

Master Thesis

Integrating SQL/PGQ in DuckDB

by

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ABSTRACT

This thesis will focus on implementing path finding algorithms in the open-source in-process SQL OLAP database management system DuckDB. This is done to further complete the integration of SQL/PGQ, an extension of SQL, in DuckDB. The algorithms, one of which is a batched version of Bellman-Ford, will make it possible to retrieve the path length between any two nodes in a (weighted) graph. Batched Bellman-Ford and other algorithms, will be implemented as user-defined functions in DuckDB, which make for a lightweight approach that does not alter the internal workings of DuckDB. In addition, possible optimizations in DuckDB for graph-like queries will be identified and implemented.

1 INTRODUCTION

There is a growing desire to perform more complex analyses on the increasing amounts of data being gathered. This data is connected, which will often be represented and thought of as a graph [44]. An extended survey of Sahu et al. [42] shows that graphs are used across a very diverse range of domains. Graphs often provide a natural way to structure data involving entities, represented as vertices, and the relationships between them, represented as edges. In turn, this increased desire has caused an increased rise in the attention given to graph database management systems (GDBMSs) [17]. These systems provide a new way of visualising and analysing graph data. However, the performance of these systems often leaves to be desired [17, 37]. The question is whether there is a need for this new method of looking at the data. Traditional relational database management systems (RDBMSs) are perfectly capable of storing graph data by using a vertex table and an edge table. Still, providing a way to perform these more complex analyses has been difficult up to now due to the limitations of the standard query language SQL [34].

In response to the limited capabilities, a plethora of GDBMSs arose, each having its own graph query language [9]. Examples of systems and their query languages are TigerGraph with GSQL [53], Neo4j with Cypher [18], Oracle Labs PGX with Property Graph Query Language (PGQL) [54]. Amazon Neptune even supports three distinct languages, namely OpenCypher, Gremlin, and SPARQL. Within these graph query languages, it is often easier to write queries containing graph pattern matching and path finding. However, each query language has a different syntax, different semantics, and different capabilities. These problems make working with graph-based data cumbersome for users.

The limited capabilities of SQL will change, as an extension of SQL is scheduled to release in March of 2023 called SQL/Property Graph Query (SQL/PGQ), making it easier to support this new type of workload related to graph data in RDBMSs [31].

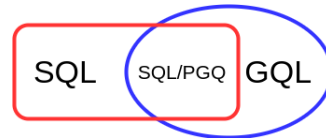


Figure 1: SQL/PGQ in relation to SQL and GQL.

For querying graph data, two functionalities are deemed most important: *graph pattern matching* and *path finding* [6]. In SQL it is possible to write queries containing graph pattern matching and path finding, but, these queries are often hard to write, understand, and inefficient to evaluate [25]. In particular, path finding requires using recursive queries. An example can be found in Listing 8, which shows a path finding query that returns all paths starting from *Person 1* [16]. Support for this was added in SQL:1999 [32], and is supported by popular RDBMSs such as PostgreSQL and SQLite. However, as shown by Michels and Witkowski [34], even relatively simple graph queries in plain SQL take up many lines and are more difficult to understand compared to the equivalent query in SQL/PGQ. The goal of SQL/PGQ is to provide a more compact syntax for graph-like data and make path finding queries, such as reachability or shortest path, more accessible.

Work on a standardization of a graph query language started in 2018 [5, 15, 21]. Currently the *ISO/IEC JTC1 SC32 WG3 Database Languages* working group is developing SQL/PGQ, which will become part of the upcoming SQL:2023 standard [31]. In SQL/PGQ a graph can be defined in terms of tables [33] and queries can contain special syntax for path finding and graph pattern matching. The standard is limited to read-only queries, as it will not be possible to modify the graph through the graph tables in SQL/PGQ. Therefore, the same workgroup is also working on Graph Query Language (GQL) [30], in which it will be possible to modify the data in addition to all the features contained in SQL/PGQ. GQL will be a superset in relation to SQL/PGQ, see Figure 1, which will have the ability to manipulate the data.

In this thesis we integrate the SQL/PGQ standard in DuckDB. DuckDB is an open-source in-process SQL OLAP database management system produced [39] by CWI [27]. With the new SQL/PGQ standard releasing soon, work on integrating it into DuckDB has already started by Singh et al. [45]. However, the standard has not been fully integrated as of now. In particular the shortest and cheapest path functions are not yet implemented. Additionally, the feature to return the nodes contained in these cheapest and shortest path functions will also be implemented. Therefore, the goal is to complete this integration by adding these functionalities. Additionally, we will identify and implement various optimizations especially related to typical graph queries in DuckDB.

The structure of this thesis proposal is as follows. Section 2 discusses relevant background information regarding graphs and SQL/PGQ. In Section 3 we formulate the research questions and the goals of this thesis. Section 4 describes how the research questions will be answered and evaluated and provides a research plan. We end with a conclusion in Section 6.

