w \) are visited first. Notice that we do not want to continue the traversal on \( w \) as it is already being processed in a higher level of the DFS tree. All we need is exploring \( e \); however this implies visiting \( w \) again, so that the properties of the linearization are satisfied. Then, we visit \( e \) and \( w \) and, in order to return to \( u \), we visit \( e \) and \( u \) again. Notice that this decreases the number of unexplored edges of \( w \). Therefore, it is more likely that \( w \) is covered before starting all the way back up on the DFS tree when the base case of the recursion takes place. Furthermore, we do not need to add such way back into \( p \) if there are no other unexplored graph elements. In practice, this condition is satisfied quite often due to the combination of the three heuristics. Notice that these heuristics make the traversal explore one region of the multigraph before visiting another one; then, the produced linearization is shorter.

3.3.2 Algorithm

The pseudocode of the Graph Linearization Algorithm (GLA) is listed in Figure 3-1. Each graph element has a boolean attribute that indicates whether it has been explored. Furthermore, the number of unexplored graph elements is stored by variable \( unexplGE \). This variable is used to avoid reinserting graph elements in the backtracking of the DFS search tree when there are no unexplored elements left. The number of unexplored adjacent edges that each node \( v \) has is stored in \( v.NumUnexplEdges \). The produced linearization is implemented as the list \( p \). These variables are used during the DFS-like traversal to apply the heuristics presented in the last section. When a graph element is inserted into \( p \) for the first time, it has to be set as \textit{explored} and the number of unexplored graph elements in the graph, \( unexplGE \), must be decreased in one unit. Furthermore, if such graph element is an edge, the number of unexplored edges of its adjacent nodes must also be decreased.

The algorithm starts by setting every graph element as \textit{unexplored}. Then, for each node, the number of unexplored edges is calculated as the total number of adjacent edges it has. The walk \( p \) is set as empty and the number of unexplored graph elements is calculated as the sum of the number of nodes and the number of edges in the graph. After these initializations, a DFS-like traversal is performed using the recursive function in \textsc{TraverseGraph()} (see Figure 3-2) starting from the node \( u \) with the lowest degree, i.e. the first call of the procedure \textsc{TraverseGraph()} is performed over \( u \). This procedure is composed of the following steps:

1. Add \( u \) into \( p \).

2. Go to already explored nodes \( v \) through unexplored edges \( e \). Add \( e \) and \( v \) into \( p \). In case there are still unexplored graph elements, it goes back to \( u \) through \( e \); this implies adding \( e \) and \( u \) into \( p \) again.
3. Go to unexplored nodes \( v \) through unexplored edges \( e \). The nodes \( v \) must be considered sorted on their number of unexplored edges. Then \( e \) is added into \( p \) and the recursive procedure \( \text{TraverseGraph}() \) is called over \( v \). If there are still unexplored graph elements after this call, then \( e \) and \( u \) must be added again into \( p \) so that other neighbours of \( u \) can be visited. When there are no other neighbours to explore, the method returns so the linearization continues at a higher level of the DFS tree.

Notice that Step (3) is represented in Figure 3-2 through lines 8 - 14. In such lines it seems that, at each iteration, we find the node \( v \) with the minimum \( \text{NumUnexploredEdges} \); however, this is presented in this way just for clarity. Instead, we can sort the couples \( e = (u, v) \) on \( v.\text{NumUnexploredEdges} \) before the loop. Then, in each iteration, we consider each of such couples \( e = (u, v) \) in ascending order on \( v.\text{NumUnexploredEdges} \). The only additional operation we perform in each iteration is checking whether the adjacent node is still uncovered. This is because the value of each \( v.\text{NumUnexploredEdges} \) does not change, unless such \( v' \) is covered. Specifically, if \( v \) is explored at a lower level of the DFS tree traversal, we will not come back to this loop until \( v \) is covered. Checking if a node is covered takes constant time. Consequently, the complexity of this operation is the initial sorting which is \( O(d \log d) \) where \( d \) is the maximum degree of the nodes in \( V \).

The algorithm terminates when the first call to the recursive procedure finishes, i.e., when it goes back to the root of the DFS tree and there are no unexplored adjacent edges. At this point, \( \text{unexplGE} = 0 \) and \( p \) contains the linearization. The linearization produced by GLA for the graph presented in Figure 1-3(a) is \( Ae_1Be_3Ce_4De_5Ce_5De_6Be_2De_6E \); its length is 17.

---

**Algorithm 1**: GLA Algorithm

**Input**: \( G = (V, E) \)

**Output**: \( p \)

1. for every \( e \in E \) do \( e.\text{Explored} \leftarrow \text{false} \)
2. for every \( v \in V \) do
3. \( v.\text{Explored} \leftarrow \text{false} \)
4. \( S \leftarrow \{(u, v) \mid u \in V \land (u, v) \in E\} \)
5. \( v.\text{NumUnexploredEdges} \leftarrow |S| \)
6. choose \( u \in V_p \) with \( \min(u.\text{NumUnexploredEdges}) \)
7. \( p \leftarrow \langle \rangle, \text{unexplGE} \leftarrow |V| + |E| \)
8. \( \text{TraverseGraph}(G, u, p, \text{unexplGE}) \)
9. return \( p \)

---

**Figure 3-1.**: GLA algorithm.
### Algorithm 2: TRAVERSEGRAPH() Procedure

**Input:** $G = (V, E), u, p, \text{unexplGE}$

1. $p$.Add($u$), $u$.Explored $\leftarrow$ true, unexplGE--
2. for every $e \in E$ such that $e = (u, v)$ do
   3. if $!e$.Explored $\land$ v.Explored then
      4. $p$.Add($e$), $e$.Explored $\leftarrow$ true, unexplGE--, $p$.Add($v$)
      5. $u$.NumUnexplEdges--, $v$.NumUnexplEdges--
   6. if unexplGE $> 0$ do
      7. $p$.Add($e$), $p$.Add($u$)
   8. while there are unexplored edges $e = (u, v)$
      9. choose $e$ with $\min (v$.NumUnexploredEdges$)$
      10. $p$.Add($e$), $e$.Explored $\leftarrow$ true, unexplGE--
      11. $u$.NumUnexplEdges--, $v$.NumUnexplEdges--
      12. TraverseGraph($G, v, p, \text{unexplGE}$)
   13. if unexplGE $= 0$ then break
   14. $p$.Add($e$), $p$.Add($u$)

**Figure 3-2:** TRAVERSEGRAPH() procedure.

### 3.3.3. Correctness Proof

In this section, we first show that the output walk $p = p_1...\ell$ produced by GLA contains all the nodes and edges in the input multigraph $G = (V, E)$ (condition (3) of Definition 2). Then, we show that $p = p_1...\ell$ is indeed a linearization of $G$, i.e., we show that conditions (1) and (2) of Definition 2 are also satisfied.

The analysis for condition (3) uses two properties:

- **Property 1.** If a node is visited, all of its adjacent edges must be visited as well. According to lines 2 - 4 and lines 8-10 of TRAVERSEGRAPH() in Figure 3-2, all the unexplored edges of a node will be visited when such node is visited.

- **Property 2.** If a node is visited, all its adjacent nodes must be visited as well. This property is guaranteed by the code in lines 8 – 12 of TRAVERSEGRAPH() in Figure 3-2.

With the two properties above, we can prove the following lemma:

**Lemma 1.** Given a connected undirected multigraph $G = (V, E)$, the output walk $p$ produced by GLA includes all its nodes and edges.
Proof. We can prove the lemma by contradiction under two cases:

- **Case 1.** Suppose that all the nodes are explored but there exists at least one edge \( e = (u, v) \) that is not visited by GLA. This hypothesis contradicts Property 1: when the node \( u \) is visited, all its adjacent edges must be visited as well.

- **Case 2.** Suppose that there exists at least one node \( u \) that is not visited. Assume that we start the linearization from node \( v_0 \). Since the multigraph is connected, there is a walk from \( v_0 \) to \( u \); let us denote such walk as \( \langle v_0, v_1, v_2, \ldots, v_k, u \rangle \). We know that, for \( 0 \leq i < k \), \( v_i \) and \( v_{i+1} \) are adjacent nodes; \( v_k \) and \( u \) are adjacent too. Since \( v_0 \) is visited, according to Property 2, we know that \( v_1 \) is visited as well. Then, \( v_1, v_2, \ldots, v_k \) and \( u \) are visited as well. Thus, we have that \( u \) is also visited, which contradicts the hypothesis.

Therefore, all the nodes and edges in \( G \) must have been visited by GLA and included in its output linearization \( p \).

Then, the correctness of is proven by the following theorem:

**Theorem 2.** The Graph Linearization Algorithm (GLA) outputs a linearization \( p_{1,\ell} \) of the input multigraph \( G = (V,E) \).

**Proof.** We must prove that conditions (1), (2) and (3) of Definition 2 are satisfied for the output walk \( p = p_{1,\ell} \) generated by GLA. In particular, condition (3) is satisfied due to Lemma 1. Then, in the remainder of this proof we show that the walk \( p_{1,\ell} \) satisfies conditions (1) and (2).

Notice that when \( \text{TRAVERSEGRAPH()} \) is called for the first time (line 8, Figure 3-1), a node \( u \) is inserted (line 1, Figure 3-2). Then, if the \textit{if} statement of line 3 (Figure 3-2) is evaluated as \texttt{true}, edges \( e \) and nodes \( v \) are alternatively added for \( e = (u, v) \) where \( e \) is unexplored and \( v \) is explored (line 4). After this, the \textit{if} statement of line 6 is evaluated. If such evaluation yields \texttt{true}, the algorithm finishes; in such case, \( p_{1,\ell} \) is an alternating sequence of nodes and edges where the odd indices correspond to nodes and the even indices correspond to edges. Furthermore each \( p_i \), for even values in \( 1 \leq i \leq \ell \), is an edge that connects \( p_{i-1} \) and \( p_{i+2} \); this is because the graph elements inserted in \( p \) are adjacent (lines 2, 8, 9). Thus, the conditions (1) and (2) of Definition 2 are satisfied. Now, let us consider the case where the \textit{if} statement of line 6 is evaluated as \texttt{false}. In such case, \( e \) and \( u \) are added into \( p \) again before visiting the next neighbours; thus conditions (1) and (2) are still satisfied for the current fragment of the walk \( p \).

Then, in the loop of line 8, unexplored edges \( e = (u, v) \) that lead to unexplored nodes \( v \) are considered. Edge \( e \) is inserted in line 10. When procedure \( \text{TRAVERSEDGRAPH()} \) is called in line 12, \( v \) is inserted as well. Then, if the \textit{if} statement of line 13 is evaluated as \texttt{true}, conditions (1) and
3.3.4. Length of GLA Linearization

Theorem 3 shows that given the multigraph $G = (V, E)$, the length of the walk generated by GLA is at most 2 times the length of an optimal linearization. Therefore, the length produced by GLA is asymptotically optimal.

**Theorem 3.** GLA is 2-approximate with respect to the length of the length-optimal linearization.

**Proof.** Any linearization algorithm, including length-optimal algorithms, must traverse each edge of the multigraph at least once. Thus, for a multigraph $G = (V, E)$, where $n = |V|$ and $m = |E|$, the number of edges in its linearization is at least $m$. Since a linearization has the format of alternating between nodes and edges, a linearization with $k$ edges has $k + 1$ nodes. Hence, the optimal linearization $p^*$ has at least $m$ edges and $m + 1$ nodes. Therefore, $|p^*| \geq 2m + 1$.

When GLA linearizes a multigraph, it visits any edge at most twice. This is because, when procedure TRAVERSEGRAPH() is executed over node $u$, an unexplored edge $e$ that leads to any explored or unexplored node $v$ is added once into $p$ (lines 4 and 10, respectively, Figure 3-2). If after executing the next instructions there are still unvisited graph elements, it is necessary to go back to $u$ through $e$; this means that $e$ and $u$ are added into $p$ again (lines 7 and 14, Figure 3-2). After this, $e$ is not visited ever again given that only unexplored edges are considered (lines 3 and 8, Figure 3-2). Therefore, the number of edges in the linearization is at most $2m$. Again, since a linearization has the format of alternating between nodes and edges, the linearization $p^{GLA}$ has at most $2m$ edges and $2m + 1$ nodes. Therefore, we have $|p^{GLA}| \leq 4m + 1$. It leads to the approximation ratio of GLA,

$$\frac{|p^{GLA}|}{|p^*|} \leq \frac{4m + 1}{2m + 1} \leq 2$$

Notice that this theorem is based on the fact that each edge in the multigraph $G$ appears at most twice in the linearization $p = p_1...\ell$ generated by GLA. Then, $\ell$ is compared to a lower bound that visits each edge only once to show worst-case approximation ratio. However, even an optimal linearization may not achieve the lower bound for many graph structures. Thus, for average cases in practice, GLA linearization is much closer to the optimal, as elaborated with empirical examples in Section 3.3.5.
3.3.5. Empirical Comparison on the Length of Different Linearization Algorithms

This section compares GLA with the linearization methods presented in Section 3.2: BFS-L and DFS-L. Furthermore, we consider a length-optimal algorithm, which we denote as OPT-L. Specifically, we compare the algorithms on the length of the linearizations produced for multigraphs of different topologies.

![Graphs](image)

**Figure 3-3**: Examples of graphs with different topology: (a) cyclic graph, (b) complete graph, (c) star graph, (d) neuron graph, (e) balanced tree and (f) unbalanced tree.

Table 3-1 shows the length of the linearizations produced by each algorithm for each graph in Figure 3-3. We can see that GLA performs better than both DFS-L and BFS-L for all cases. In some of them, the difference is quite significant. Moreover, DFS-L is much better than BFS-L because the length of the linearizations produced by BFS-L is considerably increased by the long walks that connect the node being examined and the next node to be examined; in DFS-L those walks are much shorter because of the traversal order. Even though GLA is similar to DFS-L, GLA’s heuristics significantly reduce the length of the produced linearizations; they improve the locality of the walk so that GLA is likely to finish exploring one region of the graph before moving to another one, instead of shuffling among different regions that prolong linearization walks. In Table 3-1, we can see that these heuristics are effective: GLA performs close to OPT-L: GLA produces the optimal optimization for all the graphs in the table except for the unbalanced tree for which its result is close to the one of OPT-L.
3.3 Graph Linearization Algorithm - GLA

Table 3-1.: Comparison of the output length of different linearization algorithms.

<table>
<thead>
<tr>
<th>Graph</th>
<th>Figure</th>
<th>Nodes</th>
<th>Edges</th>
<th>BFS-L</th>
<th>DFS-L</th>
<th>GLA</th>
<th>OPT-L</th>
</tr>
</thead>
<tbody>
<tr>
<td>Square graph</td>
<td>Figure 3-3(a)</td>
<td>4</td>
<td>4</td>
<td>29</td>
<td>17</td>
<td>9</td>
<td>9</td>
</tr>
<tr>
<td>Complete graph</td>
<td>Figure 3-3(b)</td>
<td>4</td>
<td>6</td>
<td>35</td>
<td>25</td>
<td>17</td>
<td>17</td>
</tr>
<tr>
<td>Star graph</td>
<td>Figure 3-3(c)</td>
<td>5</td>
<td>4</td>
<td>31</td>
<td>24</td>
<td>13</td>
<td>17</td>
</tr>
<tr>
<td>Neuron graph</td>
<td>Figure 3-3(d)</td>
<td>12</td>
<td>12</td>
<td>124</td>
<td>49</td>
<td>35</td>
<td>35</td>
</tr>
<tr>
<td>Balanced tree</td>
<td>Figure 3-3(e)</td>
<td>7</td>
<td>6</td>
<td>53</td>
<td>25</td>
<td>17</td>
<td>17</td>
</tr>
<tr>
<td>Unbalanced tree</td>
<td>Figure 3-3(f)</td>
<td>11</td>
<td>10</td>
<td>79</td>
<td>41</td>
<td>35</td>
<td>31</td>
</tr>
</tbody>
</table>

3.3.6. Complexity Analysis

In this section, we derive the complexity of GLA for linearizing the graph $G = (V, E)$, where $n = |V|$ and $m = |E|$.

