Fast, scalable WCOJ graph-pattern matching on in-memory graphs in Spark

Master thesis draft iteration 1.2

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Abstract

Graph pattern matching with their vast number of cyclic foreign-key joins is a new challenge for data processing systems, like Spark, because their intermediary results grow over linear with regards to the inputs and are materialized by the traditionally used binary joins. Worst-case optimal join algorithms, WCOJ's, are a natural match to tackle this challenge because they do not materialize the aforementioned large intermediary results. We investigate two major open questions regarding WCOJ's. First, we develop a WCOJ specialized to graph-pattern matching, namely self-joins on a relationship with two attributes, and compare its performance with a general WCOJ. Second, we propose a novel method to distribute WCOJ's. We show in our proposal that current methods to distribute WCOJ's, suitable for Spark, do not scale well to bigger graph pattern (five vertices and more). Based on this result we propose to keep the edge relationship cached on all workers but distribute the computation using a logical partitioning. Along the line of argumentation in COST [25], we aim to provide a fast WCOJ implementation in Spark which scales well in use of computational resources by trading off scalability in memory usage. This is a reasonable trade-off as most graphs today fit in main memory [23]. Our thesis provides the first distributed, open-source implementation of a WCOJ in a data processing system widely used in industry, namely Spark.

1 Introduction

Newly developed worst-case optimal join (WCOJ) algorithms, e.g. Leapfrog Triejoin, turned conventional thinking about join processing on its head because these multi-join algorithms have provably lower complexity than classical binary joins, i.e. join algorithms that join just two tables at-a-time. In the areas of data warehousing and OLAP, this finding does not have much impact, though, since the join patterns most commonly encountered are primary-foreign-key joins, which normally take the form of a tree or snowflake and contain no cycles. The computational complexity of FK-PK joins is by definition linear in size of the inputs. In these "conventional" cases, binary joins, e.g. hash joins, work fine. However, analytical graph queries often use foreign-foreign-key joins, which can grow over linearly in the size of their inputs, and often contain cycles. For these use-cases, worst-case optimal join algorithms excel because matching a pattern consisting of multiple joins causes binary joins to generate a rapidly increasing set of intermediate results, e.g. navigating a social graph with an out-degree in the hundreds, of which many matches are useless and get eliminated by later joins, e.g. the join closing the cycle. These kinds of join

patterns are frequently found during graph analysis, e.g. for graph clustering on social network graphs for customer relationship management or recommendation systems and fraud detection in the financial sector [4, 17]. Worst-case optimal join algorithms avoid large result materialization and hence promise to be orders of magnitude faster than binary joins. Therefore, we believe that worst-case optimal join algorithms could be a useful addition to (analytical) graph database systems.

We continue with a short example for a cyclic query and compare how this query is evaluated traditionally and with the new WCOJ's in place. The simplest example of a cyclical join query enumerates all triangles in a graph. This can be formulated as the following datalog query

$$Q(a, b, c) \leftarrow R(a, b), S(b, c), T(c, a)$$
 (1)

where R = S = T are aliases for the edge relationship. Traditionally, this would be processed by using multiple binary joins:

$$R \bowtie S \bowtie T \bowtie R \tag{2}$$

Independent of the chosen order, it can be proven that there exists cases where the intermediary result size is in $\mathcal{O}(n^3)$ with n=|R|=|S|=|T|. However, it is provable that maximal output of this query is in $\mathcal{O}(n^{3/2})$ [6, 28]. Hence, binary joins materialize huge intermediary results after processing parts of the query, which are much bigger than the final result after applying all joins. The described problem has been shown to be a fundamental issue with traditional join-at-a-time approaches [6, 28]. Fortunately, worst-case optimal join algorithms can materialize cyclic joins with memory usage linear to their output size by avoiding to produce large intermediary results [35, 29]. In practice, these algorithms have been shown to be highly beneficial for cyclic queries in analytical graph workloads in an optimized, single machine system [35, 30] and later in distributed shared-nothing settings [11, 3] - we describe these systems in more detail in section 3.

We identify two challenging, novel directions for our research. First, although, all of the systems cited above focus on queries widely used in graph pattern matching, e.g. clique finding or path queries, they use WCOJ's which are developed for general multi-way joins, however, graph pattern matching uses only self-joins on a relationship with two attributes, namely the edge relationship of the graph. This raises the question if WCOJ's can be optimized by specializing them for graph pattern matching - which is so far the only use-case that has been shown to benefit from WCOJ's in the literature. Second, while the communication costs for worst-case optimal joins in map-reduce like systems (an excellent definition of the term is given in [1]) is well-understood [1, 11], their scalability has not been studied in depth. Given the high complexity of worst-case optimal joins used and the fact that their only integration in a map-reduce like system [11] exhibits a speedup of 8 on 64 nodes (an efficiency of 0.125), leads us to the conclusion that designing a scalable, distributed WCOJ for a map-reduce like system is an unsolved challenge. We believe it is time to investigate how these algorithms scale in the probably most widely used, general-purpose big data processing engine: Spark. To the best of our knowledge, this would be also the first time a WCOJ is integrated with an industrial-strength cluster computing model. We detail our research questions and goals in ?? and explain how to address challenges in ??.

2 Background

TODO introduction

2.1 Analysis of public real-world graph datasets

TODO will include histogram of graph sizes, average outdeegree, maybe clustering coefficient and further interesting metrics, with regards to the thesis, for (all?) graphs of the SNAP and Labaratory of Web Algorithms dataset collection.

2.2 Compressed sparse row representation

Compressed sparse row representation (short CSR) is a well known, low-memory representation for static graphs [10, 34]. To ease its explanation, we assume that the graph's vertices are identified by the numbers from 0 to |V| - 1. However, our implementation allows the use of arbitrary vertice identifiers in \mathcal{N} by storing the translation in an additional array of size |V|.

CSR uses two arrays to represent the edge relationship of the graph: one of size |E| which is a projection of the edge relationship onto the dst attribute and a second of size |V| + 1| which stores indices into the first array. To find all destinations directly reachable from a source $src \in V$, one accesses the second array at src for the correct index into the first array for a list of destinations.

The CSR format has two beneficial properties in the context of this thesis. First, it allows locating all destinations for a source vertice by one array lookup; hence, in constant time. Second, the representation is only, roughly, half as big than a simple columnar representation. A uncompressed columnar representation needs $2 \times |E|$ while CSR uses only |V| + 1 + |E|, note that for most real-world graph $|V| \ll |E|$ holds (see section 2.1).

3 Related Work

We first provide a short overview of worst-case optimal join algorithms, then we introduce Spark and finally summarize existing distributed implementations of worst-case optimal join algorithms.