2 BACKGROUND

2.1 Property Graph Data Model

Graphs can be used to model complex, connected data through the use of vertices and edges. The vertices, sometimes referred to as nodes, represent objects. The edges, also referred to as relationships, represent the relations between objects. The simplest form of a graph is the *simple directed graph model* [9]. In this model, the graph consists of a set of vertices, and a set of edges. Each edge has a source vertex and a destination vertex. In the case of an undirected graph, both vertices within an edge act as the source and destination vertex. Graphs can either be weighted or unweighted. In weighted graphs, an edge between two vertices contains a weight. This weight can be different for every edge. In unweighted graphs, all edges are of equal weight. This is important when determining the shortest or cheapest path between two vertices in a graph. In the case of an unweighted graph, the shortest path will also be the cheapest path, since all weights are equal. For a weighted graph, this need not be the case.

The simple graph model suffices in certain cases, such as computing the reachability of a vertex from all other vertices. However, it is not possible to store any information in either the vertices or edges. For this we introduce the *Labelled Property Graph (LPG)* model, which is often used in graph databases [9]. The simple graph model is now enriched with the ability to assign labels to both vertices and edges. A label can for example be 'Person', or 'Knows'. Depending on the specific implementation of the graph database there can either be one or multiple labels assigned to a vertex or edge [5]. In addition to labels, there can also be properties assigned to vertices and edges. They can contain more specific information to the vertex or edge than the label. For example, a property could be (Age, 23) or (Name, Daniël). Examples of database systems that have based their data model on the LPG model are Neo4j [40], TigerGraph [53], and Oracle PGX [29].

2.2 SQL/PGQ

SQL/PGQ limits itself to read-only graph queries and how to define graph views over a tabular schema [15]. The two most important graph querying functionalities are graph pattern matching and path finding as described by Angles et al. [6]. These functionalities become more accessible with the addition of SQL/PGQ and queries involving these can be more easily expressed [25, 34].

With SQL/PGQ, graphs are stored as a set of vertex tables and edge tables, where each row in a vertex/edge table represents a vertex/edge in the graph [12]. A graph can be defined using the SQL statement [46] found in Listing 1.

```
1 CREATE PROPERTY GRAPH <name> [WITH SCHEMA <schema>] [FROM
  <subquery>]
```

Listing 1: Creating a graph in SQL/PGQ

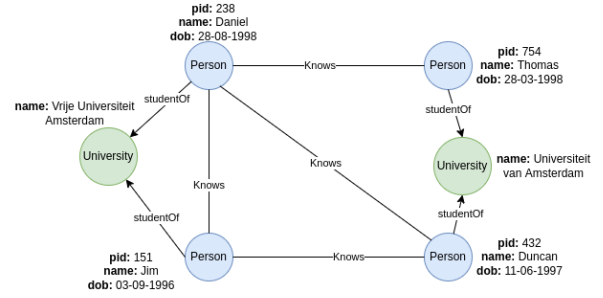


Figure 2: Graph of friend network and where they study.

For example, if we wish to create the graph of Figure 2, symbolizing a group of friends who all studied at some university, we would use the following query:

```
1 CREATE PROPERTY GRAPH friend_network
2 VERTEX TABLES ( Person PROPERTIES ( name, dob ),
3                 University PROPERTIES ( name ) )
4 EDGE TABLES ( knows SOURCE Person DESTINATION Person NO
5                 PROPERTIES,
6                 studentOf SOURCE Person DESTINATION
7                 University NO PROPERTIES )
```

Listing 2: Creating a friend network graph in SQL/PGQ

To match a pattern to this graph in SQL/PGQ, the **MATCH** syntax can be used [15], as can be seen in Listing 3. For example, the following selects the name and date of birth from persons from the graph in Figure 2 whose name is equal to 'Daniel':

```
1 SELECT p.name, p.dob
2 FROM Person GRAPH_TABLE (
3   MATCH ( a:Person WHERE a.name = 'Daniel' )
4   COLUMNS (
5     a.name ,
6     a.dob )
7 ) p
```

Listing 3: Pattern matching all nodes with the property name Daniel

Matching such a simple graph pattern is also relatively straightforward in plain SQL, however, it could be argued that the SQL/PGQ syntax feels more natural to write than the plain SQL one, since it is closer to how the human mind tends to interpret the world regarding objects and their connections [41]. As can be seen in Listing 3, we use the () notation to address a node. Addressing an edge can be done by using []. To indicate that an edge is pointing from source to destination we use (source)-[edge pattern]->(destination). This ASCII-art style is inspired by the syntax of Cypher, one of the most widely used graph query language [38].