**Time Complexity.** In Figure 3-1, line 1 takes $O(m)$ time. The time complexity of lines 2 – 5 is $O(2m)$ because every edge in the graph is visited twice (as the graph is undirected). Line 6 takes $O(n)$. However, the complexity of GLA is dominated by the traversed walk (line 8, Figure 3-1) which corresponds to the linearization $p$. Notice that $p$ has at most $2m$ edges and $2m + 1$ nodes (as presented in Section 3.3.4). Each insertion takes constant time as it is always done at the end of $p$. But when a node is inserted for the first time, it is necessary to consider the unexplored adjacent edges $e$ that lead to unexplored nodes $v$ sorted on $v.NumUnexplEdges$ (lines 8–9, Figure 3-2). This sorting operation takes $O(d \lg d)$, where $d$ is the maximum degree of the nodes in $G$; specifically $d = \max_{v \in V} v.degree$. Thus, the time complexity of GLA is $O(2m + n(d \lg d)) = O(m + dn \lg d)$.

**Space Complexity.** The space requirement of GLA is given by a list that stores the linearization $p = p_1...\ell$. Thus, the space complexity of GLA corresponds to the length of the linearization, i.e. $\Theta(\ell)$. Because a GLA linearization can have at most $2m$ edges and $2m + 1$ nodes, the space complexity of the algorithm is $\Theta(m)$. 
4. Algorithm for Graph Isomorphism

In this chapter we present a linearization-based algorithm for solving graph isomorphism. In particular, the Parameterized Matching on multi-Graphs (PMG) algorithm uses a linearization of $G_1 = (V_1, E_1)$, denoted as $p = p_1...\ell$, and matches it against $G_2 = (V_2, E_2)$ to determine whether $G_1$ and $G_2$ are isomorphic by using Theorem 1. In Section 4.1 we present the high-level idea of the algorithm. Then, in Sections 4.2 and 4.3, we respectively present the pseudocode and prove its correctness. We present the complexity analysis of the algorithm in Section 4.4 while we show experimental results in Section 4.5. Finally, we discuss how to adapt this algorithm for subgraph isomorphism in Section 4.6.

4.1. Key Ideas

PMG considers all the possible injective functions $f : \mathcal{E}_{G_1} \rightarrow \mathcal{E}_{G_2}$ to determine whether there is mapping with two properties: (i) $f$ is bijective; and (ii) there exists a walk $q = q_1...\ell$ in $G_2$ such that $q = q_1...\ell$ parameterized-matches $p_1...\ell$. These possible injective functions are explored by traversing $p$ and $G_2$ simultaneously. Specifically, a graph element $p_i$ is compared to a graph element $ge$ in $G_2$ to determine whether an injective mapping is possible. We progressively extend a successful mapping by considering $p_{i+1}$ and an adjacent graph element of $ge$. Thus, when the mapping is successful for $p_\ell$, the traversed walk in $G_2$ parameterized-matches $p$; hence $G_1$ and $G_2$ are isomorphic.

The graph elements of $G_2$ are traversed in the depth-first manner while $p$ is traversed from left to right. Let us consider the DFS tree that represents the traversal of $G_2$. Each level $i$ of the DFS tree is comprised of graph elements from $G_2$. If $i$ is odd, it is a level of nodes; otherwise, it is a level of edges. Then, the idea of this traversal of $G_2$ is considering the possible injective mappings by attempting to set $f(p_i) = ge$ where $ge \in \mathcal{E}_{G_2}$ is a graph element at level $i$ of the DFS tree. In order to guarantee that the mapping is injective, two conditions must be verified: (i) if $p_i = p_j$, for $i < j$, and $f(p_i) = ge$, then the only valid mapping for $p_j$ is $ge$; and (ii) if $p_i \neq p_j$, for $i < j$, and $f(p_i) = ge$, then $f(p_j) \neq ge$. Notice that if we consider all the assignments in the walk from the root of the DFS tree to a leaf at the level $\ell$, we obtain a bijective function $f : \mathcal{E}_{G_1} \rightarrow \mathcal{E}_{G_2}$ such that $f(p_1) \cdots f(p_\ell)$ is a walk in $G_2$ that parameterized-matches $p_1...\ell$.

Next, we show our heuristics to prune the search space. At each step of the process, a node $u \in V_2$ and a node in $p_i$ are compared. Let us say that we set $f(p_i) = u$. In order to extend the match, we use node degrees and previous assignments in $f$ to prune the search space. Specifically, we
consider two cases:

**Case 1:** Node \( p_{i+2} \) is unassigned: We consider all the possible assignments \( f(p_{i+1}) = e \) and \( f(p_{i+2}) = v \) for edges \( e = (u,v) \in E_2 \) such that: (i) both \( e \) and \( v \) are unassigned; and (ii) \( v.\text{degree} = p_{i+2}.\text{degree} \). Condition (i) is to guarantee that \( f \) is injective; condition (ii) is a pruning criterion based on that fact that, if \( G_1 \) and \( G_2 \) are isomorphic, then analogous nodes must have the same degree. Notice that if \( p_{i+2} \) is unassigned, \( p_{i+1} \) is unassigned as well; this is because the assignment of an edge in \( p \) is done at the same time (or after) the assignment of its end nodes. The process continues by considering \( p_{i+2} \) and each \( v \).

**Case 2:** Node \( p_{i+2} \) is assigned to \( v \in V_2 \): There are two sub-cases. (a) Edge \( p_{i+1} \) is already assigned: it is not necessary to check adjacency as this was done when the mapping was set. We continue by considering \( p_{i+2} \) and \( v \). (b) Edge \( p_{i+1} \) is unassigned: the algorithm considers all the possible assignments \( f(p_{i+1}) = e \) for the unassigned edges \( e = (u,v) \). The process continues at \( p_{i+2} \) and \( v \).

Notice that the procedures for each of these cases guarantees that the mapping of both nodes and edges is injective. Furthermore, the DFS tree is expected to be sparse due to the pruning criteria; however, all the possible mapping functions are considered. If the algorithm reaches a successful assignment for \( p_\ell \), then the algorithm reports that the multigraphs are isomorphic.

### 4.2. Algorithm

The algorithm PMG, that determines whether \( G_1 = (V_1, E_1) \) and \( G_2 = (V_2, E_2) \) are isomorphic, is listed in Figure 4-1. The mapping function is represented as the array \( f \); namely, in such array, there is a position for each graph element \( ge \) in \( G_1 \) that stores its associated mapping \( f(ge) \) to \( G_2 \). On the other hand, boolean array \( g \) indicates if each graph element in \( G_2 \) is already assigned to a graph element in \( G_1 \) (through function \( f \)). The process starts by obtaining a linearization \( p = p_{1...\ell} \) of \( G_1 \) by means of GLA. Then, we initialize the mappings of all the graph elements in \( G_1 \) as **undefined** (which we abbreviate as **undef** in the pseudocode). Likewise, we set that none of the graph elements in \( G_2 \) has been assigned to graph elements in \( G_1 \).

After these initializations, we start the exploration of the search space. In particular, the DFS search trees are explored by calling the recursive procedure \texttt{EXTENDMATCH()} (see Figure 4-2). Each executing instance of this procedure considers a node \( p_i \), a node \( u \in V_2 \) and a copy of arrays \( f \) and \( g \). Furthermore, it is assumed that \( u \) has already been assigned to \( f[p_i] \). Then, what the procedure does is attempting to set the adjacent graph elements of \( u \), as mappings for \( p_{i+1} \) and \( p_{i+2} \), under the two cases presented in Section 4.1. The corresponding partial mappings are extended by recursive calls to \texttt{EXTENDMATCH()} according to the rules of these cases.
The roots of the DFS search trees, which correspond to the initial calls to the recursive procedure (line 8, Figure 4-1), are the nodes in $V_2$ that have the same degree as $p_1$. When we run PMG for the running example (see Figure 1-3), and $p = Ae_1Be_3Ce_4De_5Ce_5De_2Be_2De_6E$ is the linearization of $G_1$, the match is returned when either walk $q_1 = Xe'_1Ye'_3Ze'_4W'e'_5Ze'_5W'e'_2Ye'_2W'e'_6S$ or walk $q_2 = Xe'_1Ye'_3Ze'_5We'_4Ze'_4We'_2Ye'_2W'e'_6S$ is traversed. Notice that both $q_1$ and $q_2$ parameterized-match $p$. The mapping functions of these matches correspond to the functions $f_1$ and $f_2$ presented in Figure 1-3(c).

**Algorithm 3: PMG Algorithm**

**Input:** $G_1 = (V_1, E_1), G_2 = (V_2, E_2)$  
**Output:** true/false

1. $p = GLA(G_1)$
2. for every $ge \in (V_1 \cup E_1)$ do $f[ge] \leftarrow undef$
3. for every $ge \in (V_2 \cup E_2)$ do $g[ge] \leftarrow false$
4. for every $u \in V_2$ do
   5.     if $u.\text{degree} = p_1.\text{degree}$
   6.         $f' \leftarrow \text{copyOf}(f), f'[p_1] \leftarrow u$
   7.         $g' \leftarrow \text{copyOf}(g), g'[u] \leftarrow true$
   8.         if $\text{ExtendMatch}(u, p, 1, f', g', G_2) = true$
   9.             return true
10. return false

---

Figure 4-1.: PMG algorithm.

### 4.3. Correctness Proof

The correctness of the Parameterized Matching on multi-Graphs algorithm (PMG) is proven by the following theorem:

**Theorem 4.** The Parameterized Matching on multi-Graphs algorithm (PMG) determines whether two input multigraphs $G_1 = (V_1, E_1)$ and $G_2 = (V_2, E_2)$ are isomorphic.

**Proof.** We prove the correctness of PMG by showing the following invariant: when $u \in V_2$ and $p_i$ are processed in the recursive procedure $\text{EXTENDMATCH}()$, the walk $p_{1...i}$ parameterized-matches a walk $q_{1...i}$ in $G_2$ where $q_i = u$. Specifically, $q_{1...i} = f(p_1) \cdots f(p_i)$. Next, we show that this condition holds throughout the execution of the algorithm.
4.3 Correctness Proof

Algorithm 4: EXTENDMATCH() Function

Input: \( u, p = p_1\ldots e, i, f, g, G_2 = (V_2, E_2) \)
Output: \( \text{true/false} \)

1. if \( i = \ell \) then return \text{true}
2. if \( f[p_{i+2}] = \text{undef} \)
3. for every \( e = (u, v) \in E_2 \) do
4. \( g[v] = \text{false and } g[e] = \text{false and } v.\text{degree} = p_{i+2}.\text{degree} \)
5. \( f'[p_{i+1}] = e, f'[p_{i+2}] = v \)
6. \( g'[e] = \text{true, } g'[v] = \text{true} \)
7. if EXTENDMATCH\((v, p, i + 2, f', g', G_2) = \text{true} \)
8. return \text{true}
9. else
10. \( v = f[p_{i+2}] \)
11. if \( f[p_{i+1}] = \text{undef} \)
12. for every \( e = (u, v) \in E_2 \) such that \( g[e] = \text{false} \)
13. \( f'[p_{i+1}] = e \)
14. \( g'[e] = \text{true} \)
15. if EXTENDMATCH\((v, p, i + 2, f', g', G_2) = \text{true} \)
16. return \text{true}
17. else
18. if EXTENDMATCH\((v, p, i + 2, f, g, G_2) = \text{true} \)
19. return \text{true}
20. return \text{false}

Figure 4-2.: EXTENDMATCH() function.

- **Initialization:** When \( i = 1, p_1 \) parameterized-matches \( u \) given that \( f(p_1) = u \) was set in line 6 - Figure 4-1 before calling EXTENDMATCH\((u, p_1) \) (line 8, Figure 4-1).

- **Maintenance:** Let us assume that the invariant holds for \( p_i \) and \( u \). Then, we have to consider two cases: (i) \( f(p_{i+2}) \) has not been defined; and (ii) \( f(p_{i+2}) = v' \). Let us consider first case (i). Notice that, because of lines 3 – 4 in Figure 4-2, EXTENDMATCH\((v, p_{i+2}) \) is only called for unassigned nodes \( v \) such that \( e = (u, v) \in E_2 \) is also unassigned; thus, adding \( f(p_{i+1}) = e \) and \( f(p_{i+2}) = v \) (line 5, Figure 4-2) maintains the injective property of \( f \). Hence, when EXTENDMATCH\((v, p_{i+2}) \) is executed, \( p_1\ldots i + 2 \) parameterized-matches the walk \( f(p_1) \ldots f(p_{i+2}) \) in \( G_2 \) and \( f(p_{i+2}) = v \). For the case (ii), we have two sub-cases: that \( p_{i+1} \) is already assigned or that it is not. In the former, we need not change the function and, thus, the invariant will hold when EXTENDMATCH\((v, p_{i+2}) \) is called (line 18, Figure 4-2)). In the latter, the algorithm only considers unassigned edges \( e = (u, v') \) for the mapping of \( f(p_{i+1}) \) (lines 12 – 13, Figure 4-2)); hence, when EXTENDMATCH\((v', p_{i+2}) \) is called in line 15, the mapping \( f \) will still be injective and
Algorithm for Graph Isomorphism

\[ f(p_{i+2}) = v'. \]

- **Termination:** When no matches can be extended until \( p_\ell \), all the calls of \( \text{ExtendMatch()} \) return \( false \) (see line 20); hence, PMG returns \( false \) as well (line 10, Figure 4-1). This is correct as all the possible mapping functions were considered. Namely, (i) the search was started from all the valid nodes (lines 5 – 10, Figure 4-1); and (ii) all the possible mappings were considered at all stages of the search (lines 3 – 5 and 12, Figure 4-2). On the contrary, if a match is extended in any branch of the search and reaches position \( \ell \) (line 1, Figure 4-2), we obtain that \( p_{1...\ell} \) parameterized-matches the walk \( q_{1...\ell} = f(p_1) \cdots f(p_\ell) \) in \( G_2 \); then, PMG returns \( true \) (line 9, Figure 4-1)). In such case, considering Theorem 1, we conclude that \( G_1 \) and \( G_2 \) are isomorphic.

\[ 4.4. \text{ Complexity Analysis} \]

In this section, we analyse the complexity of determining isomorphism of multigraphs \( G_1 = (V_1, E_1) \) and \( G_2 = (V_2, E_2) \), where \( n = |V_1| = |V_2| \) and \( m = |E_1| = |E_2| \), using PMG.

**Time Complexity.** The PMG algorithm performs two basic steps: (i) linearizing multigraph \( G_1 \) into a walk \( p = p_{1...\ell} \); and (ii) matching the linearization \( p = p_{1...\ell} \) against multigraph \( G_2 \). The former takes \( O(m +dn\log d) \), where \( d \) is the maximum degree of the nodes in \( G_1 \), as presented in Section 3.3.6. The complexity of the latter is studied in this section.

The initialization of the arrays \( f \) and \( g \) takes \( O(n+m) \) (lines 2 – 3, Figure 4-1). However, this cost is negligible with respect to the number of executions of the recursive procedure \( \text{ExtendMatch()} \); each execution requires constant time. This number is equal to the number of nodes and edges in the DFS search trees. The number of edges in a DFS tree is equivalent to the number of nodes: each node, except the root, is associated to an edge that leads to its parent. Then, the asymptotic behavior of PMG depends on the number of nodes in the DFS trees. Next theorem gives an upper bound for this number.

**Theorem 5.** Let \( p = p_{1...\ell} \) be a linearization of multigraph \( G_1 \). Also, let \( d \) be the maximum degree of the nodes in multigraph \( G_2 \); specifically \( d = \max_{v \in V_1} v.\text{degree} \). The DFS tree that represents the traversal of \( G_2 \) done by PMG has at most \( O(d^{\lfloor \ell/2 \rfloor}) \) nodes.

**Proof.** Let \( n \) and \( m \) be the cardinality of the sets of nodes and edges, respectively, of both \( G_1 \) and \( G_2 \) (i.e., \( n = |V_1| = |V_2| \), \( m = |E_1| = |E_2| \)). In the worst case, all the nodes in \( V_1 \) and \( V_2 \) have the same degree; in such case, the pruning criterion based on only matching the nodes \( v \in V_2 \) with the same degree of \( p_i \) is not useful. The DFS tree rooted at one of the nodes of multigraph \( G_2 \) has \( \lfloor \ell/2 \rfloor \) levels of nodes and \( \lfloor \ell/2 \rfloor \) levels of edges; we just consider the levels of nodes. The root has
4.5 Experimental Evaluation

one node and the second level has \( d \) nodes. Each of these \( d \) nodes is associated to \( d - 1 \) nodes in the third level (as the edges that lead to nodes in upper levels of the tree are not considered); thus, the third level has \( d(d - 1) \) nodes. Similarly, the fourth level has \( d(d - 1)(d - 2) \) nodes. In general, level \( i \) of the tree has \( \prod_{j=0}^{i-2}(d - j) \) nodes. Thus, the total number of nodes of a DFS tree is:

\[
1 + \sum_{i=2}^{\lceil \ell/2 \rceil - 2} \prod_{j=0}^{i-2}(d - j) = O(d^{\lceil \ell/2 \rceil - 1})
\]

Since a linearized walk alternates between nodes and edges while starting and ending at a node, \( \ell \) is odd. Thus, \( O(d^{\lceil \ell/2 \rceil - 1}) = O(d^{\lceil \ell/2 \rceil}) \).