The development of worst-case optimal joins started in 2008 with the discovery that the output size of a relational query is bound by the fractional edge number of its underlying hypergraph [6]. In 2012, Ngo, Porat, Re and Rudra published a join algorithm matching this bound [29]. In the same year, Veldhuizen proved that the algorithm "Leapfrog Triejoin" used in LogicBlox, a database system developed by his company, is also worst-case optimal with regards to the fractional edge number bound. Both alorithms have been shown to be instances 'Generic Join' in 2013 Ngo et al. [28]. Leapfrog Triejoin is the only worst-case join algorithm that has been implemented and benchmarked numerous times in widely different settings, e.g. Oxford course work, published research and a commercial database system [35, 2, 30, 18, 3, 33].

Spark is the probably most widely used and industry accepted cluster computing model. It improves over former computing models, e.g. MapReduce [14], Hadoop or Haloop [9], by allowing to cache results in memory between multiple queries, using so-called resilient distributed datasets [36]. However, although Spark implements most operations in memory, shuffles are an exception which still writes and reads the whole intermediary dataset to and from disk. Consequently, shuffling remains to be a highly expensive operation which is the bottleneck of many workflows. Still, Spark provides the user with a general distributed data processing model with strong fault- and struggler tolerance on commodity shared-nothing clusters.

Due to its generality, widespread acceptance in the industry, the ability to use cloud hardware and its fault tolerance by design, it is an attractive target for big graph processing. For example, GraphFrames [13], GraphX [16] (a Pregel [24] implementation) or graph query languages as G-CORE [4] and openCypher with their 'Cyper for Apache Spark' [31] all aim to ease graph processing on Spark. The last two technologies translate their graph specific operations to the relational interface of Spark (SparkSQL) to profit from Spark's relational query optimizer Catalyst [5]. Hence, we believe that the WCOJ's, with their efficiency for analytical graph queries, are a valuable addition to Spark's built-in join algorithms in general and these graph-on-spark systems in particular.

In 2014, a Leapfrog Triejoin variant, dubbed "Tributary Join", was used as a distributed join algorithm on a shared-nothing architecture called "Myria" [11]. They use Tributary Join as a

local, serial worst-case optimal join algorithm, combined with the Hypercube shuffle algorithm to partition the data between their machines [1]. However, it is not obvious how well Hypercube shuffles scales because it replicates many of its input tuples [11]. The combination of Hypercube shuffles and Tributary Join in Myria does not scale well (speedup of 8 on 64 workers compared to the time it takes on 2 nodes) which, although unlikely to be optimal, is not investigated in great detail; we therefore, explain Hypercube shuffles in detail and show that they tend to replicate all data to all nodes for bigger graph patterns section 3.1. Their approach is directly applicable to Spark.

A second distributed version of worst-case optimal joins was published in 2018 based on a Timely Data Flow system [3, 27]. In Timely Data Flow, shuffling is a streaming, asynchronous operation which requires no disk access¹. Therefore, the number of shuffle operations is less important than in Hadoop or Spark. The authors take advantage of this fact by using a uniform, non-replicating partitioning scheme for their relationships. Then they implement a worst-case optimal join using distributed data flow operators [27], e.g. min and intersection. Similar to us, the authors focus on scalability and efficiency in their work but due to the use of a streaming, in memory shuffles and a fine-grained batched processing scheme their approach is unlikely to be successful in Spark. Hence, their and our research share the same goals, however, we aim to achieve it in a more restrictive, but widely used and industrial accepted, system.

3.1 Hypercube shuffle

We first explain how the Hypercube shuffle algorithm, HC, partitions data. After, we provide an analysis of its scaling with regards to graph patterns with an increasing number of vertices.

HC partitions the input relationships for a multi-way join over w worker nodes, such that, all tuples, which could be joined, end up on the same worker in a single shuffle round. Hence, it allows running any multi-way join algorithm locally after one shuffle round. The output of the join is the union of all local results. HC realizes this partitioning by logical organizing all workers in a hypercube with one dimension per join variable; we call the number of variables A and use a_i to reference a single variable. Each dimension has a size p_i , the p_i 's have to be chosen such that the constraint $w \ge \prod_i p_i$ is satisfied. In other words, each worker can be addressed by its coordinate in the hypercube of the form $\{1..p_1\} \times ... \times \{1..p_A\}$.

With this topology in mind, it is straightforward to find a partitioning for all tuples from all relationships such that tuples that could join are sent to the same node. We choose a hash function h_i for each join variable which maps its values in the range of $\{1..p_i\}$. Then each worker determines where to send the tuple it holds by hashing its join variables. This results in a coordinate in the hypercube which is fixed for all join variables, which occur in the tuple, and unbounded for join variables not bound by the tuple. Then the tuple is sent to all workers with a matching coordinate. For example, assume a join with three variables a_1 , a_2 and a_3 and tuple t that binds a_1 and a_2 then we get the coordinates $h_{a1}(t_{a1}) \times h_{a2}(t_{a2}) \times \{0..p_{a3}\}$ and send the tuple to all workers with a coordinate matching the first two attributes and arbitrary third component of the coordinate; the tuple is replicated across p_{a_3} machines.

Next, we analyse the scalability of HC on growing graph patterns - that is, joins over a single relationship, the edge relationship of the graph E and with two variables per atom. In this context, atoms of the join can be seen as the edges of the pattern and variables as vertices. In the following we consider the join query represented by the Datalog rule $Q(a_1, ..., a_A) = R_1(a_1, a_2), ..., R_k(a_{A-1}, a_A)$, we call atoms(Q) the set of all atoms in Q, $p_1(R)$ and $p_2(R)$ the p_i corresponding to the variables of R. We use the method described in [11] to calculate optimal shares allocation $p_1...p_A$.

¹The most commonly used cluster computing engines (Hadoop and Spark) implement shuffling as a synchronizing operation that requires to write and read all tuples from disk.

Pattern	Edges	workload [64]/[128]
Triangle	3	0.18 / 0.12
4-clique	6	0.59 / 0.44
5-clique	10	0.9 / 0.82
House	5	0.42 / 0.32
Diamond	8	$0.76 \ / \ 0.67$

Table 1: Workload, on 64 and 128 workers, in percentage of tuples of the edge table assigned to each worker, using Poison binominial distribution to estimate the workload and the method from [11] to determine the optimal shares configuration.

Each worker receives $\sum_{R \in atoms(Q)} |R|/(p_1(R) * p_2(R))$ tuples under the assumption of uniform data distribution and good hash functions. Our argument is that the tuples of each R are divided onto $p_1(R) * p_2(R)$ workers - the workers that form the hypercube planes of its two variables.