A more complex graph pattern, involving both nodes and undirected edges could look like the following:

```
1 MATCH ( a:Person WHERE a.name = 'Daniel' )-[ e:knows ]-(
  b:Person )-[ f:studentOf ]-( c:University )
```

Listing 4: Pattern matching using nodes and edges

This statement would extract all patterns that match node *a* being a Person with the name Daniel, who knows a Person who is a studentOf a University. Within every node or edge, we can filter

the possibilities by adding a **WHERE** statement, as can be seen in Listing 4.

One of the features of SQL/PGQ will be the ability to match a single edge pattern or a parenthesized path pattern for an arbitrary length [15]. An example where we want to find paths of length 2 to 5 of *knows* edges:

```
1 MATCH ( a:Person )-[ e:knows ]->{2,5}( b:Person )
```

Listing 5: Path length of 2 to 5 knows edges

Finding such a path in plain SQL is significantly more difficult [34]. SQL/PGQ is not limited to quantifying the upper-bound of the path length in such a path finding query. Similar to regular expressions, it is possible to use the Kleene star (*) operator, to indicate that the pattern can occur 0 or more times. Additionally, matching the pattern 1 or more times is possible using the Kleene plus (+) symbol. The following is an example of a pattern using the Kleene star operator:

```
1 MATCH ( a:Person )-[ e:knows ]->*( b:Person )
```

Listing 6: Path length of arbitrarily many knows edges

Another addition in SQL/PGQ will be the ability to return the path corresponding to a query. Alongside returning every path that adheres to a given pattern, it will be possible to return the cheapest or shortest path. The cheapest path will return the path in which the total weight of the edges along the path are the lowest. The shortest path returns the path containing the least amount of hops from the starting node to the end node. In case the graph is unweighted, cheapest and shortest path will provide equivalent results, since the weight of any edge will be equal to 1.

2.3 Graph traversal algorithms

2.3.1 Multi-Source Breadth-First Search (MS-BFS). In order to compute the shortest path of unweighted graphs, the batched variant of the MS-BFS algorithm developed by Then et al. [51] can be used. The pseudo-code of the algorithm is provided in Algorithm 1. The algorithm is an example of a bulk algorithm that fits well in the vectorized execution engine of DuckDB [11]. It is able to run multiple BFSs concurrently on the same graph in a single CPU core. Furthermore, it can make use of Single Instruction Multiple Data (SIMD) instructions, such as AVX-512 [23], that are available in modern CPUs. This lets us handle 512 BFS steps in one CPU cycle, further increasing the efficiency in CPU usage. Additionally, it has the ability to scale up as the number of CPU cores increases. Since there are no dependencies between the various BFSs, they can be divided over multiple cores. It makes use of the small-world property [2] that occurs when the diameter of a graph is small in comparison to the total number of nodes. This means that each BFS discovers most vertices in a few iterations, and concurrent BFSs have a high chance of overlapping sets of discovered edges in the same iteration. This allows access to be shared among the multiple BFSs and reduce the chance of cache misses, reducing the overall computation time [51].

2.3.2 Cheapest path. To find a cheapest path in weighted graphs either the Dijkstra or Bellman-Ford algorithms can be used. The Dijkstra algorithm has a worst-case time complexity of $O(|E| +$

Algorithm 1 MS-BFS

```

Input:  $G, \mathbb{B}, s$ 
2:  $seen_{s_i} \leftarrow \{b_i\}$  for all  $b_i \in \mathbb{B}$ 
    $visit \leftarrow \bigcup_{b_i \in \mathbb{B}} \{(s_i, \{b_i\})\}$ 
4:  $visitNext \leftarrow \emptyset$ 
   while  $visit \neq \emptyset$  do
6:   for each  $v$  in  $visit$  do
        $\mathbb{B}'_v \leftarrow \emptyset$ 
8:     for each  $(v', \mathbb{B}') \in visit$  where  $v' = v$  do
            $\mathbb{B}'_v \leftarrow \mathbb{B}'_v \cup \mathbb{B}'$ 
10:    end for
       for each  $n \in neighbours_v$  do
12:          $\mathbb{D} \leftarrow \mathbb{B}'_v \setminus seen_n$ 
           if  $\mathbb{D} \neq \emptyset$  then
14:              $visitNext \leftarrow visitNext \cup \{(n, \mathbb{D})\}$ 
                $seen_n \leftarrow seen_n \cup \mathbb{D}$ 
16:             Do BFS computation on  $n$ 
           end if
       end for
18:   end for
20:    $visit \leftarrow visitNext$ 
    $visitNext \leftarrow \emptyset$ 
22: end while

```

$|V| \log |V|)$ [19], which is better than Bellman-Ford's worst-case time complexity of $O(|V| \cdot |E|)$ [7]. However, the expected runtime of Bellman-Ford is $O(|E|)$ in large dense graphs with low diameter [57]. Then et al. [50] propose a Batched Bellman-Ford-based algorithm, shown in Algorithm 3, to find the shortest distance between two nodes in a graph. This algorithm can make use of SIMD instructions to increase the CPU usage efficiency, which is not possible with the standard Bellman-Ford algorithm [8], which is shown in Algorithm 2.