As we have a DFS tree starting at each node in \( G_2 \), the total number of nodes visited, and hence the time complexity of PMG, is \( O(nd^{\lceil \ell/2 \rceil}) \). Thus, it is important to have a short linearization of \( G_1 \). Note that if \( G_2 \) is complete, i.e., \( d = n - 1 \), the time complexity is \( O(n(n-1)^{\lceil \ell/2 \rceil}) = O(n^{\lceil \ell/2 \rceil}) \).

However, it is important to remark that Theorem 5 gives an upper bound for the worst-case complexity. It assumes that, at every level of nodes, all the possible neighbors are explored. The average-case situations are often not that “bad” because: (i) when a node \( p_i \) has already been assigned, only such assigned node is considered; and (ii) when the multigraph has varied node degrees, the pruning criterion highly reduces the number of adjacent nodes to be visited. Thus, in practice, our algorithm has a better performance than the given worst-case bound.

**Space Complexity.** PMG compares the linearization \( p \) of \( G_1 \) with its potential parameterized-matching walks in \( G_2 \). Notice that only one of such walks is considered at a time. Therefore, PMG only needs to store the mapping table of the linearization with respect to the current walk being considered. Such table contains mapping for all the nodes and edges. Hence, the space complexity of PMG is \( \Theta(n + m) \).

4.5 Experimental Evaluation

We assess the performance of our proposed approach experimentally. We implement our proposed framework in both Python and C# and release them as public resources \(^1\) to help future comparison studies. We compare our approach to VF2, using an optimized implementation from the networkX library \(^2\). Both are implemented in Python and open-sourced.

\(^1\)http://ids.postech.ac.kr/graph

\(^2\)http://networkx.github.io
As datasets, we employ a set of public benchmark graphs (Section 4.5.1) and synthetic graphs (Section 4.5.2). All evaluations are performed on a server running under a Windows platform on a 3.40GHz CPU with 16GB memory.

### 4.5.1. Benchmark Graphs

In this section, we study performance over a collection of public benchmark graphs used for evaluating isomorphism papers, to validate the generality of GLA. More specifically, we consider the following three widely adopted families of graphs in the public repository. Each family has graphs of size up to one thousand nodes.

- Strongly regular graphs: Graphs with high regularity. The repository has 87 pairs, covering families of Steiner Triple, Latin Square, Paley, Lattice, and Triangular graphs used in prior literature.

- Component-based graphs: Graphs connecting regular graphs as a component. The repository has 84 pairs, covering a union of regular graphs, cliques, or tripartite graphs used in existing work.

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3https://sites.google.com/site/giconauto/home/benchmarks
4.5 Experimental Evaluation

- Graphs based on Miyazaki’s construction: Graphs following Miyazaki’s construction to deliberately add complexity to the problem. There are 195 instances available in the repository.

Figure 4-3 first illustrates the response time differences of VF2 and GLA for all 366 instances. *Test id* is assigned, in ascending order, on GLA response time (X-axis); the Y-axis represents the response time of the two algorithms. Observe that in 181 pairs, about half of the datasets, GLA terminates earlier. This is encouraging as the benchmark datasets are intentionally biased into “strongly regular” graphs. For example, nodes in the graphs in the strongly regular graph group are all of the same degree, and node degrees in the component-based graphs also coincide into few values (e.g., degree of a node within the regular component and degree of a connecting node). Our target scenario is supporting graphs that are more structurally heterogeneous. The third group of Miyazaki’s construction adds some such variation, from which GLA clearly excels VF2. In summary, GLA shows fast response in the half of all benchmark graphs.

Table 4-1 and 4-2 show the break down of this result. In Table 4-1, our empirical finding is consistent with the above analysis, as VF2 excels in regular graphs, while GLA is significantly faster in 65% (126 out of 195) in Miyazaki-based constructed graphs. This is an interesting result since Miyazaki-constructed graphs constitute one of the hardest cases for graph isomorphism algorithms [141]. Table 4-2 shows cases where GLA is not short-running, or takes 2+ minutes. However, in the majority of 272 short-running cases, namely 66% of such cases, GLA runs faster. This opens up a possibility of a hybrid algorithm that selects between these two algorithms, either statistically-based on the graph topology or dynamically after running for some time, which we leave as future work.

<table>
<thead>
<tr>
<th></th>
<th>Strongly Regular graphs</th>
<th>Component-Based Graphs</th>
<th>Miyazaki’s construction</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>VF2</td>
<td>63</td>
<td>53</td>
<td>69</td>
<td>185</td>
</tr>
<tr>
<td>GLA</td>
<td>24</td>
<td>31</td>
<td>126</td>
<td>181</td>
</tr>
</tbody>
</table>

4.5.2. Synthetic Graphs

In this section, we study performance over synthetically generated graphs to isolate the factors that positively and negatively affect the performance of the two algorithms. For graph generation, we deliberately avoid the “trivial cases”. For example, consider a graph where node $v_i$ is connected to $v_1, \ldots, v_{i-1}$, for $1 < i \leq n$. As the degree of each node is unique, testing isomorphism can be done trivially by using a simple heuristic like sorting nodes by degree. In contrast, we consider cases
Table 4-2: Ratio of short-running cases of GLA and VF2 on the benchmark graphs.

<table>
<thead>
<tr>
<th></th>
<th>Strongly Regular graphs</th>
<th>Component-Based Graphs</th>
<th>Miyazaki's construction</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>VF2</strong></td>
<td>100% (87/87)</td>
<td>100% (84/84)</td>
<td>100% (195/195)</td>
<td>100% (366/366)</td>
</tr>
<tr>
<td><strong>GLA</strong></td>
<td>55.2% (48/87)</td>
<td>94.0% (79/84)</td>
<td>74.4% (145/195)</td>
<td>74.3% (272/366)</td>
</tr>
</tbody>
</table>

where no such simple heuristic can be used. Graphs where every node has identical degree would be much more challenging in that sense.

Meanwhile, we also randomize the node degrees to complement benchmark studies focusing on regular topologies. Given that the complexity of VF2 is reported to vary significantly over degree, from $O(n^2)$ to $O(nn!)$ [44], we consider both low- and high-degree cases to evaluate algorithms in a wide spectrum of settings. The lower end of this spectrum is observed when the matching graphs are early found in a sparse graph, while the opposite case of dense graphs often leads to long running times. More specifically, we generate sparse and dense identical-degree graphs as follows: 1-Sparse: We generate a random graph $G$, with $n$ nodes and $3n$ edges, where every node has degree three. We first build a random binary tree with $n-1$ edges. Then, the nodes with the degree less than three are connected to another such node randomly chosen. 2-Dense: We generate graph $G'$ by subtracting $G$ from a complete graph. Every node of $G$ has the same degree (i.e., $n-4$).

In each setting, we vary the number of nodes from 16 to 256 to evaluate the response time of GLA and VF2. For each point in the figures, we randomly generate 45 graphs and report the median response time. We choose median response time as our performance metric because the running time on different graphs significantly varies over graph complexity (as discussed above) while the optimization margin is narrow for easy cases and hard extremes. Our target problems are thus neither of these. Namely, using the average or min/max as the main performance metric would bias the results to represent either extreme. In contrast, median would filter out extreme results.

Figure 4-4(a) and (b) show the results for sparse graphs and dense graphs, respectively. The $X$-axis is the number of nodes (in log scale) and the $Y$-axis is the median response time in milliseconds. Note the two figures have different scales. Furthermore, the number of edges is linear with the number of nodes for sparse graphs and quadratic for dense graphs. In Figure 4-4(a), the median running time of GLA remains more or less constant to 10 milliseconds, despite the increase in graph size. As a result, when $n = 256$, GLA outperforms VF2 by an order of magnitude. In Figure 4-4(b), we observe a consistent trend, except that the performance gap is larger. In particular, for $n = 256$, GLA is faster by two orders of magnitude. These figures show that GLA has low res-
4.6 PMG-SI: Solution for Subgraph Isomorphism

In this section we present an adaptation of PMG to solve the subgraph isomorphism problem. Specifically, we determine whether graph \( G_1 = (V_1, E_1) \), where \( n_1 = |V_1| \) and \( m_1 = |E_1| \), is response time, less than VF2. This can be explained by the effective pruning of the notoriously large search space, which is guided by the heuristics employed during the linearization and matching phases.

**Figure 4-4:** Response time of GLA and VF2: (a) on sparse graphs; and (b) on dense graphs.

response time, less than VF2. This can be explained by the effective pruning of the notoriously large search space, which is guided by the heuristics employed during the linearization and matching phases.

4.6. PMG-SI: Solution for Subgraph Isomorphism

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Algorithm for Graph Isomorphism

isomorphic to a subgraph in $G_2 = (V_2, E_2)$, where $n = |V_2|$ and $m = |E_2|$. We solve this problem by means of the linearization approach, which is correct due to the Corollary 1 of Theorem 1. In Section 4.6.1 we present the algorithm and in Section 4.6.2 we present some experimental results.

4.6.1. Algorithm

The algorithm is essentially the same as the one presented in Figure 4-1, considering that we linearize the smallest graph (i.e., $G_1$). However, some aspects must be taken into account. For instance, the length of array $f$ is $n_1$ while the length of array $g$ is $n$. Furthermore, the heuristic to prune on node degrees must be modified: instead of an equality constraints, we must use inequalities. In particular, the condition of line 5 in Figure 4-1 is replaced by $u.\text{degree} \geq p_1.\text{degree}$; similarly, the corresponding constraint of line 4 in Figure 4-2 is replaced by $v.\text{degree} \geq p_{i+2}.\text{degree}$. This might decrease the pruning power of the node degree heuristic, especially when the node degrees in $G_2$ are significantly higher than the ones of $G_1$.

Algorithm 5: PMG-SI Algorithm

**Input:** $G_1 = (V_1, E_1), G_2 = (V_2, E_2)$

**Output:** $\mathcal{R}$

1. $p = \text{GLA}(G_1), \quad \mathcal{R} = \emptyset$
2. for every $ge \in (V_1 \cup E_1)$ do $f[ge] \leftarrow \text{undef}$
3. for every $ge \in (V_2 \cup E_2)$ do $g[ge] \leftarrow \text{false}$
4. for every $u \in V_2$ do
5. if $u.\text{degree} \geq p_1.\text{degree}$
6. $f' \leftarrow \text{copyOf}(f), \quad f'[p_1] \leftarrow u$
7. $g' \leftarrow \text{copyOf}(g), \quad g'[u] \leftarrow \text{true}$
8. $\text{ExtendMatchSI}(u, p, 1, f', g', G_2, \mathcal{R})$
9. return $\mathcal{R}$

**Figure 4-5:** PMG-SI algorithm.

Figure 4-5 presents a variation of the algorithm, called PMG-SI, that reports all the subgraphs of $G_2$ that are isomorphic to $G_1$ in the set $\mathcal{R}$. The DFS traversal is performed by the recursive procedure $\text{ExtendMatchSI}()$ (see Figure 4-6). The correctness proof and the complexity analysis presented in Sections 4.3 and 4.4, respectively, also apply for PMG-SI. Thus, its time complexity is $O(nd^{\ell/2})$, where $d$ is the maximum node degree in $G_2$ and $\ell$ is the length of $G_1$’s linearization.

4.6.2. Experimental Evaluation

We evaluate the performance of our approach for solving subgraph isomorphism under a variety of graph sizes. We first show the experimental setup and then we show the results.
Algorithm 6: EXTENDMATCHSI() Procedure

Input: u, p = p₁..., ℓ, i, f, g, G₂ = (V₂, E₂), ℛ

1. if i = ℓ then return ℛ. Add(f)
2. if f[pᵢ+2] = undef
3. for every e = (u, v) ∈ E₂ do
4.   if g[v] = false and g[e] = false and v.degree ≥ pᵢ+2.degree
5.     f' ← copyOf(f), f'[pᵢ+1] ← e, f'[pᵢ+2] ← v
6.     g' ← copyOf(g), g'[e] ← true, g'[v] ← true
7.     ExtendMatchSI(v, p, i + 2, f', g', G₂, ℛ)
8. else
9.   v = f[pᵢ+2]
10. if f[pᵢ+1] = undef
11. for every e = (u, v) ∈ E₂ such that g[e] = false
12.   f' ← copyOf(f), f'[pᵢ+1] ← e
13.   g' ← copyOf(g), g'[e] ← true
14.   ExtendMatchSI(v, p, i + 2, f', g', G₂, ℛ)
15. else
16.   ExtendMatchSI(v, p, i + 2, f, g, G₂, ℛ)

Figure 4-6: EXTENDMATCHSI() procedure.

Experimental Setup

Implementation. We implement the GLA algorithm and the PMG algorithm in C#.

Small graphs. We employ graphs G₁ = (V₁, E₁) with different sizes and topologies. In order to vary the topology, we consider complete graphs, path graphs, cyclic graphs and star graphs. A cyclic graph is a path in which the first node is the same as the last node. A star graph is a graph in which one node is connected to every other node.

Large graphs. We generate graphs G₂ = (V₂, E₂) using the Recursive Matrix (RMAT) model [31] that generates scale-free graphs similar to the types of graphs used in many applications. We use graphs with different sizes: |V₂| = 1024, 16384, 131072, 524288, 1048576. The number of edges of each graph is |E₂| = 5 × |V₂| representing sparse graphs.

Hardware. We perform the experiments on a commodity server with 3,300GHz Intel Xeon X5680 CPU with 24GB RAM running Windows Server 2008R2.
**Metrics.** Our main performance metric is the query response time. We report the number of graph element comparisons as this is the dominant factor in time complexity. For reference, on our server, 400000 comparisons are performed per second. We also report the length of the linearization generated for $G_1$.

**Experimental Results**

The small graphs $G_1 = (V_1, E_1)$ were linearized into walks $p = p_1...\ell$ by means of the GLA algorithm. Then, they were queried on each one of the large graphs $G_2 = (V_2, E_2)$, where $n = |V_2|$ and $m = |E_2|$, via the PMG algorithm. The behaviour of the matching process was very similar on all the large graphs; thus, we analyse in detail only the experiment on the large graph with $|V_2| = 1024$.

- **For complete graphs:** The length of the linearization grows linearly with number of edges in the graph $G_1$ as expected from Theorem 3 (see Figure 4-7(a)). However, it grows at a quadratic rate with $V_1$ (see Figure 4-7(b)); this can be explained by the fact that in complete graphs $|E_1| = O(|V_1|^2)$. The time taken by the algorithm grows faster for low values of $\ell$ than for greater values (see Figure 4-7(c)). This is because for low values there are more matches and, hence, more graph elements need to be explored; for greater values, mismatches are early detected.

- **For path graphs:** The length of the linearization is linear respect to both the number of nodes and edges in $G_1$ (see Figure 4-8(a,b)). This is because in paths $|E_P| = O(|V_P|)$. The time taken by PMG grows exponentially on the length of the linearization of $G_1$ (see Figure 4-8(c)). This verifies the time complexity analysis of PMG in Section 4.4, which concludes that its time complexity is $O(nd^{\ell/2})$ where $d$ is the maximum degree of the nodes in $G_1$. Given that, as the experiment is done on a fixed $G_2$, both $n$ and $d$ are constant. Consequently, we obtain an exponential function of $\ell$. Given that for cyclic and star graphs, $|E_P| = O(|V_P|)$, the performance for such graphs is similar.

In Figure 4-9, we show a comparison of the performance among all the different small graphs $G_1$ against two large graphs $G_2$. In Figure 4-9(a) we show the linearization length of the different types of graphs $G_1$. Because of the number of edges, the linearization of the complete graphs is significantly much longer in all cases. Figure 4-9(b,c) show the time taken by PMG for all the types of graphs $G_1$ in the graphs $G_2$ of $|V| = 1024$ and 1,048,576, respectively.