In the special case of graph pattern matching where all atoms of the query are pointing to the same relationship, we can optimize HC shuffle such that a tuple is only sent once to a worker, although it might be assigned to it via multiple atoms. If we apply this optimization, we can predict the probability with which each tuple is assigned to a worker using the Poisson binomial distribution. The Poisson binomial distribution $Pr(n, k, u_0, ..., u_n)$ allows us to calculate the likelihood that k out of n independent, binary and differently distributed trials succeed, under the condition that the i'th trial succeeds with a probability of u_i . We then use n = |atoms(Q)|, k = 0 and $u_i = 1/(p_1(R_i) * p_2(R_i))$ to calculate the probability that a tuple is not assigned to an arbitrary, fixed worker w. This allows us to predict the number of tuples assigned to each worker by $|E| * (1 - Pr(|atoms(Q)|, 0, u_0, ..., u_{|atoms(Q)|})$.

Table 1 shows the expected percentage of tuples from E assigned to each node for graph patterns of different sizes calculated using Poison binomial distribution and optimal shares assignments according to the method used in [11]. As we can see in this table, the number of tuples assigned to each worker grows over linear in the size of the graph pattern and that doubling the number of workers is inefficient to counter this growth.

The second observation has two reasons. First, doubling the number of workers does not allow to double the dimensions of the hypercube - a hypercube always needs $p_1 \times ... \times p_A$ workers to be built. Second, the number of replicated tuples increases with a growing hypercube because each tuple from R_i is replicated to $\prod_{R_j \in atoms(Q)/R_i} p_1(R_j) * p_2(R_j)$; due to the fact that each tuple binds only two out of A variables the tuple is replicated over many dimensions, e.g. let the bounded varibales be a_p and a_s then we get $|\{0..p_0\} \times ... a_p... \times ... a_s... \times \{0..p_A\}|$ matching worker coordinates.

In light of the numbers presented in table 1 and in line [3], we conclude that the communication costs for HC converge towards a full broadcast for bigger graph patterns and scaling becomes increasingly inefficient. Anyhow, HC is proven to be communication cost optimal for general multi-way joins in map-reduce like systems in a single round of shuffling in multiple settings [7, 8, 20]. Therefore, we decide to follow a different direction for distributing WCOJ's which we explain in ??.

4 Implementation

4.1 General sequential version (seq)

We implement the Leapfrog Triejoin [35] as our general sequential version of a WCOJ. However, instead of using B-Trees as a backing data structure, we use sorted arrays and a binary search, which has been described in [11] and is called Tributary join in their paper. Our Leapfrog Triejoin is implemented in three components which we explain in order below: LeapfrogJoin, ArrayTrieIterable and LeapfrogTriejoin.

The Leapfrog join is a variant of the sort-merge join for unary relationships, originally described in [15, 19]. To join k unary relations $A_1(x)$, $A_2(x)$, ..., $A_k(x)$ it takes one iterator per input relations and offers an iterator interface that yields the intersection of all relations. It requires that it's input iterators offer a key method in $\mathcal{O}(1)$, a next method and a leastUpperBound(key: Int) both in $\mathcal{O}(\log n)$ (n defined as the size of the input relationship). leastUpperBound moves the iterator to the first position of the sought key or the first position of the next higher value. An idiotmatic implementation of a Leapfrog join is shown in listing 1, for the optimized implementation see leapfrogTriejoin.LeapfrogJoin in our repository.

```
class LeapfrogJoinIdiomatic(var iterators: Array[LinearIterator]) {
     var atEnd: Boolean = false
     var p = 0
     var key = OL
     def init(): Unit = {
       atEnd = iterators.exists(li => li.atEnd)
       p = 0
       key = -1
        iterators.sortBy(_.key)
11
        leapfrogSearch()
12
13
14
15
     def leapfrogNext(): Unit = {
       iterators(p).next()
       p = (p + 1) % iterators.length
18
       leapfrogSearch()
19
20
21
     def leapfrogLeastUpperBound(key: Long): Unit = {
22
        iterators(p).leastUpperBound(key)
23
       p = (p + 1) % iterators.length
24
       leapfrogSearch()
25
     }
26
27
     private def leapfrogSearch(): Unit = {
28
       if (!iterators(p).atEnd) {
29
          var max = iterators((p - 1) % iterators.length).key
30
          var min = iterators(p).key
31
32
          while (min != max && !iterators(p).atEnd) {
33
            iterators(p).leastUpperBound(max)
34
            max = iterators(p).key
35
            p = (p + 1) % iterators.length
36
            min = iterators(p).key
37
38
          key = min
       atEnd = iterators(p).atEnd
41
42
43
```

Listing 1: Leapfrog join.

To support j-arity relations, $A(a_1, a_2, ..., a_j)$ we add two methods to the iterator interface that represents the input relationships: up and open; both are required to work in $\mathcal{O}(\log n)$. We call

Х	у	Z	Α						
1	1	5		/	^	\			
1	1	6	1			4			
4	5	3	1		5	8	9		
4	8	3			Ĭ				
4	9	3	5	6	3	3	3		

Figure 1: A 3-ary relationship as table (left) and trie (right), to position the iterator at the tuple (1, 1, 5) one calls *open* twice, *key* returns now 5, after a call to *next*, *key* returns 6 and *up* would lead to *key* returning 1.

this new iterator Trieiterator because it represents the relationship as a trie, see fig. 1.

The implementation of a Trieiterator backed by a columnwise representation of the relation using one array per column is straight forward, we outline the basic ideas here and refer the interested reader to leapfrogTriejoin.ArrayTrieIterable.TrieIteratorImpl in our repository for further details. It helps to think about the Trieiterator as consisting out of a linear component, containing the functions key, next and leastUpperBound, and horizontal component, made off the functions up and open, to move the linear component from one trie level to another.

First, we explain the horizontal component. They keep track of the current level of the Trieiterator and the startPosition and endPosition for in the column, e.g. in fig. 1 when the current level is 1 (or x), the key equals 4, the startPosition is 2 and the endPosition is 5 because the value 4 occurs 3 times. With these bookkeeping variables, updated by up and open, one can implement the linear part by a binary search over the current column (given by level) which is limited to startPosition and endPosition.

The Leapfrog Triejoin combines Trielterators and Leapfrog joins to join k relationships of arbitrary arity. Its input is one Trielterator per relationship, with these it builds one Leapfrog join per attribute which receives references to all Trielterator of relationships containing the attribute, e.g. for the triangle query R(a, b), S(b, c), T(a, c) the Leapfrog Triejoin receives three Trielterator, for R, S and T, and builds three Leapfrog joins, for a, b, c, which receive references to two Trielterators each. To generate the join result the Leapfrog Triejoin operates the horizontal components of the Trielterators directly and uses the Leapfrog joins to operate the linear component.

We show an idiomatic implementation of a Leapfrog Triejoin in listing 2 and 3, a performance oriented implementation can be found in our repository in leapfrogTriejoin.LeapfrogTriejoin. These listings contain two important functions: the initialization function from line 3 to line 14 and the *moveToNextTuple* function at line 16. We go through these functions in order.