Algorithm 2 Bellman-Ford

```

Initialize-Single-Source( $G, s$ )
2: for  $i \leftarrow 1$  to  $|V[G]| - 1$  do
       for each edge  $(u, v) \in E[G]$  do
4:         Relax( $u, v, w$ )
       end for
6:   end for
       for each edge  $(u, v) \in E[G]$  do
8:         if  $d[v] > d[u] + w(u, v)$  then
               Return False
           end if
10:   end for
       end for

```

2.4 DuckDB

DuckDB is a database management system specialized in OLAP workloads [39]. This means that the system is optimized more towards analytical queries, touching large data volumes using joins and aggregations. Just like SQLite, DuckDB is an in-process system, though SQLite is specialized in OLTP workloads.

Algorithm 3 Directed batched Bellman-Ford-based algorithm

```
Input: WeightedGraph G, Array<Vertex> sources
2: Output: VertexProperty<BatchVar<double>> dists
   VertexProperty<BatchVar<bool>> modified =
   falsedists = Infinite

4: for i=1..sources.length do
6:   Node v = sources[i]
   dists[v][i] = 0
8:   modified[v][i] = true
end for
10: bool changed = true
   while changed do
12:   changed = false
   for each v in G.vertices do
14:     if not modified[v].empty() then
       for each v in G.neighbours(v) do
16:         double weight = edgeWeight(v,n)
         for each i in modified[v] do
18:           double newDist = min(dists[n][i],
dists[v][i] + weight)
           if newDist != dists[n][i] then
20:             dists[n][i] = newDist
             modified[n][i] = true
22:             changed = true
           end if
       end for
     end for
   end while
```

DuckDB consists of a number of components: Parser, logical planner, optimizer, physical planner, and execution engine. The system can be accessed through a C/C++ API, as well as a SQLite compatibility layer. The SQL parser is based on the PostgreSQL SQL parser [39]. The logical planner consists of a binder and a plan generator. The binder is responsible for the expressions from the query related to anything containing a schema such as tables and views and retrieves the required columns and data types. The plan generator then creates a tree of basic logical query operators from the retrieved parse tree. Once the logical planner is done, the optimizer is used to optimize the logical plan. This will result in an optimized logical plan which is given to the physical planner where it is turned into a physical plan. The physical plan consists of operators, where each operator implements one step of the plan. An example of a unary operator is the *scan*, which scans a table and brings each tuple of a relation into main memory [20]. A join operator that makes use of two tables is an example of a binary operator.

These operators are split up into pipelines, which determines the order of operation execution. A query can consist of one or more pipelines, some of which contain a dependency on another pipeline. A pipeline with a dependency is for example, one containing a join operator. The start of a pipeline is referred to as a source and the end is referred to as the sink, which is where all the data is collected

(materialized). Only sink operators, such as sorting, building a hash table, and hash aggregation need to see all the data before they can proceed. All other operators do not need to materialize all data before proceeding. In the case of binary operators there are always two pipelines, one that builds the hash table, and one that probes this hash table. Both pipelines contain a source and a sink, and since the probing is a non-materializing operation it can be scheduled in the middle of a pipeline.

With every join, a hash table needs to be built on which the join can be performed and the operator needs to wait for the entire hash table to be built. Only then can the join be correctly executed. In the same vein, another pipeline might require the outcome of this join before it can be executed, creating a chain of dependencies.

DuckDB makes use of hash tables to perform join operations¹. Cardinality estimation is performed to assess which of the two tables is the smaller one. These estimations can be wrong, so it is not guaranteed that the smallest table is always used.

This smallest table is then used to build a hash table from, this is also referred to as the *sink* or the build side. The other, larger, table is then used to probe the hash table, looking for matching entries, this is referred to as the *source* or the probe side. Whenever the two tables are of equal size, a random one of the two is chosen to be the sink.

The execution engine of DuckDB is vectorized [39]. The use of vectors is more CPU efficient than the more common tuple-at-a-time execution found in other DBMSs [13]. ACID-compliance (Atomic, Consistent, Isolated, and Durable) in DuckDB is provided by Multi-Version Concurrency Control [39]. To allow for persistent storage, DuckDB uses the DataBlocks storage layout, which is read-optimized [39]. A useful feature of DuckDB is the allowance of extension modules, also referred to as Scalar user-defined functions (UDFs). These Scalar UDFs are as fast as the built-in functions of DuckDB due to the vectorized query processing, meaning that the parallelization of the UDF is handled by DuckDB.