Notice that the time taken by the cyclic graphs is considerably greater than the other types of graphs $G_1$ on both large graphs $G_2$. This is because, even though paths, star and cyclic graphs of a given $|V_1|$ have a similar topology, their number of edges are $|V_1| - 1$, $|V_1| - 1$ and $|V_1|$, respectively. So let us compare the matching of a cyclic graph with both the path and the star graphs. A cyclic graph is the same as a path but with an additional edge that connects the first node with the last one.
4.6 PMG-SI: Solution for Subgraph Isomorphism

Figure 4-7.: Subgraph isomorphism for complete graphs $G_1 = (V_1, V_2)$ on a graph $G_2 = (V_2, E_2)$ where $|V_2| = 1024$. (a) Linearization obtained by GLA for different values of $E_1$. (b) Linearization obtained by GLA for different values of $|V_1|$. (c) Time taken by PMG to solve the subgraph isomorphism problem for different values of the linearization length.

Figure 4-8.: Subgraph isomorphism for path graphs $G_1 = (V_1, E_1)$ on a graph $G_2 = (V_2, E_2)$ where $|V_2| = 1024$. (a) Linearization obtained by GLA for different values of $E_1$. (b) Linearization obtained by GLA for different values of $|V_1|$. (c) Time taken by PMG to solve the subgraph isomorphism problem for different values of the linearization length.

Figure 4-9.: Experimental results of different types of graphs $G_1 = (V_1, E_1)$ (complete, path, cyclic and star graphs) on different graphs $G_2 = (V_2, E_2)$. X-axis is $|V_1|$. (a) Linearization length obtained by GLA for all the graphs $G_1 = (V_1, E_1)$. Number of graph element comparisons used in PMG for graphs $G_2 = (V_2, E_2)$ with (b) $|V_2| = 1024$, (c) $|V_2| = 1048576$.

Thus, the linearization of a cyclic graph of a given $V_1$ exceeds in one the length of the linearization of a path with the same number of nodes. Consequently, the time required to solve the problem for a cyclic graph with $|V_1|$ nodes is equivalent to the time required for a path with $|V_1| + 1$ nodes (as
it can be verified in Figure 4-9(b,c)).

Even though the number of edges of a graph with \(|V_1|\) nodes is \(|V_1| - 1\) for a star graph and \(|V_1|\) for a cyclic graph, the linearization of the star graph can be longer. This is because most edges of the star graphs are visited back and forth in order to return to the central node. However, because the mapping of a given edge \(e \in E_1\) is established only once, the second occurrence of the edge in the linearization is skipped. Therefore, the time taken for a star graph of a given \(|V_1|\) is similar to the time of a path of the same size but much less than the time of a cyclic graph of the same size. On the other hand, even though complete pattern graphs have long linearizations, the time required for the matching phase is low due to the early detection of mismatches.
Part II.

Queries on Attributed Graphs Solved through Parameterized Matching
5. Generalized Pattern Queries

In the last decades, the use of multigraphs to represent information of different types has widely spread. In particular, attributed multigraphs have been used to represent social networks [114], communication networks [20] and bioinformatics structures [92], to name some. In this chapter, we define the attributed multigraph model considered in this thesis (Section 5.1). Furthermore, we introduce a new type of queries on attributed multigraphs: generalized pattern queries. We show that our queries support both reachability and pattern match queries and even queries that cannot be represented under these models (Section 5.2).

5.1. Graph Model

We consider an attributed multigraph, a graph in which nodes and edges have attributes; furthermore, more than one edge may connect the same pair of nodes representing different relationships. Next, a more formal definition is presented:

**Definition 5 (Attributed Multigraph).** An attributed multigraph $G = (V, E, f_V, f_E)$ is a multigraph where:

- $V$ is a set of nodes.
- $E$ is a multiset of edges on $V \times V$.
- $f_V$ is a node-attribute function defined on the set of nodes $V$ such that, for each $v \in V$, $f_V(v) = (A_1 = a_1, \ldots, A_d = a_d)$ where $A_k$ is an attribute and $a_k$ is the value assigned to the corresponding attribute.
- $f_E$ is an edge-attribute function defined on $E$ in a similar way as $f_V$ is defined on the set of nodes $V$.

Furthermore, we assume that each node $v$ has at least two attributes: $v.id$ which is a unique identifier, and $v.type$ which is a label (or categorical attribute) called type. Figure 5-1 shows an example of an attributed multigraph modelling a social network. It contains nodes of two types: person and photo, and edges of three types: friend, colleague and tag. A person node may have the gender attribute; its value is male for the nodes with ids “Mike” and “Nick” and female for the nodes with ids “Alice”, “Mary”, “Rose” and “Kate”.
Unlike previous literature that does not consider attributes for edges, we support edges with attributes. This can be useful, for instance, to consider the date when the friendship or the tag was established. In Figure 5-1, we can see that Rose and Nick became friends on December 2, 2013.

![Diagram](image)

**Figure 5-1.** Example of an attributed multigraph that represents a social network. Node type is *person* or *photo*, and edge type is *friend*, *colleague* or *tag*.

As graph processing generally requires random data access (no locality), we assume that the graph is managed in main memory to allow fast random access using pointer-base representation. Furthermore, if the graph is directed, each node has references to both inbound and outbound edges to allow queries to traverse both directions.

5.2. Query Model

We propose a novel query model, which allows to find paths, walks, and subgraphs that satisfy reachability requirements. We call our queries *generalized pattern queries*. In this section, we first introduce a few new concepts, and we use them to define our query model. We also present examples and discuss the expressive power of our queries. We show that our queries can be used to express both reachability queries and pattern match queries, and beyond.

5.2.1. Prerequisites

Our model supports not only structural requirements, but also attribute predicates and reachability. In particular, our queries contain nodes and edges that are associated to *attribute predicates* and...
edges that are associated to *reachability expressions*. In this section we formalize these concepts.

An *attribute predicate* is a set of conditions on the attributes established for a node or an edge in a graph query. Specifically, an attribute predicate is either called a *node predicate* or an *edge predicate* depending on whether it is associated to a node or an edge, respectively. We formalize this definition as follows:

**Definition 6 (Attribute Predicate).** An attribute predicate on a node (or edge) of an attributed multigraph is an expression, with conjunctions and disjunctions, that evaluates some conditions on its attributes. Let us denote as $u\text{.pred}$ the predicate associated to the node (or edge) $u$. An attribute predicate $u\text{.pred}$ associated to the node (or edge) $u$ is drawn from the following grammar:

\[
P \rightarrow \text{Attribute Op Constant} \mid P \land P \mid P \lor P \\
\text{Op} \rightarrow < \mid \leq \mid > \mid \geq \mid = \mid \neq
\]

A node/edge $v$ in the attributed multigraph satisfies the predicate $u\text{.pred}$, denoted as $v \sim u$, if $u\text{.pred}$ is true when evaluated with $v$’s attributes.

Furthermore, our model generalizes the traditional definition of reachability in graphs — we introduce the concept of $(u, v, \rho)$-*reachability*. Here $u$ and $v$ represent nodes associated to node predicates and $\rho$ represents an edge that is associated to a *reachability expression expression*, which we define next.

**Definition 7 (Reachability Expression).** Let $e$ and $n$ represent edges and nodes that are associated to predicates, respectively. We define a reachability expression as an expression defined over $Q$:

\[
Q \rightarrow e \mid Q(nQ)^* \mid (Qn)^*Q \mid Q \cup Q
\]

A reachability expression is associated to an edge in our query. We denote it as $\rho\text{.re}$ when it is associated to edge $\rho$.

Note that a reachability expression represents a reachability relation. Hence, the corresponding edge can be mapped to a path with arbitrary length of the attributed multigraph.

**Definition 8 ((u, v, $\rho$)-Reachability).** Let us consider the nodes $u$ and $v$ associated to predicates, and an edge $\rho$ associated to a reachability expression. Given an attributed multigraph $G = (V, E, f_V, f_E)$ and two nodes $u', v' \in V$, $v'$ is said to be $(u, v, \rho)$-reachable from $u'$ iff:

1. $u' \sim u$;
2. $v' \sim v$; and
3. there exists a path drawn from $\rho\text{.re}$ that connects $u'$ and $v'$.

Note that a reachability expression $\rho\text{.re}$ must be an alternating sequence of edges and nodes associated to predicates where the Kleene star and union operators are supported. We use $W_G(u, v, \rho)$
to represent the set of all paths in \( G = (V, E, f_V, f_E) \) that connect any \( u' \in V \) to any \( v' \in V \) such that \( v' \) is \((u, v, \rho)\)-reachable from \( u' \).

In order to determine whether a particular \( v' \in V \) is \((u, v, \rho)\)-reachable from a given \( u' \in V \), we can either match forward from \( u \) to \( v \) through \( \rho.re \) (forward approach), or match from \( v \) to \( u \) backward through \( \rho.re^R \), which is the reversal of the reachability expression \( \rho.re \) (backward approach). In the forward approach, outbound edges of the nodes in the attributed multigraph are considered; in the backward approach, inbound edges of the nodes are traversed. Table 5-1 shows how to obtain the reversal of different reachability expressions drawn from the context-free grammar \( Q \) under different rules of production (see Definition 7).

**Table 5-1:** Reversal of reachability expressions drawn from \( Q \) under different rules of production (see Definition 7).

<table>
<thead>
<tr>
<th>( \rho.re )</th>
<th>( \rho.re^R )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( e )</td>
<td>( e )</td>
</tr>
<tr>
<td>( Q(nQ)^* )</td>
<td>( (Q^Rn)^Q^R )</td>
</tr>
<tr>
<td>( (Qn)^*Q )</td>
<td>( Q^R(nQ^R)^* )</td>
</tr>
<tr>
<td>( Q \cup Q )</td>
<td>( Q^R \cup Q^R )</td>
</tr>
</tbody>
</table>

### 5.2.2. Definition of Generalized Pattern Queries

Now, we are ready to define generalized pattern queries and their matches:

**Definition 9** (Generalized Pattern Query). A generalized pattern query \( G_P = (V_P, E_P) \) is a weakly connected directed multigraph\(^1\) where:

- \( V_P \) is a set of nodes associated to node predicates.
- \( E_P \) is a multiset of edges, on \( V_P \times V_P \), that are associated to reachability expressions.
- Each edge \( e' = (u, v) \in E_P \) establishes a \((u, v, e')\)-reachability requirement.

A match of a generalized pattern query \( G_P = (V_P, E_P) \) is a set of nodes in the attributed multigraph that bijectively satisfy the predicates associated to the nodes in \( V_P \). Furthermore, each pair of nodes in the match must satisfy the reachability requirements established for the edges that connect their corresponding nodes in \( G_P \). Hence, our model considers structural requirements, predicates and reachability. We present a more formal definition next.

\(^1\)A directed multigraph is weakly connected if replacing all its directed edges with undirected edges produces a connected (undirected) multigraph.
Definition 10 (Match of a Generalized Pattern Query). Let the generalized pattern query $G_P = (V_P, E_P)$, where $V_P = \{u_1, u_2, \ldots, u_n\}$ and $n = |V_P|$. Also let $G = (V, E, f_V, f_E)$ be an attributed multigraph and $V' \subseteq V$. The tuple $(v_1, \ldots, v_n)$, for $v_i \in V'$ and $1 \leq i \leq n$, is a match of $G_P$ in $G$ iff there exists a bijective function $f : V_P \rightarrow V'$ such that:

1. $v_i = f(u_i)$ and $v_i \sim u_i$ for every $1 \leq i \leq n$.
2. $f(u_j)$ is $(u_i, u_j, e)$-reachable from $f(u_i)$ for every $e = (u_i, u_j) \in E_P$, $1 \leq i, j \leq n$.

We say that $(v_1, \ldots, v_n)$ is a match of $G_P$ in $G$ under $f$.

In this thesis, we study the next problem:

Problem 4 (Generalized Pattern Query Problem (GPQP)). Given a generalized pattern query $G_P = (V_P, E_P)$ and an attributed multigraph $G = (V, E, f_V, f_E)$, the Generalized Pattern Query Problem (GPQP) consists of finding the set $G_P(G)$ of all the matches of $G_P$ in $G$, i.e.,

$$G_P(G) = \{(f_k(u_1), \ldots, f_k(u_n)) \mid f_k \in \mathcal{M}_{G_P \rightarrow G}\},$$

where $\mathcal{M}_{G_P \rightarrow G}$ denotes the set of mapping functions $f_k : V_P \rightarrow V'$, for a set $V' \subseteq V$, that yield matches of $G_P$ in $G$.

5.2.3. Example

An example of a generalized pattern query is presented in Figure 5-2(a). In particular, the query searches for photos in which Alice is tagged with two people: one of her colleagues and another female; furthermore, the colleague must be within Mary’s network (they must be connected through friend-edges). Figure 5-2(b) shows the output of this query on the attributed multigraph of Figure 5-1. In this case, all the edges except $e_1$ represent single edge predicates; thus their matches are edges in the attributed multigraph. As for $e_1$, we can see in Figure 5-1 that there is a path between Mary and Nick that satisfies the regular expression $([\text{type} = \text{friend}][\text{type} = \text{person}])^* [\text{type} = \text{friend}]$. The support of predicate attributes on both nodes and edges, as well as the expressive power of the reachability expressions, allows us to construct many useful and complex queries.

5.2.4. Discussion

Generalized pattern queries possess rich expressive power. It is a strict superset of reachability queries [40, 151, 84, 127, 128] and pattern match queries [159, 161]. We can use it to express reachability queries, pattern match queries, and their combination. In particular, a reachability query (RQ) is a special case of generalized pattern queries that only has two nodes (associated to predicates) and one edge (associated to a reachability expression). While a RQ establishes a single reachability requirement, generalized pattern queries allow multiple $(u, v, \rho)$-reachability
5.2 Query Model

requirements that are co-related. We support reachability queries with attribute predicates on both intermediate edges and nodes, which had not been supported by previous models.

For example, if we want to know in which photos there are men within Alice’s network, we can use a query like the one in Figure 5-3(a). The output of the query on the multigraph of Figure 5-1 includes (Alice, Photo1) and (Alice, Photo2). The set of the connecting paths associated to these results is presented in Figure 5-3(b). Furthermore, if we also want to retrieve the names of the men (tagged in the photos), we can use the query in Figure 5-4(a). The output of such query on the graph of Figure 5-1 is composed by (Alice, Nick, Photo1) and (Alice, Nick, Photo2).

A pattern match query, associated to subgraph isomorphism, is also a special case of generalized pattern queries where \( e.re \), for all \( e \in E_P \), corresponds to a single edge predicate. For instance, let us consider the query in Figure 5-5: it finds the friends of Alice that are tagged with her in a photo and also retrieve the photos. The output of this query on the graph of Figure 5-1 is presented in Figure 5-5(b). This result is a subgraph of the attributed multigraph that is isomorphic to the query and satisfies all the predicates.

Generalized pattern queries can also represent queries that cannot be modelled by either reachability or subgraph isomorphism queries. Specifically, generalized pattern queries are graphs that establish the predicates on a set of nodes of interest (through node predicates) and the reachability requirements among them (through highly-expressive regular expressions associated to the edges).
Generalized Pattern Queries

\[
W_G(u, v, \rho) = \{\langle Alice, friend, Mike, friend, Rose, friend, Nick, tag, Photo 1 \rangle, \\
\langle Alice, friend, Mike, friend, Rose, friend, Nick, tag, Photo 2 \rangle\}
\]

**Figure 5.3:** Example of a reachability query expressed as a generalized pattern query. (a) Query. (b) Set of the connecting paths associated to the output of this query on the graph of Figure 5.1.

\[
e_1.re = \langle friend - [type=person]* - friend \rangle \quad e_2.re = \langle [tag] \rangle
\]

**Figure 5.4:** Example of a reachability query with intermediate nodes of interest expressed as a generalized pattern query.

We show an example in Figure 5.2(a); this query cannot be expressed by either of the previous query models. In the next sections, we present an algorithm that finds all the matches of a generalized pattern query; hence, this algorithm also solves reachability queries, pattern match queries, and more.
6. Linearization on Generalized Pattern Queries

In this chapter, we extend the concept of linearization presented in Section 3.1 to represent generalized pattern queries in a linear manner. We then define match of a query linearization against the attributed multigraph, and show that each match of the query linearization corresponds to a match of our query. Therefore, processing our query requires two steps: (1) linearize the query, and (2) find the matches of the query linearization. In Section 6.1, we present the main definitions and prove the correctness of our approach. Then, in Section 6.2, we propose an algorithm that takes into account the attributed multigraph statistics to produce a query linearization that entails low time requirements during the matching phase.