The initializer gets two arguments: a mapping from variables to TrieIterators (each TrieIterator belongs to the list of attributes of its relationship) and the global variable ordering as a sequence of Strings. First, it creates one LeapfrogJoin per variable (line 5) which receives references to each TrieIterator operating on a relationship with this attribute. Then it builds a mapping from variables to all TrieIterators acting on a relationship with an attribute of the same name (line 8). Finally, it initializes maxDepth, action, depth, bindings and atEnd (line 10 to line 14). depth and bindings are an internal variable storing the index of the variable to bind currently and the current bindings for all variables up to depth. atEnd signals that the join has been completed to the client.

Action	Condition	Next action	Yields
	lf.atEnd	UP	no
NEXT	$\neg lf.atEnd \land reachedMaxDepth$	NEXT	yes
	$\neg lf.atEnd \land \neg reachedMaxDepth$	DOWN	no
DOWN	lf.atEnd $\neg lf.atEnd \land reachedMaxDepth$ $\neg lf.atEnd \land \neg reachedMaxDepth$	UP NEXT DOWN	no yes no
	depth = 0, means highest $lf.atEnd$ is true	- (done)	yes
UP	lf.atEnd	UP	no
	$\neg lf.atEnd$	NEXT	no

Table 2: Summarizes which action follows from the current action under which condition. *If* abbreviates the *LeapfrogJoin* of the current variable. *reachedMaxDepth* is true if we currently find bindings for the last variable in the global order. The columns *Yields* details if the main loop of the state machine yields before computing the next action, this is the case, when the *bindings* array contains a complete tuple.

move To Next Tuple implements a depth-first traversal of the Trie Iterators. This traversal needs to stop each time when a complete tuple has been found to support the iterator interface of the join. Therefore, it is implemented as a state-machine which stops each time the deepest level is reached and all variables bound (loop condition in line 35). The next action of the state machine is determined by the outcome of the current action. Hence, we can characterize the state machine by describing each possible action and its possible outcomes. There are three possible actions: NEXT, DOWN, UP. We summarize the possible actions, conditions for the next action and if the main loop of the state machine yields the next tuple in table 2 and describe each action below.

NEXT moves the LeapfrogJoin at the current depth to the next possible binding for its variable (line 2 in listing 3). If the LeapfrogJoin reached its end, we continue with the UP action (line 4), otherwise we set the binding and continue by another NEXT action, if we are at the deepest level or by moving to the next deeper level by the DOWN action (line 7).

DOWN moves to the next variable in the global variable ordering by opening all related *TrieIter-ators* (line 20) A DOWN can be followed by an UP if the LeapfrogJoin is atEnd (line 22), by a NEXT action if the trie join is at its lowest level (line 25), or by another DOWN to reach the last level.

UP can signal the completion of the join if all bindings for the first variable in the global ordering have been explored, or in other words, the first LeapfrogJoin is atEnd (condition depth == 0 $\land action == UP_ACTION$ line 28). Otherwise, all TrieIterators corresponding to the current variable are moved upwards by calling triejoinUp (line 31) which also updates depth and bindings. Then, this action is followed by another UP or a NEXT depending on atEnd of the current LeapfrogJoin (lines 32).

```
class LeapfrogTriejoinIdiomatic(trieIterators: Map[Set[String], TrieIterator],
                                     variableOrdering: Seq[String]) {
     // One LeapfrogJoin per variable with references to all TrieIterators
     // which relationships have an attribute of the same name
     private val leapfrogJoins: Array[LeapfrogJoin]
5
     // A mapping of each variable to all TrieIterators related to it
     private val variable2TrieIterators: Map[String, Seq[TrieIterator]]
     private val maxDepth = variableOrdering.length - 1
     private var action = DOWN_ACTION // The important line
11
     private var depth = -1
12
     private var bindings = Array.fill(variableOrdering.size)(-1L)
13
     var atEnd: Boolean = trieIterators.values.exists(i => i.atEnd)//
14
15
     def moveToNextTuple(): Array[Long] = {//
16
       if (action == NEXT_ACTION) action = nextAction()
       do {
18
         if (action == DOWN_ACTION) {
19
           triejoinOpen()
20
           if (leapfrogJoins(depth).atEnd) {
21
              action = UP_ACTION
22
           } else {
23
              bindings(depth) = leapfrogJoins(depth).key
24
              action = if (depth == maxDepth) NEXT_ACTION else DOWN_ACTION
           }
26
         } else if (action == UP_ACTION) {
27
           if (depth == 0) {
28
              atEnd = true
29
           } else {
30
              triejoinUp()
31
              action = if (leapfrogJoins(depth).atEnd) UP_ACTION else nextAction()
           }
33
34
       } while (!((depth == maxDepth && bindings(maxDepth) != -1) || atEnd))
35
       bindings
36
     }
37
   }
38
```

Listing 2: Shows the main methods of LeapfrogTriejoin, the initializer and moveToNextTuple functionality helper methods are detailed in listing 3.

```
private def nextAction(): Int = {
     leapfrogJoins(depth).leapfrogNext()
     if (leapfrogJoins(depth).atEnd) {
3
        UP ACTION
4
     } else {
5
       bindings(depth) = leapfrogJoins(depth).key
6
        if (depth == maxDepth) NEXT_ACTION else DOWN_ACTION
     }
   }
9
10
   private def triejoinOpen(): Unit = {
11
     depth += 1
12
     val trieIterators = variable2TrieIterators(variableOrdering(depth)).foreach(_.open())
13
     leapfrogJoins(depth).init()
14
15
16
   private def triejoinUp(): Unit = {
17
     variable2TrieIterators(variableOrdering(depth)).foreach(_.up())
18
     bindings(depth) = -1
19
     depth -= 1
20
   }
21
```

Listing 3: LeapfrogTriejoin helpers.

4.1.1 Optimizations

A simple, idiomatic Scala implementation of the Tributary join is not able to beat Spark's BroadcastHashjoin on any other query than the triangle query. Hence, we spent roughly 2 weeks to optimize our first implementation. After, we are able to beat Spark's BroadcastHashjoin on nearly all queries and datasets. In this section, we discuss the implemented optimization and give a rough estimate of how important each of these is. In total, we improved the WCOJ running time from 248.2 seconds to 44.5 seconds on the unfiltered 5-clique query on the Amazon-0602 dataset. We list all optimizations in table 3 and label them 'very important', 'important' and 'minor' based on the performance improvement directly after applying it.

It is not helpful to give more detailed information on the effect of single optimization because they are not independent of each other. Hence, they might have a hugely different effect when applied in a different order, e.g. we first applied an optimization to the binary search and then optimized the *LeapfrogJoin.next* method to avoid many searches Hence, giving detailed runtime measurements for the binary search optimization would overestimate its value. It is out of the scope of this work to study the dependency and order of the optimization to gain correct runtime measurements.