2.5 Current state of SQL/PGQ in DuckDB

This thesis will continue upon the work done by Singh et al. [45]. They identified several challenges that needed to be addressed. DuckDB is primarily intended for tabular workloads and wants to limit its core features to those required for the tabular types of workloads. One of the first challenges to be tackled was successfully parsing SQL/PGQ queries. The SQL/PGQ queries are transformed into plain SQL queries, for example, see Listing 9 and Listing 10. It can be observed that Listing 10 contains a Kleene star, making it more difficult to transform into a plain SQL query. Since SQL/PGQ contains new statements (e.g. GRAPH, LABEL, PROPERTIES, see Listing 2), these had to be added to the parser of DuckDB. Therefore, minimal changes to the parser and transformer were made to allow correct parsing of SQL/PGQ queries. Another consequence of limiting the core features was the decision to implement the new operators, such as the batched MS-BFS algorithm [51], as UDFs. With MS-BFS it is possible to compute the reachability of any number of nodes given one or more source nodes. DuckDB handles the

¹ Whenever the ids of the smaller table are dense, meaning the maximum id is not much larger than the size of the table, an array is used instead, eliminating the need to build a hash table. This is referred to as a perfect join [1].

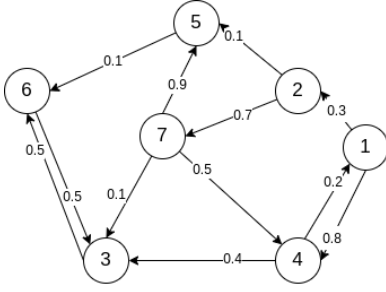


Figure 3: An example of a weighted graph

parallelization for Scalar UDFs using the morsel-driven method [26], which helps when scaling the batched MS-BFS algorithm. Another benefit of implementing these operators as Scalar UDFs instead of expressions is the fact that little to no changes have to be made to the internals of DuckDB. For example, to implement reachability, no further changes needed to be made to the parser and no new logical and physical operators had to be introduced.

Another challenge Singh et al. looked at was the access patterns. Typically, a pointer-based data structure can be used for $O(1)$ lookup. In this case, every node contains a mini-index to all nearby nodes [55]. However, this becomes inefficient in the case of multiple traversals due to poor memory locality. An alternative is to make use of a Compressed Sparse Row (CSR) data structure. With this data structure, the rowids of both the vertex and edge tables are condensed into integers in the range $[0, |V|)$ and $[0, |E|)$. Most importantly are the edge tables, which are usually much larger, leading to a higher chance of poor memory locality. Thus, Singh et al. created a UDF to create such CSR-like data structures, which are then used during the execution of the MS-BFS UDF. A weighted graph is represented in Figure 3. The rows in the table represent the source vertices, while the columns represent the destination vertices. Vertices that share an edge contain a weight, ranging between $[0, 1)$ in this example in their respective cell. For example, the edge from vertex 1 to vertex 2 has a weight of .3. Figure 4 shows the CSR representation for the graph in Figure 3. The row pointers point to the index of the first column for which there is an edge going from the row index to the column index. For example, in Figure 3 we observe the first edge from row 1 goes to column 2. For row 4, the first edge goes to column 1, therefore, the row pointer 4 goes to column index 1. The offset between two row pointers can be used to determine the number of edges for a row. The value corresponding to the edge can be found in a separate array using the same index as the column index. This condenses the matrix-like data structure into regular arrays, improving the memory locality.

3 GOALS

The goal of this thesis is to build on the work done by Singh et al. described in Section 2.5. To further complete the integration of SQL/PGQ into DuckDB several UDFs need to be implemented. The next step after being able to compute the reachability will be to compute the minimal length of a path between any two nodes. The paths can either have a weight equal to 1, or a custom weight which is a positive number. In either case, the minimal path length can be

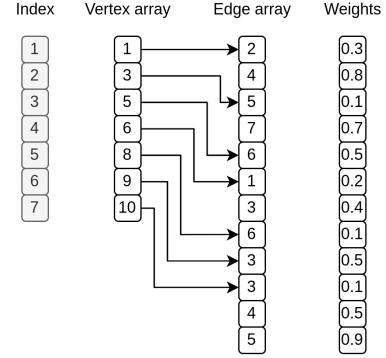


Figure 4: CSR representation of graph in Figure 3

computed using either Bellman-Ford or Dijkstra’s algorithm. Once computing this is possible, we will work on retrieving all the nodes and edges contained in the shortest / cheapest path and return this in a query.

Another goal of this thesis is to identify and implement various optimizations in DuckDB especially related to graph queries. One of these optimizations has already been identified and considers the case when multiple identical hash tables are built due to duplicate join statements. These duplicate joins can occur in plain SQL queries, though are more common in SQL/PGQ queries due to the common occurrence of many-to-many relationships in graphs. For example, first joining a person table with a knows table (for person1_id), after which the person table is again joined with the knows table (for person2_id). In both instances, the same hash table is built for an equivalent join, resulting in equivalent hash tables, which causes redundant work. This is wasted computation time and storage, given that one hash table would suffice for both joins. The optimization will be to identify whenever these identical joins are done and ensure that only the minimally required amount of hash tables are built.