6.1. Query Linearization

In this section, we present the definition of query linearization and how it can be used to solve the generalized pattern queries.

Definition 11 (Query Linearization). Let $G_P = (V_P, E_P)$ be a generalized pattern query. An undirected walk $p = p_1\ldots\ell$ on $G_P$ is a query linearization of $G_P$ iff:

1. $p_i$ is a node $v \in V_P$ if $i$ is odd, $1 \leq i \leq \ell$.

2. $p_i$ is an edge $e \in E_P$ if $i$ is even, $1 \leq i \leq \ell$, such that either $e = (p_{i-1}, p_{i+1})$ or $e = (p_{i+1}, p_{i-1})$.

3. Each node $v \in V_P$ and each edge $e \in E_P$ appears at least once in $p$.

Notice that this definition is similar to Definition 2. The difference is that, as the graph is a generalized pattern query, the nodes are associated to attributes and each edge defines a $(u, v, \rho)$-reachability requirement. Then, the objective of a query linearization $p$ of $G_P$ is representing the structure, the reachability requirements and the attribute predicates of $G_P$ in a linear manner. We use an alternating sequence of adjacent nodes (associated to predicates) and edges (associated to reachability expressions) that starts and ends at a node. All the nodes and edges in $G_P$ must appear in $p$ at least once so that the complete set of adjacency relations in $E_P$ is represented.
We use undirected walks, as there may not exist a directed walk including all the nodes and edges. However, the direction of each edge $e \in E_P$ is denoted in the query linearization as $e$.direction and is considered during the matching phase. Namely, for the edges $p_i$ in $p$, $p_i$.direction = forward if $p_i = (p_{i-1}, p_{i+1})$ and $p_i$.direction = backward if $p_i = (p_{i+1}, p_{i-1})$, for even values of $i$ in $1 \leq i \leq \ell$. We refer to them as forward and backward edges, respectively. A forward edge $p_i$ establishes a $(p_{i-1}, p_{i+1}, p_i)$–reachability requirement; similarly, a backward edge $p_i$ establishes a $(p_{i+1}, p_{i-1}, p_i)$–reachability requirement, for even values in $1 \leq i \leq \ell$. Then, the forward and backward edges are matched in the attributed multigraph using the forward and backward approach, respectively.

There may be more than one query linearization that represents the same generalized pattern query. Figure 6-1 depicts an example of two possible linearizations for the query in Figure 5-2(a).

![Figure 6-1](image)

**Definition 12 (Match of a Query Linearization).** Consider the generalized pattern query $G_P = (V_P, E_P)$ and an attributed multigraph $G = (V, E, f_V, f_E)$. Also, let $p = p_1...p_\ell$ be a query linearization of $G_P$ and $q = q_1...q_k$ be an undirected walk on $G$ where $k \geq \ell$. Furthermore, let $q'$ be a subsequence of nodes in $q$ such that $q' = (q_1', q_3', ..., q_{\ell}' = (q_{s_1}, q_{s_3}, ..., q_{s_\ell})$ where $s_1 = 1$, $s_\ell = k$ and $s_i < s_{i+2}$ for odd values of $i$ in $1 \leq i \leq \ell$. Then, the subsequence $q'$ of nodes is a match of the query linearization $p$ iff:

1. There exists a bijective mapping function $f : V_P \rightarrow V'$, where $V'$ is the set of nodes in the subsequence $q'$, such that $q_{s_i} = f(p_i)$ and $q_{s_i} \sim p_i$, for odd values of $i$, $1 \leq i \leq \ell$.  

Because processing a walk is simpler than processing a graph, we use query linearization to solve the generalized pattern queries problem by comparing undirected walks of $G$ with a query linearization $p$. Then, we define match of a query linearization:
2. \( q_{s_{i+2}} \) is \((p_i, p_{i+2}, p_{i+1})\)-reachable from \( q_s \), for odd values of \( i \), \( 1 \leq i < \ell \), for which \( p_{i+1}.direction = \text{forward} \). We denote this as \( q_s \xrightarrow{p_i \ p_{i+2}} q_{s_{i+2}} \).

3. \( q_s \) is \((p_{i+2}, p_i, p_{i+1})\)-reachable from \( q_{s_{i+2}} \), for odd values of \( i \), \( 1 \leq i < \ell \), for which \( p_{i+1}.direction = \text{backward} \). We also denote this as \( q_s \xleftarrow{p_{i+2} \ p_i} q_{s_{i+2}} \).

We say that \( q' \) is a match of \( p \) in \( G \) under function \( f \).

Notice that a match \( q' = (q_{s_1}, q_{s_3}, \ldots, q_{s_k}) \) is a subsequence of nodes contained in an undirected walk \( q \) on \( G \); such nodes are connected in an manner that satisfies the structural constraints, the predicates and the reachability requirements associated to the query linearization \( p = p_1...\ell \).

Specifically, Condition (1) validates the predicates established by the nodes in the query linearization and their bijective association with the nodes in \( q' \). Furthermore, Conditions (2) and (3) validate the reachability requirements established by the edges in the linearization. It is important to remark that for edges whose direction is \textit{backward}, the reversal of its associated reachability expression must be considered. An example of a match of the query linearization \( p = p_1...13 \) of Figure 6-1(a) on the attributed multigraph of Figure 5-1 is presented in Figure 6-2. Specifically, the match \( q' = (Mary, Nick, Alice, Photo2, Nick, Photo2, Kate) \) is contained in the undirected walk \( q = q_1...19 \).

![Figure 6-2](image)

\textbf{Figure 6-2.} \( q' = (Mary, Nick, Alice, Photo2, Nick, Photo2, Kate) \) is a match of the the linearization \( p = p_1...13 \) presented in Figure 6-1(a). It is equivalent to finding a match \((Mary, Nick, Alice, Photo2, Kate) \) of \( G_P \) in Figure 5-2(a).

The core idea of our approach to find all the matches of the query \( G_P \) in \( G \) is as follows. Let \( p \) be a query linearization of \( G_P \). Recall that, \( p \) represents the structural requirements, the reachability requirements and the predicates of \( G_P \). If a subsequence \( q' \) of nodes in \( G \) is a match of \( p \), then \( q' \) satisfies the requirements and predicates established in \( p \) (and, hence, in \( G_P \)); thus, \( q' \) is an
occurrence of $G_P$ in $G$. More formally, the following theorem allows us to find the matches of a
generalized pattern query in an attributed multigraph by means of query linearization.

**Theorem 6.** Consider the generalized pattern query $G_P = (V_P, E_P)$, where $V_P = \{u_1, u_2, \ldots, u_n\}$,
and an attributed multigraph $G = (V, E, f_V, f_E)$. Also, let $p = p_1, \ldots, \ell$ be a query linearization of
$G_P$ and $V' \subseteq V$. Then, the subsequence $q'$ of nodes is a match of $p$ under function $f : V_P \rightarrow V'$,
where $V'$ is the set of nodes contained in $q'$, if and only if $(f(u_1), \ldots, f(u_n))$ is a match of $G_P$
under bijective mapping function $f$ in $G$.

**Proof.** Let us consider the attributed multigraph $G = (V, E, f_V, f_E)$, the generalized pattern query
$G_P = (V_P, E_P)$ and the query linearization $p = p_1, \ldots, \ell$ of $G_P$. Furthermore, $V_P = \{u_1, u_2, \ldots, u_n\}$
and $n = |V_P|$. Also, let $V'$ be a subset of $n$ nodes in $G$ ($V' \subseteq V$). In order to prove the theorem,
we need to show that (i) if $q'$ is a match of $p$ in $G$ under the bijective function $f : V_P \rightarrow V'$, then
$(f(u_1), \ldots, f(u_n))$ is a match of $G_P$ in $G$ under $f$; and (ii) if $(f(u_1), \ldots, f(u_n))$ is a match of $G_P$
in $G$ under function $f : V_P \rightarrow V'$, then $q'$ is a match of $p$ in $G$ under $f$.

First we prove (i). According to Definition 12, if $q'$ is a match of $p$ in $G$, there exists a bijective mapping function $f : V_P \rightarrow V'$ such that $q_{s_i} = f(p_i)$ and $q_{s_i} \sim p_i$, for odd values of $i$, $1 \leq i \leq \ell$ (condition (1) of Definition 12). Given that $p$ contains all the nodes in $V_P$ (see Definition 11), condition (1) of Definition 10 is satisfied. Furthermore, because $p$ also contains all the adjacency relations of $G_P$, conditions (2) and (3) of Definition 12 imply that for every pair $e = (u_1, u_2) \in E_P$, $f(u_2)$ is $(u_1, u_2, e)$–reachable from $f(u_1)$ (condition (2) of Definition 10).

Now we prove (ii). Let us consider $V' = \{v_1, v_2, \ldots, v_n\}$. According to Definition 10, if $(v_1, \ldots, v_n)$ is a match of $G_P$ in $G$, then there exists a bijective mapping function $f : V_P \rightarrow V'$ such that $v_i = f(u_i)$ and $v_i \sim u_i$, for all $1 \leq i \leq n$ (condition (1) of Definition 10). Then, condition (1) of Definition 12 is satisfied. Moreover, for every pair $e = (u_1, u_2) \in E_P$, $f(u_2)$ is $(u_1, u_2, e)$–reachable from $f(u_1)$ (condition (2) of Definition 10). Given that all the edges in $E_P$ are contained at least once in $p$ (see Definition 11), for each $e = (u_i, u_j) \in E_P$, there exists at least a walk $\langle f(u_i), \ldots, f(u_j) \rangle \in W_G(u_i, u_j, e)$. Therefore, conditions (2) and (3) of Definition 12 are satisfied as well.

Notice that this theorem is based on the following facts: (i) a query linearization represents all the
adjacency relations of the corresponding generalized pattern query; and (ii) the matches of both the
generalized pattern query and the query linearization involve the existence of a bijective mapping
of the nodes that satisfies the node predicates and the reachability requirements.

In other words, this theorem states that the matches of the generalized pattern query $G_P =
(V_P, E_P)$ in the attributed multigraph $G = (V, E, f_E, f_E)$ are defined with the same bijective
mapping functions of the matches of $p$ in $G$, where $p$ is a query linearization of $G_P$. Each match of
6.2 Enhanced Graph Linearization Algorithm — E-GLA

$p$ in $G$ is associated to a match of $G_P$ in $G$ under a common function $f$. For example, for the attributed multigraph in Figure 5-1, the mapping function of both $p$ and $G_P$ (presented in Figures 6-2 and 5-2(b), respectively) are associated to the same mapping function:

$$f : (u_1, u_2, u_3, u_4, u_5) \rightarrow (Mary, Nick, Alice, Photo2, Kate)$$

Then, finding the matches of $G_P$ is equivalent to finding the matches of $p$ in $G$. Based on this, our approach to find the matches of a generalized pattern query consists of two steps: computing a linearization $p$ of $G_P$ and finding the matches of such linearization in $G$. We describe each of these steps in the next sections.

6.2. Enhanced Graph Linearization Algorithm — E-GLA

In this section, we present an algorithm for identifying a query linearization with low matching cost, out of many possible linearizations, as illustrated in Figure 6-1. In particular, our algorithm is called Enhanced-Graph Linearization Algorithm (E-GLA).

6.2.1. Baseline: GLA for Length-Optimality

As a baseline, we consider GLA (Graph Linearization Algorithm), which was presented in Section 3.3. This algorithm aims at selecting a length-optimal linearization. For example, Figure 6-1(a) is a length-optimal linerization: a linearization with minimum length among all possible linearizations (see Definition 4). As exact computation incurs prohibitive cost, GLA implements simple heuristics on a depth-first search (DFS) of the query graph. Specifically, the search can be represented as a DFS tree. The main heuristics of GLA is to cover the nodes in the highest levels of the DFS tree to keep the depth minimal; this is done by prioritizing visits to the nodes with minimum number of unexplored edges when there are choices. Its heuristics achieve $2$-approximate optimality (see Theorem 3).

Though length-optimality is a reasonable cost model for the isomorphism problem, this is not as effective in a more complex query model like ours where selectivity varies. Specifically, it is important to take into account that we match the query linearization against the attributed multigraph using a depth-first search (DFS), where the top levels of the DFS tree correspond to the first elements of the query linearization. Then, we want to put the predicates with fewer matches at the beginning of the query linearization in order to prune the search space by detecting mismatches at an early stage of the process.

To illustrate this, let us consider the generalized pattern query of Figure 5-5 and the attributed multigraph of Figure 5-1. A query linearization can be started from any of the nodes in the generalized pattern query; however, $u_1$, $u_2$ and $u_3$ have 1, 6 and 3 matches during the matching phase,
respectively. Therefore, we start the query linearization from $u_1$ in order to consider only one DFS search tree in the matching phase. For the next steps, let us consider the following statistics: (i) the average number of photos a person is tagged in is 1,16; and (ii) the average number of friends a person has is 1,66. These statistics are obtained by simple counting on the attributed multigraph of Figure 5-1. Hence, it is better to continue the query linearization with $u_3$ as it is likely to lead to fewer matches in the second level of the DFS search trees of the matching phase. Thus, the query linearization that most likely reduces the search space is $u_1e_3u_3e_2u_2e_1u_1$. We already observe the cost benefit of using statistics in this example even though the graph is tiny. It becomes much more important when the graph size increases, as different query linearizations may incur orders of significantly different costs.

### 6.2.2. Key Ideas

In this section, we develop the key ideas of the Enhanced-Graph Linearization algorithm (E-GLA). The cost of the matching phase is represented as DFS search trees where the roots are the possible matches of $p_1$. The goal of E-GLA is pruning such trees by placing the nodes in $V_P$ with fewer matches in $G$ (i.e., nodes with low selectivity) at the beginning of the query linearization $p$. As it was mentioned in last section, this causes that such few matches are placed at the top levels of the DFS search trees during the matching phase. Hence, the search space is pruned at an early stage by avoiding partial matches of $p$ that will not lead to complete matches.

More formally, we define the **selectivity** for each node $u \in V_P$ in the generalized pattern query. The selectivity of the node $u$ with respect to the attributed multigraph $G = (V, E, f_V, f_E)$ is denoted as $\text{select}_G(u)$. It is calculated as the probability of selecting a node from $G$ that satisfies the predicate $u_{\text{pred}}$. In this sense, E-GLA gives preference to the nodes in the query with the lowest selectivity for starting (or continuing) the linearization.

The traversal performed by E-GLA on the generalized pattern query is a DFS search with the following heuristics: (1) the traversal starts from the node with the lowest selectivity; (2) the unexplored edges that lead to already explored nodes are visited before than the ones that lead to unexplored nodes; and (3) the edges that lead to unexplored nodes are considered sorted, in ascending order, on the selectivity of such nodes. Notice that heuristics (1) and (3) aim to optimize for selectivity, by placing the nodes of the generalized pattern query that will have many mismatches at the beginning of the query linearization; this helps to prune the search space at an early stage of the matching phase. Moreover, heuristic (2) optimizes for the length of the linearization as in GLA. Thus, E-GLA produces a linearization that not only considers the length but also takes into account the statistics of the attributed multigraph to reduce the matching-phase time.
6.2 Enhanced Graph Linearization Algorithm — E-GLA

6.2.3. Algorithm

The pseudocode in Figure 6-3 describes E-GLA. Each node and edge has a boolean flag that indicates whether it has been explored. All these flags are initialized with false. Furthermore, the number of unexplored nodes/edges is stored in variable unexplVE. This variable is used to avoid reinserting graph elements in the backtracking of the DFS search tree when there are no unexplored elements left. The query linearization is represented as a list p.

Notice that the selectivity of each node v, denoted as v.selectivity, is populated by means of the function COMPUTESELECTIVITY(). The parameters of this function are the corresponding node predicate v.pred and the attributed multigraph \( G = (V, E, f_v, f_e) \). This function can be implemented in different ways: (i) counting the nodes in the attributed multigraph that satisfy v.pred; (ii) using previous information on the attributed multigraph statistics; or (iii) designing formulas based on the distribution of the node attributes. Thus, the complexity of this function can vary from constant to linear on the number of nodes in the attributed multigraph.