We discuss the optimization in categories: Leapfrog Triejoin specific, binary search specific, Spark related, Scala related and general. We conclude the section with some changes we tried that do not improve performance.

Binary search specific optimizations become a category on its own because the sorted search is the most expensive operation in the Tributary join. According to profiler sessions, the join spends more than 70% of its time in this method. This result is in line with the observation that 'in the Tributary join algorithm, the most expensive step is the binary search' from [11].

Category	Optimization	Impact
LEGI	LeapfrogJoin.init avoid sorting iterators	very important
LFTJ	$ArrayTrieIterable.next$ in $\mathcal{O}(1)$ for deepest level	very important
Dinany goonah	linear search for short search spaces	important
Binary search	avoiding unnecessary conditions	important
Spark	direct use of arrays instead of ColumnVector	important
	use while instead of map, foreach, exists, etc	very important
Scala	use Array instead of Scala's collections	very important
Scala	use of private[this]	minor
	enable compiler optimization	minor
	remove array lookups from the critical path $column(depth)(position) \rightarrow$	very important
General	current Column (position)	
	use $Array$ instead of Map if keys are integers and dense	important
	strength reduction $(i + 1)$ % 5 \rightarrow if $(i == 4)$ 0 else $i+$ 1	important

Table 3: Summary of all optimizations used for seq and an estimate of their impact.

We applied two Tributary join specific optimizations. The first in the class leapfrogTriejoin.LeapfrogJoin (see also listing 1) and the second in the leapfrogTriejoin.ArrayTrieIterable.TrieIteratorImpl.

The LeapfrogJoin.init method is origingally described in [35] to sort its TrieIterators. However, the method can be improved by avoiding to sort the TrieIterators (line 11). We can start moving the TrieIterator without sorting them and arrive at an ordered array in $\mathcal{O}(n)$ steps - n defined as the size of iterators. This approach improves over the original algorithm in two ways: (1) it starts moving the TrieIterators to their next intersection immediately without sorting them first and (2) orders the array in fewer steps than traditional sorting algorithms.

To implement this we find the maximum key value in all iterators and store the index of this TrieIterator in p. Then we move the TrieIterator at p+1 to the least upper bound of this max (by calling seek) and store the result as the new maximum. We proceed with this process - wrapping p around when it reaches iterators.length - until p equals the original maximum index. Now, we are either in a state in which all TrieIterators point to the same value and we are done - the LeapfrogJoin is initialized - or we arrived at a state in which the iterators array is sorted according to key and can proceed as in the original LeapfrogJoin.init method. To apply this optimization one replaces the call to sort in line 11 with the procedure explained above².

The second Leapfrog Triejoin specific optimization is to change the ArrayTrieIterable.TrieIteratorImpl.next method. This method moves the iterator to the next value on the same level of the trie. Hence, it generally runs in $\mathcal{O}(\log n)$, n being the number of tuples in the relationship, because it needs to find the least upper bound of key + 1. However, under the assumption that all tuples are unique - which is fulfilled for the use-case of an edge relationship - the last level of the trie is unique. Hence, we can move to the next value by simply increasing the position by one, which is an operation in $\mathcal{O}(1)$.

The binary search is the most expensive operation of the Leapfrog Triejoin. Hence, special attention needs to be paid while implementing it. Our most important optimization is to change to a linear search once we narrowed the search space to a certain threshold - currently at 60 values. We experimented with values from 0 to 400 and found that 60 was optimal but even

²The implementation of Scala's array sort for objects is slow because it copies the array twice and casts the values to *Java.Object* such that it can use Java's sorting methods. Before we applied the sorting optimization above, we replaced Scala's sort method with an optimized insertion sort, which was faster than Scala's sorting method - the *iterator* array contains normally at most 20 items.

going as high as 120 values would not change the performance much.

Another important optimization is to avoid unnecessary if-statements in the loop of the binary search, e.g. the implementation on Wikipedia and many other example implementations use an if-statement with three branches for smaller, bigger and equal but two branches for greater than and less-or-equal suffice for a least upper bound search.

A similar optimization can be applied to a linear search on a sorted array: intuitively one would use the while-loop condition $array(i) > key \land i < end$ with key being the key to find the least upper bound for, i the loop invariant and end the exclusive end of the search space. Anyhow, it is faster to check for key > array(end - 1) once before the loop and return if this is the case because the value cannot be found in the search space. This obviously circumvents the main loop of the linear search; additionally, it simplifies the loop condition to array(i) > key.

The Spark infrastructure uses the interface ColumnVector to represent columns of relationships. The implementation OnHeapColumnVector is a simple wrapper around an array of the correct type with support for *null* values and *append* operations. First, we used this data structure to represent our columns but we could see a clear increase in performance by replacing it by an implementation that exposes the array to allow the binary search to run on the array directly. This is likely due to saving virtual function calls in the hottest part of our code. The implementation is straightforward and can be found in our repository in leapfrogTriejoin.ExposedArrayColumnVector; we implemented it only for the Long datatype.

We found many standard optimizations and Scala specific optimizations to be really useful. Most likely these are the optimizations that brought the biggest performance improvements. However, they are well-known, so we mention them only in the table 3. For Scala specific optimizations one can find good explanations at [12].

Apart from the aforementioned very useful optimizations, we investigated multiple other avenues in hope for performance improvements which did not succeed, we list these approaches here to save others the work of investigating:

- reimplement in Java
- use of a Galloping search before the binary search
- unrolling the while-loop in Leapfrog Triejoin.move To Next Tuple
- predicating the action variable in Leapfrog Triejoin.move To Next Tuple

Finally, we believe that code generation for specific queries that combines the functionality of LeapfrogTriejoin, LeapfrogJoin and ArrayTrieIterator into one query specific function would lead to noticeable performance improvements. The reason for this belief is that our implementation takes about 3.46 seconds for a triangle query on the Twitter social circle dataset while a triangle query specific Julia implementation, of a colleague of ours, needs only half a second. The main difference between our implementation and his are: the language used (Julia is a high-performance, compiled language) and the fact that his implementation has no query interpretation overhead but cannot handle any other query than the triangle query.

However, a code generated Leapfrog Triejoin is out of scope for this thesis, also, we are aware of efforts by RelationalAi to write a paper about this specific topic. We are looking forward to seeing their results.

4.2 User interface

```
val sparkSession = SparkSession.builder
  .master("local[1]")
  .appName("WCOJ-spark")
  .getOrCreate()
// read a dataframe
val df = sparkSession.read
  .option("inferSchema", "true")
  .csv("/path/to/edge/relationship")
// df needs columns called 'edge_id', 'src' and 'dst'
// Use WCOJ to find a triangle pattern
val triangles = df.find(
  """(a) - [] -> (b);
    |(b) - [] -> (c);
    |(a) - [] -> (c)""".stripMargin,
  Seq("a", "b", "c")
triangles.limit(10).show()
// Shows a dataset with 3 columns: 'a', 'b', 'c' being node ids
```

Listing 4: Example usage of a WCOJ to find triangles in graph.