The research questions of this thesis will be:

- (1) How can shortest and cheapest paths best be implemented using MS-BFS?
 - (a) What are the trade-offs between using Bellman-Ford and Dijkstra’s algorithms?
 - (b) How can the number of states during computation be minimized to allow for optimal execution?
- (2) What are the performance bottlenecks in SQL/PGQ?
 - (a) How can these performance bottlenecks be experimentally evaluated?

4 METHODOLOGY

4.1 Shortest path

Computing the shortest path between two vertices in an unweighted graph can be done using MS-BFS. In the current implementation by Singh et al., it is possible to determine the reachability of a node. However, to determine the shortest path, some form of state needs to be kept, such that it can be tracked how many hops have been

done before reaching the destination node. Often, graphs have the small-world property, the distance between any two nodes is very small compared to the size of the graph [51]. In addition, the shortest distance is always smaller than or equal to the diameter, the greatest distance between any two nodes, of the graph. Therefore, in unweighted graphs, the size of the values needed to be stored in memory can also be relatively small. For every source node for which the shortest distance is computed, it will likely be enough to store the distance within 8 bits. Since the distance can never be negative, an unsigned 8-bit integer would often suffice, in the case that the diameter of the graph is less than 255. However, we should care not to exceed this limit as the integer will overflow, causing it to wrap around to 0. Whenever that would happen we should copy the values into 16-bit or 32-bit unsigned integers to ensure no overflow errors occur. This does reduce the efficiency of the MS-BFS algorithm compared to the reachability implementation. Computing the reachability only requires a 1-bit bool value per BFS, which is set to 1 in case the destination node is reachable, 0 otherwise. Using the AVX-512 operation, it is possible to execute 512 reachability searches at once. However, since computing the shortest path requires at least 8 bits to keep track of the distance, it will only be possible to execute $512/8 = 64$ BFS searches at a time. If we were to use 32-bit integers, only 16 BFS searches could be run. Unsigned integers are preferred over signed, since the length of a path can never be negative, there is no need for negative integers. Denoting that a node is unreachable can be done by using `UINT_MAX` [35]. During execution, we keep track of the depth of the MS-BFS. Whenever this depth reaches `UINT_MAX`, we know there is a chance for an overflow and make sure this is handled correctly.

To compute the cheapest path in weighted graphs, there are typically two algorithms that can be used, Dijkstra’s algorithm using a Fibonacci heap and the Bellman-Ford algorithm. Then et al. [50] propose a batched Bellman-Ford algorithm to compute the geodesic distance in weighted graphs. They find that the performance of the Bellman-Ford algorithm is 3-10x higher compared to a batched variant of Dijkstra’s algorithm [50]. Therefore, we will first look to implement the batched Bellman-Ford algorithm proposed by Then et al. as shown in Algorithm 3. Implementing a batched version of Dijkstra’s algorithm will be more complex. During the batched version multiple instances of Dijkstra’s algorithm will run, each having its Fibonacci heap. Given that each heap will be of a different structure, the effectiveness of SIMD instructions will be reduced. Therefore, it is not likely that a batched version of Dijkstra’s algorithm will outperform the batched version of the Bellman-Ford algorithm. To test the performance of the implementations we will be using the LDBC Graphalytics benchmark [24], which is designed to test various algorithms such as single-source shortest path (SSSP) and BFS. Additionally, we will be using the LDBC Social Network Benchmark [4] to further test the performance of the implementations.

4.2 Shared hash join

Queries can contain multiple expensive joins which are in essence identical. This also happens in graph-like queries, where it is common to join the vertex table twice on the edge table. Once for the source of the edge, and once for the destination of the edge.

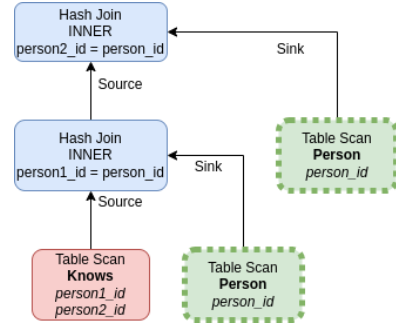


Figure 5: Example of a duplicate sink state in the physical plan

In the current state of DuckDB, for each join a hash table is built from the smaller table. However, this is wasteful in the case where queries containing multiple join operations have identical sinks. For example, there exists a vertex table *Person* containing the column *person_id*, and an edge table *Knows* containing the columns *person1_id* and *person2_id*. These tables are used to build a CSR representation for the edge table. Thus, it is necessary to perform two joins, namely $\text{person1_id} \bowtie \text{person_id}$ and $\text{person2_id} \bowtie \text{person_id}$. In this case, the right side is identical in both joins. See Figure 5 for the relevant part of the physical plan of this example query. An optimization is to build this hash table only once, and reuse it for any identical joins containing the same sink. This will eliminate the need to build the same hash table multiple times.