With such selectivity information, we find the node u with lowest selectivity (line 5, Figure 6-3). Then, the DFS traversal is performed by calling the recursive procedure STATSRAVERSE() over node u (see Figure 6-4). This procedure is essentially the same as TRAVERSEGRAPH() (the recursive procedure that performs the traversal for GLA); the difference is that edges that lead to unexplored nodes are sorted, in ascending order, on their selectivity (lines 7–8, Figure 3-2). The algorithm terminates when the first call to STATSRAVERSE() finishes.

Algorithm 7: E-GLA Algorithm

<table>
<thead>
<tr>
<th>Input:</th>
<th>( G_P = (V_P, E_P) ), ( G = (V, E, f_v, f_e) )</th>
<th>Output:</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. for every ( e \in E_P ) do ( e.\text{Explored} \leftarrow \text{false} )</td>
<td></td>
<td>2. for every ( v \in V_P ) do</td>
<td></td>
</tr>
<tr>
<td>3. ( v.\text{Explored} \leftarrow \text{false} )</td>
<td></td>
<td>4. ( v.\text{selectivity} \leftarrow \text{ComputeSelectivity}(v.\text{pred}, G) )</td>
<td></td>
</tr>
<tr>
<td>5. choose ( u \in V_P ) with ( \min(u.\text{selectivity}) )</td>
<td></td>
<td>6. ( p \leftarrow \langle \rangle ), unexplVE \leftarrow</td>
<td>V</td>
</tr>
<tr>
<td>7. StatsTraverse(( G_P, u, p, \text{unexplVE} ))</td>
<td></td>
<td>8. return ( p )</td>
<td></td>
</tr>
</tbody>
</table>

Figure 6-3.: E-GLA algorithm.

Example. To illustrate how E-GLA works, Table 6-1 shows the selectivity of the nodes in the query of Figure 5-2(a). This can be obtained from the statistics on the attributed multigraph of Figure 5-1, e.g., the number of nodes that satisfy each predicate divided by the total number of...
Algorithm 8: \textsc{StatsTraverse()} Procedure

\begin{algorithmic}[1]
\Procedure{StatsTraverse}{$G_P = (V_P, E_P), u, p, \text{unexplVE}$}
\State $p.\text{Add}(u)$, $u.\text{Explored} \leftarrow \text{true}$, $\text{unexplVE} \leftarrow -$
\For{every $e \in E$ such that $e = (u, v)$ or $e = (v, u)$}
\If{$!e.\text{Explored} \land v.\text{Explored}$}
\State $p.\text{Add}(e)$, $e.\text{Explored} \leftarrow \text{true}$, $\text{unexplVE} \leftarrow -$, $p.\text{Add}(v)$
\EndIf
\EndFor
\If{$\text{unexplVE} > 0$}
\State $p.\text{Add}(e)$, $p.\text{Add}(u)$
\EndIf
\While{there are unexplored edges $e = (u, v)$ or $e = (v, u)$}
\State \begin{varitemize}
\item \textbf{choose} $e$ with $\min(v.\text{selectivity})$
\end{varitemize}
\State $p.\text{Add}(e)$, $e.\text{Explored} \leftarrow \text{true}$, $\text{unexplVE} \leftarrow -$
\State \text{StatsTraverse($G_P, v, p, \text{unexplVE}$)}
\If{$\text{unexplVE} = 0$} \textbf{break} \EndIf
\State $p.\text{Add}(e)$, $p.\text{Add}(u)$
\EndWhile
\EndProcedure
\end{algorithmic}

\textbf{Figure 6-4.} \textsc{StatsTraverse()} procedure.

nodes. E-GLA may start the linearization from the nodes with lowest selectivity, namely $u_1$ and $u_3$: if the starting node is $u_1$, the linearization obtained is the one presented in Figure 6-1(a); if the starting node is $u_3$, the linearization is slightly longer as presented in Figure 6-1(b).

\textbf{Table 6-1.} Selectivity of the nodes in the generalized pattern query of Figure 5-2(a) with respect to the attributed multigraph of Figure 5-1.

<table>
<thead>
<tr>
<th>Node</th>
<th>$u_1$</th>
<th>$u_2$</th>
<th>$u_3$</th>
<th>$u_4$</th>
<th>$u_5$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Selectivity</td>
<td>1/9</td>
<td>2/3</td>
<td>1/9</td>
<td>1/3</td>
<td>4/9</td>
</tr>
</tbody>
</table>

6.2.4. Correctness Proof

The correctness of this algorithm is proven in the following theorem:

\textbf{Theorem 7.} The Enhanced-Graph Linearization Algorithm (E-GLA) outputs a query linearization of the input generalized pattern query $G = (V_P, E_P)$.

\textbf{Proof.} Let $G = (V_P, E_P)$ be a generalized pattern query and $p = p_1...\ell$ the output walk produced by E-GLA. Notice that, disregarding the predicates associated to the nodes in $V_P$ and the reachability expressions associated to the edges in $E_P$, $G_P$ is a regular graph. Similarly, a query linearization of $G_P$, without the node predicates and the reachability expressions of the edges, is a simple linearization (as defined in Definition 2). Then, we prove Theorem 7 by showing that the
Let us recall that the recursive procedure `STATESTRAVERSE()` is essentially the same as the recursive procedure `TRAVERSEGRAPH()`. These procedures are in charge of the DFS traversal performed by GLA and E-GLA, respectively. Their only difference is the order in which adjacent edges that lead to unexplored nodes are considered. Hence, even though the traversal performed by E-GLA can be different from the one of GLA, such traversal maintains the same properties. Namely, conditions (1), (2) and (3) of Definition 2 are satisfied due to Theorem 2. That is, the output walk \( p \) is a linearization of \( G_p \). Thus, if the corresponding predicates and reachability expressions are respectively included in the nodes and edges of the walk \( p \), then such walk is a *query linearization* of \( G_p \).

**6.2.5. Length of E-GLA Linearization**

In this section, we formally discuss the strength of E-GLA. Theorem 8 shows that given the generalized pattern query \( G_p = (V_p, E_p) \), the length of the linearization generated by E-GLA is at most 2 times the length of a length-optimal linearization. That is, E-GLA optimizes for selectivity, yet gives the same asymptotic length guarantee as GLA.

**Theorem 8.** E-GLA is 2-approximate with respect to the length of a length-optimal linearization.

**Proof.** Let \( G_p = (V_p, E_p) \) be a generalized pattern query and \( p = p_1 \ldots \ell \) the output walk produced by E-GLA. Any query linearization algorithm, including length-optimal algorithms, must traverse each edge of the multigraph at least once. Thus, the number of edges in a query linearization of \( G_p \) is at least \( |E_p| \). Since a linearization has the format of alternating between nodes and edges, a query linearization with \( k \) edges has \( k + 1 \) nodes. Hence, the optimal query linearization \( p^* \) has at least \( |E_p| \) edges and \( |E_p| + 1 \) nodes. Therefore, \( |p^*| \geq 2|E_p| + 1 \).

Like GLA, E-GLA also visits any edge at most twice during the DFS traversal. This is because, when procedure `STATESTRAVERSE()` is executed over node \( u \), an unexplored edge \( e \) that leads to any explored or unexplored node \( v \) is added once into \( p \) (lines 4 and 9, respectively, Figure 6-4). If after executing the next instructions there are still unvisited graph elements, it is necessary to go back to \( u \) through \( e \); this means that \( e \) and \( u \) are added into \( p \) again (lines 6 and 12, Figure 6-4). After this, \( e \) is not visited ever again given that only unexplored edges are considered (lines 3 and 7, Figure 6-4). Therefore, the number of edges in the query linearization is at most \( 2|E_p| \). Again, since a linearization has the format of alternating between nodes and edges, the query linearization \( p^{E-GLA} \) has at most \( 2|E_p| \) edges and \( 2|E_p| + 1 \) nodes. Therefore, we have \( |p^{E-GLA}| \leq 4|E_p| + 1 \). It leads to the approximation ratio of E-GLA,

\[
\frac{|p^{E-GLA}|}{|p^*|} \leq \frac{4|E_p| + 1}{2|E_p| + 1} \leq 2.
\]
This worst-case approximation ratio is obtained by comparing the length of E-GLA linearizations with a lower bound where it is assumed that each edge in the query appears only once in the query linearization. However, even a length-optimal linearization may not achieve this lower bound for many graph structures, because it may require including an element more than once. Thus, for many cases, in practice, E-GLA linearizations are much closer to length-optimal linearizations than the ratio given by this theorem.

6.2.6. Complexity Analysis

This section studies the computational complexity of E-GLA to generate a query linearization \( p = p_1...p_\ell \) of the generalized pattern query \( G_P = (V_P, E_P) \) for the attributed multigraph \( G = (V, E, f_V, f_E) \), where \( n = |V| \) and \( m = |E| \).

**Time Complexity.** Initializing the edges and nodes as unexplored takes \( O(|E_P|) \) and \( O(V_P) \), respectively (lines 1–3, Figure 6-3). As discussed in Section 6.2.3, the time complexity for computing the selectivity of each node can vary from constant to linear on \( n \). In this analysis, we assume that we have information on the distribution of the node attributes; thus, the cost of executing this procedure is \( O(1) \) per node and, hence, \( O(|V_P|) \) for all nodes in \( G_P \) (line 4, Figure 6-3). Finding the node with lowest selectivity takes \( O(|V_P|) \) (line 5, Figure 6-3). Initializing \( p \) and unexpVE takes constant time (line 6, Figure 6-3).

Notwithstanding, the time complexity of E-GLA is dominated by the undirected walk traversed (line 7, Figure 6-3) which corresponds to the length of the query linearization. In Section 6.2.5, we showed that \( p \) has at most \( 2|E_P| \) edges and \( 2|E_P| + 1 \) nodes. Each insertion takes constant time as it is always done at the end of \( p \). But when a node is inserted for the first time, it is necessary to consider the unexplored adjacent edges \( e \) that lead to unexplored nodes \( v \) sorted on their selectivity (lines 7–8, Figure 6-4). This sorting operation takes \( O(d \lg d) \), where \( d \) is the maximum degree of the nodes in \( G_P \) (including both incoming and outgoing edges). Thus, the time complexity of E-GLA is \( O(2|E_P| + |V_P|(d \lg d)) = O(|E_P| + d|V_P| \lg d) \).

**Space Complexity.** The space complexity is given by the list that stores the linearization, i.e., by the length of the linearization. Because the linearization can have at most \( 2m \) edges and \( 2m + 1 \) nodes, the total space complexity is \( \Theta(m) \).
7. Solution of Generalized Pattern Queries

We develop a matching algorithm for generalized pattern queries, which we call GPQM. This algorithm uses a query linearization $p = p_1 \ldots \ell$ of the generalized pattern query $G_P = (V_P, E_P)$ to find the matches of $G_P$ in the attributed multigraph $G = (V, E, f_V, f_E)$. In Section 7.1, the high-level ideas of the algorithm are presented. Then, we go through the details in Section 7.2. Its correctness is proven in Section 7.3 while the complexity analysis is derived in Section 7.4. Finally, some experimental results are presented in Section 7.5.

7.1. Key Ideas

The GPQM algorithm searches for matches of the query linearization $p = p_1 \ldots \ell$ of the generalized pattern query $G_P = (V_P, E_P)$ in the attributed multigraph $G = (V, E, f_V, f_E)$. According to Theorem 6, each match represents a match of $G_P$ in $G$. This section presents the key ideas of the algorithm.

We search all the possible undirected walks in $G$ that may contain matches of a linearization $p = p_1 \ldots \ell$ in a DFS manner. Our search starts from $p_1$: the roots of the candidate DFS search trees are the nodes that satisfy $p_1$. Then, we try to associate each node $p_i$ to every node $u$ in these DFS search trees that satisfies both the node predicates and the reachability requirements. Each undirected walk from the root of the search tree to $u$ is associated to an injective function $f$ that establishes a mapping between the nodes in the linearization and certain nodes on the walk. The injective property ensures that two different nodes in $p$ are not assigned to the same node $u \in V$; likewise, two different nodes in $V$ are not associated to the same node in $p$.

More specifically, the query linearization $p = p_1 \ldots \ell$ is traversed from $p_1$ to $p_\ell$ while we consider the possible assignments for each $p_i$, $1 \leq i \leq \ell$ in the graph $G$. Let us assume that $f(p_i) = u$ was set. We recursively extend the current partial match under $f$ where there are two cases:

**Case 1:** Node $p_{i+2}$ is unassigned: We consider all the possible assignments $f(p_{i+2}) = v$ for all the nodes $v \in V$ such that $u \overset{p_i+p_{i+2}}{\longrightarrow} v$ (or $u \overset{p_i+p_{i+2}}{\longleftarrow} v$, if $p_{i+1}$ is a backward edge). Let us establish the predicate $p'_i,\text{pred} : [ID = u.id]$. Because $u$ was assigned to $p_i$, we can find the nodes $v$ by
finding the nodes $v$ such that $u \xrightarrow{p_i, p_{i+2}} v$ (or $u \xleftarrow{p_i, p_{i+2}} v$). Notice that, if $p_{i+1}$ is a backward edge, solving $u \xleftarrow{p_i, p_{i+2}} v$ by taking the backward approach prunes the search space as there is only a single source, i.e., $u$. This is an advantage of using the query linearization approach instead of separately solving each reachability requirement and then computing their intersections. Furthermore, among the possible nodes $v$, we only consider the ones that are unassigned; this is to guarantee that $f$ is injective. In case that there exists no node $v$ that satisfies the requirements, the match cannot be extended. Otherwise, the process continues at $p_{i+2}$ and each $v$.

**Case 2:** Node $p_{i+2}$ is assigned to $v \in V$: There are two sub-cases. (a) The reachability requirement established by $p_{i+1}$ has already been evaluated: it is not necessary to evaluate reachability again; we continue by considering $p_{i+2}$ and $v$. (b) The reachability requirement established by $p_{i+1}$ has not been evaluated: we establish the node predicates $p_i, pred : [ID = u.id]$ and $p_{i+2}, pred : [ID = v.id]$ and evaluate whether $u \xrightarrow{p_i, p_{i+2}} v$ (or $u \xleftarrow{p_i, p_{i+2}} v$, if $p_{i+1}$ is a backward edge, which is evaluated using the backward approach). If the requirement is satisfied, the process continues at $p_{i+2}$ and $v$. Otherwise, the match cannot be extended.

The above procedures guarantee that: (i) all the reachability requirements established by the edges in $G_P$ are satisfied; (ii) all the possible mapping functions (or all the possible matches) are considered; and (iii) because of Case 2, the reachability requirement defined by each edge is evaluated only once. If the algorithm reaches a successful assignment for $p_\ell$, then the algorithm reports that the corresponding mapping function $f$ is associated to a match of $G_P$ in $G$.

## 7.2. Algorithm

Figure 7-1 lists the pseudocode of GPQM. Three arrays are used to store the information of a partial (or full) match. (i) The mapping function is represented as the array $f$ where each position is associated to a node $u \in V_P$, which will eventually contain the mapping of $u$. In the beginning, all the mappings are *undefined* (which we denote as *undef*). (ii) We have a boolean array $h$, where each position is associated to an edge $e \in E_P$ to establish whether the reachability requirement established by $e$ has already been checked. (iii) We have a boolean array $g$ where each node in $V$ is associated to a position in the array. Specifically, $g[v] = true$, for a node $v \in V$, if a mapping for $v$ has already been established (through $f$). Intuitively, all the positions of arrays $g$ and $h$ are initialized with *false*.

The algorithm starts by computing a query linearization $p = p_{1...\ell}$ of $G_P$ and initializing the set $R$ of matches as empty. Then, all the nodes $v$ in $V$ such that $v \sim p_1$ are considered as roots of the DFS trees. In particular, the DFS traversal is performed by the recursive procedure `PROCESSNODE()` (see Figure 7-2). Each execution instance of this procedure considers a node $p_i$, a node $u \in V$ and a copy of $f, g$ and $h$. It is assumed that $u$ was assigned to $f(p_i)$ before the execution instance
was called. Thus, there is a partial match of $p_{1...i}$ in $G$ under $f$. Then, what the procedure does is attempting to extend this partial match according to the cases presented in last section.