As one can see in line 16 of listing 4, we support a clean and precise DSL to match patterns in graphs. This DSL is inspired by GraphFrames [13]. The user can define a pattern by its edges, each edge is written as (a) - [] -> (b) where a is the source vertice and b is the destination, multiple edges are separated by a semicolon. A connected pattern is expressed by defining multiple edges with the same source or destination. One should be aware, that a named source or destination is not guaranteed to be a distinct element in the graph, e.g. (a) - [] -> (b); (b) - [] -> (c) could be a linear path of size two or a circle between a and b; in the second case a and c are the same element. The reader might wonders, why we chose to stay with the GraphFrame syntax for edges of - [] ->, although, we could have went with something simpler, like ->. However, sticking to the more verbose syntax allows us to include labels inside of the squared brackets in future extensions, e.g. for our stretch goal of integration with CAPS.

The second parameter to findPattern allows the user to specify the variable ordering used in the WCOJ algorithm. Furthermore, the user interface takes multiple optional arguments, e.g. to apply to common filters to the output of the result, specify different relationships to be used as input for each edge of the pattern. The filters are distinctFilter, ensuring that each vertice can occur only as binding for one variable, and smallerThanFilter to allow only output bindings were the values decrease with regards to the specified variable ordering, e.g. the binding [1, 2, 3] but not [2, 1, 3] for the triangle query above. We experienced that these queries are typical for graph queries and that the performance greatly benefits from pushing them into the join. Implementing the possibility to push general filters into the join would be a valuable addition but we decided against it because it a pure engineering task.

4.3 Spark integration

We integrated our WCOJ implementation into Spark such that it can be used as function on Datasets. Therefore, we build all components necessary to execute a WCOJ in Spark's structured queries, provided by Catalyst, this includes a LogicalPlan, a Strategy to convert this logical operator into a SparkPlan, a physical operator to materialize RDD's in sorted fashion into a columnar representation that supports a least upper bound operation in $\mathcal{O}(\log n)$ and finally a general implementation of Spark's zipPartition operation to support more than 5 children. We explain these components below; before we outline some limitations of our integration.

It is not in the scope of this work to integrate WCOJ's into the SQL parser of Spark. Hence, WCOJ can only be used by Spark's Scala functional interface and not through Spark's SQL queries.

We do not integrate it into the query optimization components of Catalyst, e.g. we do not provide rules or cost-based strategies to decide when to use a WCOJ or a binary join. It is up to the user to decide when to use a WCOJ or a binary join. However, our integration allows the user to intermix these freely. The reason for this decision is, that at time of writing no published paper existed that systematically studies which queries benefit from WCOJ's in general, nor, does research exist that studies the combination of WCOJ's and binary joins. Only a month after we decided on our scope for Spark integration Salihoglu et al. published an arXiv paper [26] that tackled these problems for the first time. The lack of peer-reviewed papers and the high complexity of the arXiv paper clearly illustrate that deeper integration with Spark's optimizer is out of scope for this thesis.

To allow the user to call findPattern (as defined in section 4.2) on a Dataset, we added a new class to the package org.apache.spark.sql called WCOJFunctions and an implicit conversion from Dataset to WCOJFunctions. This function parses the pattern - the necessary code is borrowed from GraphFrame [13] - into a representation of a WCOJ join, JoinSpecification, and creates new AttributeReferences named accordingly to the join variables.

Due to our decision not to build optimizer rules to introduce WCOJ's, we have to implement a new *LogicalPlan* to represent WCOJ's. This plan has as many children as relationships joined by the WCOJ. Its *output* are the AttributeReferences produced by findPattern.

The physical operator for WCOJ's, WCOJExec, is an implementation of the trait SparkPlan. It expects as many children as the relationships in the join; these children have to support a Trieiterator interface as specified in section 4.1. The biggest challenge in the implementation of WCOJExec is that Spark does not support to zip the partitions of more than 5 RDD's. However, to operate synchronously on the partitions of all children, we need this operation for an arbitrary number of children. Fortunately, although, not publicly available, Spark does implement *zipPartitions* in a general manner in the abstract class ZippedPartitionsBaseRDD. Hence, supporting this functionality is a simple matter of subclassing this class publicly to expose its generality to our code (see sparkIntegration.GeneralZippedPartitionsRDD).

We need to materialize the input RDD's for WCOJExec, to support the Trieiterator interface. This is done by ToTrieIterableRDD, a SparkPlan, with a single child. It iterates over the child and stores all tuples in a columnar representation, a ColumnarBatch³ ⁴. Furthermore, our Trieiterator

³To all but highly experienced Spark developers, the construct used to allow us to create an RDD which controls how its partitions are stored might look unintuitive. Normally, the type parameter of the RDD class is used to refer to the type of its elements, e.g. Sparks Catalyst module uses mostly RDD[InternalRow]. However, it is possible - and practised in the Spark project itself - to use this type parameter to give the type of the partitions of an RDD. This is what we do in the class TrieIterableRDD, which at the same time is a subclass of RDD[InternalRow], so it can be used in Catalyst.

⁴Later we noticed the RDD function glom which allows to operate on the partitions of RDD's an array. However, by then we had already implemented our own solution and decided to keep it for better control.

implementation demands that the tuples are sorted. Therefore, ToTrieIterableRDD requires its child to be ordered using the requiredChildOrdering functionality of Spark.

The last component of our Spark integration is a simple Strategy to translate a WCOJ logical operator into a WCOJExec physical operator. This is a one-to-one translation from WCOJ to WCOJExec as an implementation of Spark's abstract class SparkStrategy. Additionally, this Strategy also introduces a ToTrieIterableRDD operation before the WCOJ for every child.

The Spark integration of this work has been inspired by the IndexedDataframes [indexed-dataframes] work of Alex Uta and the supervisors of this thesis.

4.4 Operator to create a CSR from a DataFrame

To be defined and written.

4.5 Graph pattern sequential version (graph-pattern-seq)

4.5.1 Combining LFTJ with CSR

For our graph pattern matching specialized LeapfrogTriejoin version we choose CSR (see section 2.2) as backing data structure. This data structure is typically used for static graphs and we show that it is a great match for LFTJ. In this section, we shortly describe the implementation of a CSR based TrieIterator, point out the differences between this new version and a column based TrieIterator (as described in section 4.1⁵) and conclude with an experiment demonstrating the power of this optimization.