The plan for the shared join optimization is to make the sink shareable between multiple pipelines in the case where the joins are identical. This will eliminate the need of building multiple identical hash tables since they can be reused. During the building phase of the pipelines, we will save all join operations we come across in combination with its respective pipeline. In the case where a duplicate join is encountered will the sink state of that pipeline be shared with the sink state of the pipeline seen earlier. In addition, the dependency of the parent pipeline on the current pipeline needs to be changed. The optimization can not be performed on FULL or RIGHT joins. For these joins a state needs to be kept for the hash tables to see if a particular value has been accessed or not. This access may vary between joins, and thus will result in different states. Therefore, the hash table cannot be reused for these types of joins.

In DuckDB, unit tests can be written using the *SQLLogicTests framework* to test the correctness of the implementation [28]. Assuming that the implementation works correctly, we will be testing the performance of the optimization using the CSR creation queries used by Singh et al. [45]. These queries contain a duplicate join on the vertex table. By performing tests with and without the optimization, we should see that the queries using the optimization have a lower run-time.

4.3 Planning

See Table 1.

Month	Goals
February	Work on literature study Start on optimization shared hash tables
March	Work on optimization hash tables Start on shortest path using MS-BFS
April	Finish optimization hash tables Finish shortest path and experiment
May	Start on cheapest path using batched Bellman-Ford / Dijkstra
June	Finish cheapest path Start on experimenting
July	Finish up experimenting Finish up writing

Table 1: Initial planning

5 RELATED WORK

5.1 Resource Description Framework

Another form in which graph data can be modelled, aside from the property graph model, is *Resource Description Framework* (RDF). Data is modelled in the form of triples, which is composed of a *subject*, *predicate*, and an *object* [43]. The nodes within an RDF can be of three types: Internationalised Resource Identifiers (IRIs) used to globally identify entities and relations, literals, and blank nodes [22]. Node attributes are modelled as an extra outgoing edge (being the predicate) to another node which is the object stored as a literal. One way of modelling edge attributes is described by Sun et al. [48], where every edge attribute requires four new edges being added to the graph. This method of modelling edge attributes is generally applicable, though verbose and inefficient in terms of storage [48]. Database management systems such as BlazeGraph [10] and Amazon Neptune [3] have based their data model on RDF. Storing RDF data in relational databases has some challenges [14]. One way of storing data is by using a single table with three columns: subject, predicate, object. However, large-scale RDF data runs into performance and scalability issues [36]. In particular, queries containing joins prove to be difficult to handle [36, 37].

5.2 Self-joins

Self-joins occur whenever a table joins with itself. For example, a knows table containing two columns (person1_id, person2_id) is joined in the following query:

```

1 SELECT knows1.person1_id, knows2.person2_id
2 FROM knows k1
3 JOIN knows k2 ON k1.person1_id = k2.person2_id;
```

Listing 7: Self-join on the knows table

As described by Szarnyas [49], there are several challenges with self-joins. An important one is related to RDFs that are stored as one table, which requires a join for every attribute queried [37]. Another challenge is related to the hash-joins that are often used by database systems to perform join operations. Whenever a join between two tables is performed, the smaller of the two tables is used to build a hash table. The other table is then used to probe the hash table and look for matches. However, with self-joins, both

tables are equal in size, negating the advantage of building a hash table on a smaller table. In addition, the resulting table will probably be large in size when the table is of a many-to-many relation.

5.3 Mapping from Graph to Relational queries

Research has been conducted on the translation from a given query language to another one. Work that is particularly interesting to this thesis is translating graph query languages to SQL. An example of this is *Cytosm* (Cypher to SQL Mapping) by Steer et al. [47]. It acts as an application to execute graph queries on non-graph databases. In addition, they introduced *gTop*, a format that is able to capture the structure of property graphs and allow a mapping between graph query languages and a variety of database management systems. Steer et al. find that the translated SQL queries show comparable performance to GDBMSs [47].

Another example is GraphGen [56] that acts as an abstraction layer on top of an RDBMS. Underlying relational datasets are transformed and defined as graphs (*Graph-Views*). These graphs can then be queried using a graph API. The GraphGen framework has two main functionalities. First, users can define the structure of a graph using GraphGenDL, a Datalog-like domain-specific language (DSL). The other functionality is taking queries and executing them against the Graph-Views. The queries are specified by GraphGenQL, which is loosely based on SPARQL, Cypher, and PGQL [56].