**Algorithm 9: GPQM Algorithm**

**Input:** $G_P = (V_P, E_P)$, $G = (V, E, f_V, f_E)$  
**Output:** $R$

1. $p = E - GLA(G_P, G)$, $R = \emptyset$
2. for every $v \in V_P$ do $f[v] \leftarrow undef$
3. for every $e \in E_P$ do $h[e] \leftarrow false$
4. for every $v \in V$ do $g[v] \leftarrow false$
5. for every $u \in V$ do
6. if $u \sim p_1$ then
7. $f' \leftarrow copy(f)$, $g' \leftarrow copy(g)$, $h' \leftarrow copy(h)$
8. $f'[p_i] = u$, $g'[u] = true$
9. $ProcessNode(u, p, 1, f', g', h', G, R)$
10. return $R$

**Figure 7-1.**: GPQM algorithm.

We efficiently evaluate the reachability requirements in the query linearization by means of the function $\text{FindReachableNodes}()$ (see Figure 7-3). Let us consider an edge $p_{i+1}$ in the query linearization. We first tackle the case where $p_{i+1}.direction = forward$. Then, in order to evaluate the corresponding $(p_i, p_{i+2}, p_{i+1})$–reachability requirement, this function constructs a deterministic finite automaton (DFA) that accepts all the paths $\langle u, \ldots, v \rangle$ such that $v \in V$ is $(p_i, p_{i+2}, p_{i+1})$–reachable from $u \in V$. These walks are explored in a DFS manner. The roots of the DFS search trees are the nodes that satisfy $p_{i+1}.pred$. Starting from these nodes, we continue the DFS traversal constrained by the DFA. The process terminates when all the paths that lead to the final state, associated to $p_{i+2}.pred$, are considered.

If, on the contrary $p_{i+1}.direction = backward$, the $(p_{i+2}, p_i, p_{i+1})$–reachability requirement is validated in a similar manner. However, we use the backward approach to enhance efficiency. Specifically, notice that if we have already processed a node $p_i$ in the query linearization, then a particular node $u \in V$ such that $u \sim p_i$ has already been found. Hence, following the backward approach makes the search start from a single node (i.e., $u$) rather than all the possible nodes $v$ such that $v \sim p_{i+2}$. Note that many paths starting from valid nodes $v$ may not lead to $u$, so they should not be considered at this point. Then, the search is represented by a single DFS tree rooted at $u$. The DFA must accept the paths in $G$ that satisfy the reversal of the corresponding reachability expression (i.e., $p_{i+1}.re^R$). Thus, such paths are traversed inversely: for each node on a path, the incoming edges are considered. For a given reachability requirement, disregarding the direction of its corresponding edge, the function $\text{FindReachableNodes}()$ returns the set of nodes $v \in V$
Algorithm 10: PROCESSNODE() Procedure

Input: $u, p = p_1...\ell, i, f, g, h, G = (V, E, f_V, f_E), \mathcal{R}$

1. if $i = \ell$ then $\mathcal{R}.Add(f)$
2. else
3. $p'_i.pred : [ID = u.id]$
4. if $f[p_{i+2}] = \text{undef}$
5. $Q \leftarrow \text{FindReachableNodes}(p'_i, p_{i+2}, p_{i+1})$
6. for every $v \in Q$ do
7. if $g[v] = \text{false}$ then
8. $f' \leftarrow \text{copy}(f)$, $g' \leftarrow \text{copy}(g)$, $h' \leftarrow \text{copy}(h)$
9. $f'[p_{i+2}] \leftarrow v$, $g'[v] \leftarrow \text{true}$, $h'[p_{i+1}] \leftarrow \text{true}$
10. PROCESSNODE($v, p, i + 2, f', g', h', G, \mathcal{R}$)
11. else
12. $v = f[p_{i+2}]$
13. if $h[p_{i+1}] = \text{true}$ then
14. PROCESSNODE($v, p, i + 2, f, g, h, G, \mathcal{R}$)
15. else
16. $p'_{i+2}.pred : [ID = v.id]$
17. $Q \leftarrow \text{FindReachableNodes}(p'_i, p'_{i+2}, p_{i+1})$
18. if $Q \neq \emptyset$
19. $h \leftarrow \text{copy}(h)$, $h'[p_{i+1}] \leftarrow \text{true}$
20. PROCESSNODE($v, p, i + 2, f, g, h', G, \mathcal{R}$)

Figure 7-2.: PROCESSNODE() procedure.

Algorithm 11: FINDREACHABLENODES() Function

Input: $u, v, e, G = (V, E, f_V, f_E)$  
Output: $Q$

1. if $e.direction = \text{forward}$ then $\text{reachReq} = (u, v, e)$
2. else then $\text{reachReq} = (v, u, e)$
3. $Q \leftarrow \text{QueryConstrainedDFA}(\text{reachReq}, G, e.direction)$
4. return $Q$

Figure 7-3.: FINDREACHABLENODES() function.

that satisfy the corresponding reachability requirements with respect to a given $u \in V$.

Example. Figure 7-4 shows the DFS search tree traversed by GPQM to find the matches of the
7.2 Algorithm

query of Figure 5-2(a) on the attributed multigraph of Figure 5-1 by using the linearization of Figure 6-1(a). Notice that we associate each node in the linearization to nodes in the attributed multigraph that satisfy the reachability requirements. For example, the people associated to $p_3$ ($[\text{type} = \text{person}]$) are the ones that can be reached from Mary ($[\text{id} = \text{Mary}]$) through a path satisfying the expression $([(\text{type} = \text{friend})[\text{type} = \text{person}])^*[\text{type} = \text{friend}]$. Such paths are explored using the DFS constrained by the automaton presented in Figure 7-5.

Moreover, the example demonstrates that both reachability requirement evaluation and injective assignments prune the search space. For instance, there is just one possible mapping for $p_{11} = u_4$ in each branch of the search; because $p_7 = u_4$, then the only possible mapping for $p_{11}$ is the same mapping established for $p_7$, i.e., Photo 1 in the left branch and Photo 2 in the right branch. The highlighted walk (notes with double lines in Figure 7-4), which has mappings for all the elements in the query linearization, corresponds to the undirected walk presented in Figure 6-2. The mapping function associated to this walk constitutes a match of $p$ (and hence of $G_P$) in $G$ (the one of Figure 5-2(b)).

![Figure 7-4: DFS search tree traversed by GPQM to find the matches of the query of Figure 5-2(a) on the attributed multigraph of Figure 5-1 by using the linearization of Figure 6-1(a).](image_url)
Figure 7-5.: Deterministic Finite Automaton (DFA) corresponding to the reachability expression 
[[id = Mary](type = friend)[type = person]*)[type = friend][type = person] in Figure 5-2(a).

7.3. Correctness Proof

We formally prove the correctness of the GPQM algorithm by means of the following theorem:

**Theorem 9.** Given the attributed multigraph \( G = (V, E, f_V, f_E) \) and the generalized pattern query \( G_P = (V_P, E_P) \), the GPQM algorithm reports all the matches of \( G_P \) in \( G \).

**Proof.** We prove this theorem by showing the following invariant: when \( u \in V \) and \( p_i \) are processed in the recursive procedure \( \text{PROCESS NODE}(u) \), there exists a match of \( p_1...i \) in \( G \) under some bijective function \( f \). Let us denote such match as \( (v_1, v_3, ..., v_{i-2}, u) \), where \( v_j \in V \). Then, \( f(p_j) = v_j \), for odd values of \( j \) in \( 1 \leq j < i \), and \( f(p_i) = u \). Next, we show that this condition holds throughout the execution of the algorithm.

- **Initialization:** Let us consider the pseudocode presented in Figure 7-1. Because of the execution of lines 6 and 8 before calling \( \text{PROCESS NODE}(u, p_1) \) (in line 9), there exists a match \( (u) \) of \( (p_1) \) under \( f \) (i.e., \( f(p_1) = u \)).

- **Maintenance:** Let us assume that the invariant holds when \( \text{PROCESS NODE}(u, p_i) \) is executed (see Figure 7-2). Thus, there exists a match \( (v_1, v_3, ..., v_{i-2}, u) \) of \( p = p_1...i \) under a given function \( f \). If \( i < \ell \), we have to consider two cases: (i) \( f(p_{i+2}) \) has not been defined; and (ii) \( f(p_{i+2}) = v \), where \( v \in V \).

Let us consider first case (i). Notice that, because of lines 5 – 7, \( \text{PROCESS NODE}(v', p_{i+2}) \) is only called for unassigned nodes \( v' \in V \) such that \( v' \) is \((p_i, p_{i+2}, p_{i+1})\)-reachable from \( u \) (or \( u \) is \((p_{i+2}, p_i, p_{i+1})\)-reachable from \( v' \), if \( p_{i+1} \) is a backward edge). Thus, adding \( f(p_{i+2}) = v' \) (line 9, Figure 7-2) maintains the injective property of \( f \) and satisfies the corresponding reachability requirements. Hence, when \( \text{PROCESS NODE}(v', p_{i+2}) \) is executed (line 10, Figure 7-2), the invariant is satisfied given that \((v_1, v_3, ..., v_{i-1}, u, v')\) is a match of \( p_1...i+2 \).

For the case (ii), we have two sub-cases: the case where the reachability requirement established by \( p_{i+1} \) has already been evaluated and the case where it has not. In the former, we need not change the function and the invariant will hold when \( \text{PROCESS NODE}(v, p_{i+2}) \) is called (line 14, Figure 7-2). In the latter, \( \text{PROCESS NODE}(v, p_{i+2}) \) is only called if the node \( v \) is \((p_i, p_{i+2}, p_{i+1})\)-reachable from \( u \) (or if \( u \) is \((p_{i+2}, p_i, p_{i+1})\)-reachable from \( v \), if \( p_{i+1} \) is
a backward edge). Furthermore, the mapping function $f$ is not changed. Therefore, when \textsc{ProcessNode}(\(v, p_{i+2}\)) is called (line 20, Figure 7-2), \((v_1, v_3, \ldots, v_{i-1}, u, v)\) is a match of \(p_{1\ldots i+2}\); thus, the invariant holds.

- **Termination:** When a partial match is extended in any branch of the search and reaches position $\ell$ (line 1, Figure 7-2), we know that the function $f$ contains a match of $p_{1\ldots \ell}$ because of the invariant. Furthermore, all the matches of $p$ in $G$ are inserted in $R$ as all the possible mapping functions are considered. Namely, (i) the search is started from all the valid nodes (lines 5 – 9, Figure 7-1); and (ii) in all the stages of the search, all the possible injective mappings that satisfy the reachability requirements are considered (lines 5 – 10, 13 – 14 and 17 – 20, Figure 7-2). This set of matches also corresponds to the set of matches of $G_P$ in $G$ (see Theorem 6).

\[\square\]

## 7.4. Complexity Analysis

This section establishes the worst-case time complexity of GPQM. As we know that subgraph isomorphism, a simpler problem than Problem 4, is NP-complete [70], matching generalized pattern queries is expensive in the worst case. However, we also show that GPQM often performs better in practice than its worst-case bound and elaborate the reason.

Let us consider the generalized pattern query $G_P = (V_P, E_P)$ and the attributed multigraph $G = (V, E, f_V, f_E)$, where $n = |V|$ and $m = |E|$. As it was shown in Section 6.2.6, finding a convenient linearization $p = p_{1\ldots \ell}$ of $G_P$ takes $O(d|E_P| \lg d)$ where $d$ is the maximum degree of the nodes in $G_P$ (line 1, Figure 7-1). The initialization of $f, g$ and $h$ takes $O(|V_P|)$, $O(|E_P|)$ and $O(n)$, respectively (lines 2 – 4, Figure 7-1). Notwithstanding, these costs are insignificant with respect to the cost incurred by repeatedly executing the procedure \textsc{ProcessNode}().

In order to calculate the time complexity of GPQM, we first find an upper bound for the number of executions of the recursive procedure \textsc{ProcessNode}(). This number is equal to the number of nodes that are associated to a given $p_i$ in the DFS search trees. The next theorem establishes an upper bound.

**Theorem 10.** Let $p_{1\ldots \ell}$ represent a linearization of a generalized pattern query, and let $G = (V, E, f_V, f_E)$ represent an attributed multigraph, where $n = |V|$. A DFS search tree that represents the traversal of $G$ done by GPQM has $O(n^{\lfloor \ell/2 \rfloor})$ nodes associated to the different nodes $p_i$ in the linearization, for odd values of $i$ in $1 \leq i < \ell$.

**Proof.** Let us consider the undirected walks from the root of a DFS search tree to the lowest leaves, \textit{i.e.} the undirected walks that have $\lceil \ell/2 \rceil$ nodes associated to a $p_i$. These walks are the
ones that determine the height of the tree with the greatest number of executions of the procedure \textsc{ProcessNode}().

Next we calculate the number of possible assignments for each node \(p_i\) that requires the execution of the procedure \textsc{ProcessNode}(). There is one possible assignment for \(p_1\) in a given DFS search tree. According to the reachability requirement established by \(p_2\), there can be up to \(n - 1\) nodes that could be assigned to \(p_3\). This is because the node assigned to \(p_1\) cannot be assigned to \(p_3\) because of the injective requirement. Each of these \(n - 1\) nodes that can be associated to \(p_3\) may yield to \(n - 2\) possible assignments for \(p_5\) (again, the paths that lead to the nodes assigned to \(p_1\) and \(p_3\) are not considered); thus, the total number of possible assignments for \(p_5\) in the DFS search tree is \(O((n - 1)(n - 2))\) nodes. Similarly, the total number of assignments in the tree for \(p_7\) is \(O((n - 1)(n - 2)(n - 3))\) nodes. In general, \(p_i\) has at most \(\prod_{j=1}^{\lfloor i/2 \rfloor} (n - j)\) possible assignments.

Therefore, the total number of assignments in the tree for odd values of \(i\) in \(1 \leq i < \ell\) is:

\[
1 + \sum_{i=3}^{\ell-2} \prod_{j=1}^{\lfloor i/2 \rfloor} (n - j) = O(n^{\lfloor (\ell-2)/2 \rfloor}) = O(n^{\lfloor \ell/2 \rfloor - 1})
\]

In the worst case, we can have at most \(n\) DFS search trees with these characteristics, by assigning each distinct node in \(G\) to \(p_1\). In other words, we obtain a different DFS search tree by starting the search from a different node of \(G\). Thus, the total number of times that the procedure \textsc{ProcessNode}() can be executed is: \(\tilde{O}(n \times n^{\lfloor \ell/2 \rfloor - 1}) = O(n^{\lfloor \ell/2 \rfloor}).\)

The complexity of each execution of the procedure \textsc{ProcessNode}() is dominated by the execution of the function \textsc{FindReachableNodes}() (lines 5 and 17, Figure 7-2). The complexity of this function is determined by the construction of the DFA and the DFS search constrained by it (line 3, Figure 7-3). This can be done, using the traditional technique [142], as follows. We convert the regular expression, whose length we denote as \(r\), into an NFA with \(O(r)\) nodes. Then, this NFA is converted into a DFA in \(O(2^r)\) time. Let \(O(s)\) be a tight upper bound on the length of the paths read by this DFA. The DFS search constrained by such DFA takes \(O(n^s)\); this complexity is calculated using a similar analysis as the one of Theorem 5. Thus, the total complexity of an execution of \textsc{FindReachableNodes} is \(O(2^r + n^s)\). Given that this function is executed at most once in the procedure \textsc{ProcessNode}(), the complexity of this procedure is also \(O(2^r + n^s)\).

Hence, the total complexity of \textsc{GPQM} is the number of times that the procedure \textsc{ProcessNode}() is executed multiplied by the cost of each single execution. Specifically, the total complexity is \(O(n^{\lfloor \ell/2 \rfloor} \times \max\{2^r, n^s\})\) where \(r\) is the length of the longest regular expression in the query and \(s\) is the length of the longest matching path of any regular expression in such query.
It is, however, important to remark that this analysis gives an upper bound for the worst-case complexity. It assumes that, for every node $p_i$ in the linearization, all the possible assignments will be evaluated. The average-case situations are often not that “bad” because (i) when a node $p_i$ has already been assigned, only the assigned node is considered; and (ii) the node and edge predicates as well as reachability requirements often effectively prune the search space: for each possible assignment for a given $p_i$ (where $i$ is odd) only few assignments for $p_{i+2}$ are possible. Furthermore, this analysis was made assuming that the attributed graph is complete, while many graphs in the real world are sparse. Moreover, the values of $\ell$, $r$ and $s$ are often constant which makes our algorithm polynomial for many cases. Therefore, in practice, our algorithm shows much better performance than the given worst-case bound, as demonstrated in the experiments.