The implementation of a CSR based TrieIterator is straightforward except for one design change: instead of using the vertice identifier from the graph directly, we use their indices in the CSR representation. This change is rather minor because it can be contained at any level by using a hash map for translation, e.g. in the TrieIterator itself, in the LeapfrogTriejoin or at the end of the query by an additional mapping operation. In the current system, the translation is performed by the LFTJ implementation to allow easy integration into other projects. However, it is possible to work on the indices throughout the whole system to safe the translation costs. We now outline how to implement each of the TrieIterator methods, under the assumption that all vertices have outgoing edges. Then, we drop this assumption and explain the necessary changes. The creation of the CSR data structure itself is described in section 4.4^6 .

We use the variables depth, srcPosition and dstPosition to store if we are operating on the source or destination level and the positions of the iterator on the respective level. We call the two arrays of the CSR datastructure dst for the array storing all edge destinations and dstIndices for the array storing indices into dst (refer to ?? for a complete explanation of these arrays). The open function does nothing if we open the first level; if we open the second level, it sets dstPosition to dstIndices(srcPosition). Additionally, it updates depth. The up function only updates depth. The next method increases srcPosition or dstPosition by one depending on depth. The seek(key) method sets srcPosition to key if depth equals 0 or uses binary search to find key in dst.slice(dstPosition, <math>dstIndices[srcPosition + 1]) and then sets dstPosition accordingly. The key function returns srcPosition on the first level and dst[dstPosition] if depth equals 1. The atEnd function is: if (depth == 0) srcPosition == dstIndices.length - 1 else dstPosition == dstIndices(srcPosition + 1).

< Alternative representation of the TrieIterator implementation as table. Let me know what you

⁵To be rewritten and integrated

⁶To be written.

prefer. I think the text is better after seeing both.>

Method	1st level	2nd level
open	update depth	$dstPosition \leftarrow dstIndices[srcPosition]; update depth$
up	update depth	update depth
next	srcPosition++	dstPosition++
seek(k)	$srcPosition \leftarrow k$	$dstPosition \leftarrow lub(k, dst[dstPosition:dstIndices[srcPosition +1]]$
key	srcPosition	dst[dstPosition]
atEnd	srcPosition = dstIndices.length - 1	dstPosition = dstIndices(srcPosition + 1)

Table 4: Tabular summary of TrieIterator implementation based on CSR. lub is leastUpperBound. dst[<start>:<end>] slices the array.

To resolve the assumption of no empty outgoing adjacency lists, we adapt *open*, *next* and *seek* to skip source positions without outgoing edges. This is easy to detect because then dstIndices(x) = dstIndices(x + 1). We can skip these cases by simple linear search until we find a valid position. This solution is sufficient because there are only a few vertices with no outgoing edges in real-world graphs.

The TrieIterator implementation based on CSR is much faster than the column based iterator; mainly due to the fact that the seek method on the first level can be implemented in $\mathcal{O}(1)$, instead of $\mathcal{O}(\log n)$. This optimization has huge potential because these searches are the most costly operations for a column based TrieIterator [11]. Note that searches on the second level are fast, due to the fact that most graphs have a low outdegree (see section 2.1). Additionally to this advantage, CSR based TrieIterator do less bookkeeping because they support only 2 levels and spent nearly no time on processing atEnd for the second level, while a column based TrieIterator needs to calculate the number of outgoing edges for each source vertice in its open method, to allow a fast atEnd method).

We conclude that CSR based *TrieIterator*'s are a promising match for LFTJ and graph pattern matching. The improvements of this optimization can be seen in fig. 2. It demonstrates an up to 2.6 speedup over a column based LFTJ. We also see that the optimization has a stronger impact on queries with more edges and vertices, e.g. 5-clique. For a more thorough evaluation refer to the experiment section 5.

4.5.2 Exploiting low average outdegrees

In our study of real-world graphs (??), we show that most real-world graphs have a small, average outdegree. The outdegree over all graphs is TODO and the maximum average outdegree of a single graph is TODO at TODO. These results lead to the hypothesis that the intersection of multiple adjacency lists is small, e.g. below 10 in many cases. We can exploit this fact by materializing the intersections in the *Leapfrog joins* directly in one go; instead of, generating one value at-the-time in an iterator like fashion as described in the original paper [35]. This is beneficial because it makes better use of data locality; we elaborate this statement in a later paragraph.

We structure the remainder of this section as follows. First, we shortly reiterate the most important facts about *Leapfrog joins* for this chapter (TODO just organize both sections behind each other?). Second, we analysis the intersection workload in terms of input sizes and result size to confirm our hypothesis and gain valuable insights to choose the best intersection algorithm. Third, we explain the algorithm we chose based on the analysis. Fourth, we point out differences

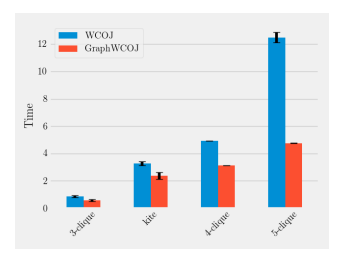


Figure 2: Barchart comparing join time of WCOJ and GraphWCOJ on multiple queries on SNB-sf1

to the original Leapfrog Triejoin. Finally, we present a short experiment showing the performance gains of this optimization.

Leapfrog joins build the intersection between multiple adjacency lists. This is done in an iterator-like fashion in their leapfrog_next method by repeatedly finding the upper-bound for the largest value in the lowest iterator. This algorithm is asymptotically optimal for the problem of n-way intersections. However, we claim that it is (1) to complex for small intersections and (2) should generate all values at once instead of one-by-one to improve performance on real-world adjacency lists.

To determine the best algorithms to build the n-way intersection in the *Leapfrog joins*, we run some experiments to characterize the workload. Towards this goal, we log the size of the full intersection, the size of the smallest iterator participating and the size of the largest intersection between the smallest iterator and any other iterator on 5-clique queries on SNB-sf-1. Figure 3 depicts these metrics as cumulative histograms. In the next paragraphs, we point out the most important observations in each of these graphs.

Figure 3c shows the size distribution of the smallest iterator, as to be expected for a social network graph, the outdegree is between 1 and 200. We do not see the long-tail distribution typical for power-law graphs because we choose the smallest iterator out of 5 and even though there are vertices with a much higher outdegree, the chance of encountering 5 of these in a single intersection is small. We note that in 80% of all cases the smallest iterator has a size lower than 80 and above that the distribution slowly increases to 100%

Figure 3b illustrates the size distribution of intersecting the smallest iterator with any other iterator, such that the intersection is maximal. We choose this specific metric to motivate one of our design choices later on. As for the smallest iterator, some of these intersections are as big as 200 but most of them are much smaller. However, unlike for the smallest iterator metric, 80% of the intersections contain less than 21 elements and the frequency increases to 100% in a steep curve. This last observation is even stronger for the size of the total intersection (fig. 3a): the size is less than 5 in 80% of all intersections and increases similarly steep to 100%. The maximum is a little lower than 200.