SQLGraph allows Gremlin queries to be converted into SQL [48]. It uses a combination of relational and non-relational data storage. The *adjacency information* makes use of relational storage, while the attributes regarding all the nodes and edges are stored with JSON storage.

Lastly, there is IBM Db2 Graph by Tian et al. [52], which is an in-DBMS graph query approach that supports synergistic and retrofitable graph queries using the query language Gremlin inside the IBM Db2 relational database. Similar to GraphGen, it has a *graph overlay* method that exposes *graph views* of the relational data.

Zhao and Yu showed that it was possible to support a large class of graph algorithms, such as BFS, Bellman-Ford, PageRank, in SQL [58]. They introduce new operations, alongside the existing ones (selection, projection, union, set difference, cartesian product, and rename) to provide explicit support for graph algorithms.

6 CONCLUSION

In this thesis we will be working on implementing the minimal path length as a UDF in DuckDB. This will be done using the batched Bellman-Ford algorithm, and/or Batched Dijkstra’s algorithm developed by Then et al. The goal of this is to further complete the integrating of SQL/PGQ in DuckDB. Alongside, we will identify and implement optimizations that are specifically useful for graph-like queries. One of such optimizations is the shared hash join, which eliminates the need of building duplicate hash tables.

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A SQL VS SQL/PGQ QUERIES

```

1 WITH RECURSIVE paths(startPerson, endPerson, path) AS (
2   SELECT -- define the path as the first edge of the
      traversal
3     person1id AS startPerson,
4     person2id AS endPerson,
5     [person1id, person2id]::bigint[] AS path
6   FROM knows
7 UNION ALL
8   SELECT -- concatenate new edge to the path
9     paths.startPerson AS startPerson,
10    person2id AS endPerson,
11    array_append(path, person2id) AS path
12   FROM paths
13   JOIN knows ON paths.endPerson = knows.person1id
14   WHERE knows.person2id != ALL(paths.path) -- detect
      cycles
15 )
16 SELECT startPerson, endPerson, path
17 FROM paths
18 WHERE startPerson = 1;

```

Listing 8: SQL query WITH RECURSIVE

```

1 select c1id, c2id, c3id
2 from GRAPH_TABLE (aml,
3 MATCH (c1 IS customer)-[IS transfers]->(c2 IS customer)-[
  t2 IS transfers]->(c3 IS customer)-[t3 IS transfers
  ]->(c1)

```

```

COLUMNS (c1.cid AS c1id, c2.cid AS c2id, c3.cid AS c3id)
gt

/* The above SQL/PGQ query is transformed into the lower
SQL query */

select c1id, c2id, c3id
from (SELECT c1.cid as c1id, c2.cid as c2id, c3.cid as
      c3id
FROM customer c1, transfers t1, customer c2, transfers t2
, customer c3, transfers t3
WHERE c1.cid = t1.from_id
AND c2.cid = t1.to_id
AND c2.cid = t2.from_id
AND c3.cid = t2.to_id
AND t3.from_id = c3.cid
AND t3.to_id = c1.cid
)

```

Listing 9: SQL/PGQ query transformed into SQL query

```

1 select gt.c1id, gt.c2id
2 from GRAPH_TABLE (aml, MATCH (c1 IS CUSTOMER)-[t1 IS
  TRANSFERS*]->(c2 IS CUSTOMER) COLUMNS (c1.cid AS
  c1id, c2.cid AS c2id)) gt
3
4 /* The above SQL/PGQ query with a Kleene star is
   transformed into the lower SQL query */
5 select c1id, c2id
6 FROM
7 (
8   WITH cte1 AS (
9     SELECT min(CREATE_CSR_EDGE(0, (SELECT count(c.cid) as
      vcount FROM Customer c),
10    CAST ((SELECT sum(CREATE_CSR_VERTEX(0, (SELECT count(c.
      cid) as vcount FROM Customer c),
11    sub.dense_id , sub.cnt )) as numEdges
12   FROM (
13     SELECT c.rowid as dense_id, count(t.from_id) as cnt
14    FROM Customer c
15    LEFT JOIN Transfers t ON t.from_id = c.cid
16    GROUP BY c.rowid
17   ) sub) AS BIGINT),
18   src.rowid, dst.rowid ) ) as temp, (SELECT count(c.cid)
19   FROM Customer c) as vcount
20 FROM
21   Transfers t
22   JOIN Customer src ON t.from_id = src.cid
23   JOIN Customer dst ON t.to_id = dst.cid
24 )
25 SELECT src.cid AS c1id, dst.cid AS c2id
26 FROM cte1, Customer src, Customer dst
27 WHERE
28 ( reachability(0, true, cte1.vcount, src.rowid, dst.rowid
   ) = cte1.temp)
);

```

Listing 10: SQL/PGQ query transformed into SQL query with Kleene star