7.5. Experimental Evaluation

This section presents the implementation and experimental results of the proposed techniques. The main objective is to show the feasibility and efficiency of executing generalized pattern queries. Specifically, this section consists of four parts. We first describe the experimental setup that includes the information of the data set, the test set and the environment used (Section 7.5.1). Second, we introduce some examples of queries and the results of their evaluation on the complete DBLP graph by means of GPQM (Section 7.5.2). Third, we evaluate the performance of GPQM when the sizes of the query and the graph are varied (Section 7.5.3). Lastly, we illustrate the efficiency of using graph statistics (Section 7.5.4).

7.5.1. Experimental Setup

Datasets. We use the DBLP graph, which is a well-known computer science bibliography that captures information on authors, papers and where they are published (e.g., journals and conferences), as well as academic citations. We obtained the data from the DBLP website [52] on December 20, 2013 to produce a graph with more than 10 million elements (1,684,750 nodes and 9,955,181 edges) containing three node types and three edge types.

To study how the query response time varies with the graph size, we also generate synthetic graphs of varying sizes as subsets of the DBLP graph. To preserve the network structure among graph nodes, we perform the following steps to generate a graph with a specific number of nodes. We randomly select an arbitrary article, that has not been selected yet, and add the article node into the synthetic graph. Furthermore, we add all its edges and immediate neighbours (the 1-hop neighbourhood) if they do not belong to the generated graph. We repeat this step until the number of nodes reaches the target. We produce three synthetic DBLP graphs with $\{250K, 500K, 1M\}$ nodes and $\{923,762,\ 2,049,242,\ 4,510,296\}$ edges, respectively.
Query workload. There is no standard benchmark for generalized pattern queries. Therefore, we form queries that possess different features (such as reachability, pattern match queries and their combinations). As performance measures, we report the query execution time and the number of graph elements accessed during the processing.

Implementation. We implement a query processing system in C#. This system includes the query linearization and matching algorithms, as well as the query language parser and compiler. We use a server with 2.79GHz Xeon CPU and 24GB main memory (RAM) running the Windows Server operating system.

7.5.2. Queries on the Complete DBLP Graph

We introduce examples of generalized pattern queries with different characteristics. Then, we discuss experimental results of performing such queries on the DBLP graph.

Examples of Generalized Pattern Queries

We present some examples of generalized pattern queries in Figure 7-1. For instance, the generalized pattern query Q1 evaluates reachability. This query makes use of the Kleene star to find paths of arbitrary length that represent the relationship between a specific author and his/her academic descendants. In particular, Q1 allows to find all the academic descendants of Jiawei Han. On the other hand, the generalized pattern query Q2 is a combination of reachability and pattern match queries. Specifically, the query allows to find the articles co-authored by two academic descendants of Jiawei Han.

The generalized pattern queries Q3 and Q4 are examples of queries with the union operator. The former searches for journals that published papers written by M.Tamer Özsu or any of his co-authors; the latter looks for articles written by H. Vincent Poor and one of his colleagues, such that the colleague is either a prolific author or has worked with the him for at least 10 times.

Notice that queries Q5 and Q6 look similar. However, they have different meaning and execution costs. In particular, Q5 searches for prolific authors who have published articles at four conferences (i.e., VLDB, PVLDB, SIGMOD and SIGKDD). In contrast, Q6 is a pattern match query where the articles at each conference also constitute nodes of the query. This makes a big difference since the output of a generalized pattern query is established by its nodes. Specifically, Q5 reports only the authors that satisfy the condition, while Q6 looks for all the articles published by the author in each conference and reports all the possible combinations. As a result, Q6 can potentially have many

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1Academic descendant: An author A is an academic descendant of another author B, if A publishes at least 10 articles with B or A is an academic descendant of an academic descendant of B.

2Prolific author: An author is a prolific author when he/she has written more than 200 articles.
### Table 7-1: Examples of generalized pattern queries on the DBLP graph.

<table>
<thead>
<tr>
<th>Query</th>
<th>Query Representation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Q1</td>
<td>(cowork[count≥10] - author -)* cowork[count≥10]</td>
</tr>
<tr>
<td>Q2</td>
<td>[exp1]: (cowork[count≥10] - author -)* cowork[count≥10]</td>
</tr>
<tr>
<td>Q3</td>
<td>(write) or (cowork - author - write)) - article - publish</td>
</tr>
<tr>
<td>Q4</td>
<td>(cowork[count≥10] - author - write) or (cowork - author[publication&gt;200] - write)</td>
</tr>
<tr>
<td>Q5</td>
<td>[exp1]: write - article - published by</td>
</tr>
<tr>
<td>Q6</td>
<td>(write - article@VLDB - write - author)≤k</td>
</tr>
<tr>
<td>Q7</td>
<td>(write - article@VLDB - write - author)≤k</td>
</tr>
<tr>
<td>Q8</td>
<td></td>
</tr>
<tr>
<td>Q9</td>
<td></td>
</tr>
</tbody>
</table>
more matches than Q5, if the authors who satisfy the condition also have many combinations of papers from the four conferences. We discuss Q7 and Q8 in Section 7.5.3, and Q9 in Section 7.5.4.

Performance of the Query Evaluation

Figure 7-6 contains three subfigures that show the experimental results of evaluating the generalized pattern queries of Table 7-1 on the complete DBLP graph. In particular, Figure 7-6a presents the execution time and Figure 7-6b shows the number of visited graph elements of each query. Similarly, Figure 7-6c shows the number of matches for each query. For example, we found 18 matches of Q1 in 0.1 second visiting 6,575 graph elements.

These results show that the number of accessed graph elements is strongly correlated to the execution time, which validates the assumption of using the number of visited elements as our complexity metric. In contrast, the number of matches does not necessarily reflect the processing time.

We show the difference on the performance between Q5 and Q6. The execution time and the number of visited graph elements for Q6 are almost 100 times greater than the ones for Q5. Also, Q6 has 980 matches, while Q5 has only one match. The reason is that, in the case of Q5, once we establish the reachability from an author to one of the conferences, we can stop exploring other paths that lead to the same conference via a different publication. In other words, we only need to find if the author has published in the conference. In contrast, Q6 needs to enumerate all the publications from the author in the conference. That is why Q6 is more expensive and returns more results. This verifies the hypothesis posed at the end of Section 7.5.2. Also, these results demonstrate that the expressive power of generalized pattern queries allows us to write the queries in a flexible way, where the trade-off between the execution time and the amount of information returned is established according to our needs.

In summary, we show that our query linearization and matching algorithms can evaluate generalized pattern queries.

7.5.3. Varying Graph and Query Sizes

We show that the processing time depends on the size of both the attributed multigraph and the generalized pattern query. We use two generalized pattern queries: Q7 and Q8 of Table 7-1.

Let us consider first query Q7. The expression $\rho^{\leq k}$ indicates that the expression $\rho$ can be concatenated with itself from 0 to $k - 1$ times, i.e., $\rho^{\leq k} = \rho \cup \rho^2 \cup \cdots \cup \rho^k$. Thus, Q7 searches for the VLDB co-authors of Madden within a ($k + 1$)-hop. Note that $k$ sets an upper bound on the length of the paths that match the corresponding reachability requirement. We use this query on the experiments with varying $k$. Figure 7-7a shows the performance of our algorithm for this query Q7 on various sizes of the DBLP graph. When the graph size is smaller than 1M, or $k$ is equal or smaller than
7.5 Experimental Evaluation

Figure 7-6.: Experimental results for queries Q1–Q6 (see Table 7-1) on the complete DBLP graph.

3, the execution time of the query is less than 2 seconds. When we use the complete DBLP graph, however, the execution time grows with respect to the query length. As a result, while we find the 6-hop co-author neighbourhood for a specific author, via VLDB paper authorship, the query requires 50 seconds for the full DBLP graph. This shows that the execution time depends on the size of graph and the query complexity.

Next, we use a star pattern match query and change the number of nodes. Query Q8 is a template for such type of query. This is a complex query with few predicates that prune the search space. It does not have various candidates of query linearization orders. We vary the number of nodes in the query by changing the number of author nodes. Figure 7-7b shows that the execution time is
strongly related to the graph size and the query size.

In conclusion, the time required for processing a generalized pattern query depends on both the size of the attributed multigraph as well as the query.

![Graphs showing response time of GPQM for queries Q7–Q8](image)

**Figure 7-7.** Response time of GPQM for queries Q7–Q8 (see Table 7-1) on subgraphs of the DBLP graph of varying sizes.

### 7.5.4. Efficiency of E-GLA

The query linearization algorithm E-GLA uses graph statistics and selectivity estimation to generate a better linearization order. We compare E-GLA with GLA, an algorithm that optimizes the length of the linearizations but does not use the attributed multigraph statistics (see Section 3.3).

Notice that both GLA and E-GLA produce the same linearization for Q8. Thus, we use the query Q9 of Table 7-1 to evaluate the difference. This query searches for articles written by three authors and published at a journal with more than 500 publications. Furthermore, the first author must have at most 4 publications.

GLA starts from either an author or a journal node and then goes to the article node. Next, it selects an arbitrary non-visited node and repeatedly goes back to the center (i.e., the article node) until all the nodes are visited. In contrast, E-GLA starts from a journal node because it has the most restrictive predicate (journal with 500+ publications). Next, it visits the article node. Then, it selects the node with the lowest selectivity, i.e., the node that establishes that the author must have at most 4
publications. Finally, it goes back to the center and visits the non-visited nodes in an arbitrary order.

The performance of the matching algorithm GPQM is different depending on the used linearization algorithm. Figures 7-8a and 7-8b respectively show the execution time and the number of visited nodes, when each linearization algorithm is used, for different graph sizes. The E-GLA algorithm results in lower execution time and the difference between E-GLA and GLA increase with the the graph size.

![Figure 7-8](image.png)

**Figure 7-8.** Experimental results for query Q9 (see Table 7-1), using GLA and E-GLA linearizations, on subgraphs of the DBLP graph of varying sizes.
8. Conclusions

This thesis presents a novel approach to determine whether multigraphs $G_1 = (V_1, E_1)$ and $G_2 = (V_2, E_2)$, where $n = |V_1| = |V_2|$ and $m = |E_1| = |E_2|$, are isomorphic. In particular, this approach is based on a string matching technique called parameterized matching. Parameterized matching is used to find strings that have the same structure, i.e., the relative distances among the occurrences of each symbol is preserved. Our solution starts by representing $G_1$ in a linear manner, which we call graph linearization $p = p_1...p_\ell$. Then, we search for the walks in $G_2$ that parameterized-match the linearization. If there exists at least one of such walks, we conclude that the graphs are isomorphic. The correctness of our approach is formally proven.

We develop a Graph Linearization Algorithm called GLA. This algorithm does a DFS–like traversal on $G_1$ guided by heuristics that consider the number of unexplored adjacent edges that nodes have. The GLA algorithm produces short linearizations as illustrated through empirical examples. In fact, it is proven that the produced linearizations are $2$–approximate length-optimal. We show that the time complexity of GLA is $O(m + nd \lg d)$ where $d$ is the maximum degree of the nodes in the graph.

New optimizations can be included in the GLA algorithm to incur in lower time during the matching phase. In particular, the matching time does not only depend on the length of the linearization, but also on the order of comparisons. For instance, the topological graph statistics of the multigraphs can be used to produce a linearization that prunes the search space during the matching phase. For example, if the frequency of some nodes of a certain degree is low, it would be appropriate to start the linearization from such nodes. However, for clarity, in this thesis, we focus on the fundamental approach only.

Furthermore, we devise a matching algorithm called PMG that searches for walks in $G_2$ that parameterized-matches the linearization $p = p_1...p_\ell$. Specifically, this algorithm does a DFS traversal on $G_2$ where all the feasible mappings from the graph elements in the linearization to the graph elements in $G_2$ are explored. One of the key ideas of the algorithm is to prune the search space by considering node degrees and previous assignments. The time complexity of PMG is $O(nd^{\lfloor \ell/2 \rfloor})$.

We experimentally evaluate the efficiency of our solution by comparing with a prominent graph isomorphism algorithm called VF2. Experiments on synthetic graphs show that our algorithm performs better for both sparse graphs and complete graphs, but the difference is more significant for
complete graphs. We also perform experiments on benchmark graphs. For those, our algorithm reported better time results than VF2 in about half of the datasets. More precisely, VF2 excels in regular graphs, while our algorithm is significantly faster in 65% of Miyazaki-based constructed graphs. This is an interesting result since Miyazaki-constructed graphs constitute one of the hardest cases for graph isomorphism algorithms [141]. It is important to remark that in some cases where our solution is not short running, VF2 is fast. However, in the majority of the short-running cases, namely 66% of such cases, our algorithm runs faster. This opens up a possibility of a hybrid algorithm that selects between these two algorithms, either statistically-based on the graph topology or dynamically after running for some time, which we leave as future work.

We present a straightforward adaptation of our approach to determine whether $G_1 = (V_1, E_1)$ is isomorphic to a subgraph in $G_2 = (V_2, E_2)$. The resulting algorithm, called PMG-SI, preserves the time complexity of PMG, i.e., $O(nd^{\ell/2})$ where $n = |V_2|$ and $d$ is the maximum node degree in $G_2$. We experimentally evaluate the algorithm on synthetic graphs $G_2$ of varying sizes. For the graphs $G_1$ we used path, star, cyclic and complete graphs. Experimental results verify that the matching time depends on the linearization length and the promptness in which mismatches are detected.

Moreover, we extend our approach to query attributed multigraphs. In fact, we define a new type of queries called generalized pattern queries that establish predicates, reachability and topological requirements. These queries are multigraphs that establish predicates on a set of nodes of interest (through node predicates) and the reachability requirements among them through highly-expressive regular expressions associated to the edges. Such expressions are composed by nodes and edges (associated to predicates) and the regular expression operators (i.e., concatenation, union and Kleene star). Thus, each edge $e = (u, v)$ in the query defines a complex reachability requirement from node $u$ to node $v$. It is important to remark that previous reachability query models do not support evaluation of predicates on intermediate nodes and edges nor the Kleene star operator. This new type of queries can represent pattern match queries, reachability queries and beyond.

Then, we use the linearization approach to solve the problem of finding the matches of a generalized pattern query $G_P = (V_P, E_P)$ in an attributed multigraph $G = (V, E, f_V, f_E)$. Each vertex and each edge in $G$ is associated to a set of attributes, which are defined by the functions $f_V$ and $f_E$, respectively. In order to produce a convenient linearization, we propose an algorithm, called E-GLA, that takes into account the statistics of the attributed multigraph to linearize the generalized pattern query. In particular, for each node $u \in V_P$, the selectivity of $u$ is calculated as the probability of selecting a node from $G$ that satisfies the predicates of $u$. Then, the main heuristic of E-GLA is starting (or continuing) the query linearization from the node with lowest selectivity. In this sense, the mismatches are early detected during the matching phase and, hence, the search space is pruned. Furthermore, E-GLA query linearizations are also 2–approximate length–optimal. The time complexity of the algorithm is the same as the one of GLA.
The matching algorithm, called GPQM, does a DFS traversal on the attributed multigraph. The different feasible mappings between the nodes in the query linearization and the nodes in the DFS search tree are considered. In order to evaluate reachability, we construct a deterministic finite automaton (DFA) that accepts all the paths that satisfy the corresponding reachability expression. Then, we perform another DFS constrained by such automaton. The time complexity of GPQM is $O(n d^{\lfloor \ell/2 \rfloor})$, where $n = |V|$, $d$ is the maximum node degree in $G$ and $\ell$ is the length of the query linearization.

We experimentally validate the algorithm on the DBLP graph. We formed generalized pattern queries that possess different features, including reachability, pattern match queries and their combinations. Our experiments verify that the number of visited nodes is strongly correlated with the execution time. Furthermore, the results illustrate that the expressive power of generalized pattern queries allows us to write the queries in a flexible way, where the trade-off between the execution time and the amount of information returned is established according to our needs. Other set of experiments demonstrate that the processing time of a generalized pattern query depends on both the size of the query and the attributed multigraph. In order to evaluate the effectiveness of E-GLA heuristics, we compare the performance of GPQM when either E-GLA or GLA linearization algorithm is used. The results indicate that, when E-GLA is used, the execution time is lower and the difference increase with the graph size. As a conclusion, generalized graph queries are processed efficiently by considering graph statistics and selectivity estimation for query linearization.
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