These observations confirm our hypothesis that the size of the intersections is small (below 5) and do not show the same long-tail distribution as the whole social network graph. Hence, we can materialize them without running at risk of building big intermediary results. Furthermore, the experiment shows that optimizing by taking iterator sizes into account is worthwhile but only for the smallest iterator because once we start with the smallest iterator the further intersections

are small (below 21) in the vast majority of all instances.

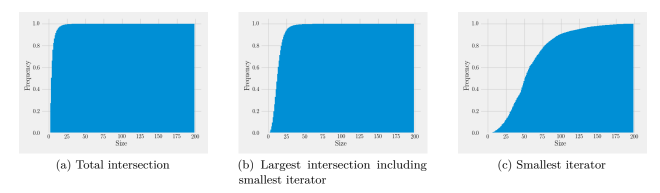


Figure 3: Cumulative histograms of total intersection sizes, largest intersection with the smallest iterator and any other, and size of the smallest iterator participating in 5-clique on SNB-sf-1.

In the coming paragraphs, we detail how to build the n-way intersection of multiple adjacency list such that we gain performance by better use of data-locality than original the *Leapfrog join*. We choose to use pairwise intersections over multi-way intersection algorithms for their simpler, linear memory access patterns.

From our analysis, we conclude that the final intersection size is strongly dependent on the smallest iterator and that the intersection of the smallest iterator with any other iterator is close to the final size. These insights translate into two design decisions. First, we start with the smallest iterator. However, we do not take the sizes of any other iterators into account because the effort for sorting the iterators by size would not pay off.

Second, we use two different tactics to build the pairwise intersections. The first intersection between two iterators is built *in-tandem*, where we seek the upper bound of the higher value in the smaller iterator. This algorithm guarantees to find the intersection between two iterators in the asymptotical fastest way. After this first intersection, the intermediary result is quite small. Therefore, we use the simpler scheme of linearly iterating the intermediary and probing the iterator by binary search with fall-back to linear search.

Finally, we point out a few exceptional cases and pitfalls for implementors:

- If all iterators of the *Leapfrog join* are on their first level, the intersection is near |V|. In this case, we fall back to the original *Leapfrog join*.
- We use an array to materialize the intersections because Scala collections are slow. Instead
 of deleting elements, we replace them with a special value.
- Allocating a new array for every Leapfrog join initialization is costly. We estimate the size
 of the intersection by the size of the smallest iterator and reuse the array whenever possible.
 We use a sentry element to mark the end of the array.

The carefully chosen operations explained above are faster than the original *Leapfrog join* algorithm for two reasons. First, although, the original algorithm uses an asymptotical better n-way intersection, multiple binary intersections are preferable with the rather small adjacency lists of graph workloads. In particular, in our situation where the size of the first binary intersection is already very close to the size of the final intersection.

Second, the *Leapfrog join* generates a single value, then yields control to other parts of the *LeapfrogTriejoin* algorithm and later touches the same adjacency lists again to generate the next

⁷We take advantage of the fact that CSR allows us to determine the size of iterators cheaply (see ??)

value. Our approach touches the adjacency lists exactly once per *Leapfrog* initialization and condenses the intersection into a much smaller array. This array is more likely to stay cached while the other parts of the *LeapfrogTriejoin* do their work.

Figure 4 shows that a materialized *Leapfrog join* out-performance better than the original algorithm. As expected, the optimization is more powerful for bigger queries because they work with more adjacency lists per *Leapfrog join*, e.g. for a triangle each *Leapfrog join* intersects only two iterators while for a 5-clique each join handles 5 adjacency lists. Anyhow, even for the triangle query, we see a small but clear improvement and a lower error due to better cache use. We refer the reader to our experiment section (5) for further experiments on our graph specialized wcoj and detailed descriptions of the datasets and queries used.

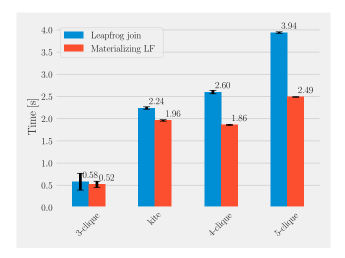


Figure 4: Barchart showing GraphWCOJ with and without *Leapfrog join* materialization enabled for different queries on SNB-sf1.

5 Experiments

TODO introduction

5.1 Setup

We run Spark 2.4.0. We use Scala 2.11.12 on Java openJDK 1.8.

So far, we run three different join algorithms. Our sequential implementation of Leapfrog Triejoin (see section 4.1), called seq and Spark's two different binary join algorithms: BroadcastHashJoin and SortMergeJoin. In the majority of our experiments, we use Spark in its standard configuration with enabled code generation. We also tune the parameters for driver and executor memory usage (spark.driver.memory and spark.executor.memory) to fit all necessary data into main memory. We adjust the "spark.sql.autoBroadcastJoinThreshold" parameter to control if Spark is using a BroadcastHashJoin or a SortMergeJoin.

5.1.1 Hardware

We run our experiments on machines of the type brick of the Scilens cluster owned by the CWI Database Architecture research group. These machines feature 32 Intel Xeon E5-2650 processors

Name	Variant	Vertices	Edges	Source
SNB	sf1		453.032	[21]
Amazon	0302	262,111	1,234,877	[22]
	0601	$403,\!394$	3,387,388	[22]
${f Twitter}$	$\operatorname{sc-d}$	81,306	1,768,149	[22]
	sc-u			[22]

Table 5: Summary of all datasets mentioned in the thesis. Explanation of them and for the variants is given in running text.

with a 32 KB 1st level cache, 250 KB 2nd level cache and 20 MB of shared third level cache and run at 2.0 to 2.8 GHz. The main memory consists of 256 GB of RAM DDR-3 memory.

5.1.2 Datasets

We run the majority of our experiments on two datasets from different use-cases, social networks and product co-purchase. We motivate our choice in the next paragraph. Table 5 includes a list of all graph datasets mentioned throughout the thesis.

The SNB benchmark [21] generates data emulating the posts, messages and friendships in a social network. For our experiments, we only use the friendships relationship (person_knows_person.csv) which is an undirected relationship. After generation only edges of the kind src < dst exist, we generate the opposing edges before loading the dataset, such that the edge table becomes truly undirected. The benchmark comes with an extensively parameterizable graph generation engine which allows us to experiment with sizes as small as 1GB and up to 1TB for big experiments and different levels of selectivity. The different sizes are called scale-factor or sf, e.g. SNB-sf1 refers to a Social network benchmark dataset generated with default parameters and scale-factor 1. We include the exact parameter used for generation in our repository under experiments/snb/params.txt.

The Amazon co-purchasing network contains edges between products that have been purchased together and hence are closely related to each other [22]. This is a directed relationship from the product purchased first to the product purchased second, both directions of an edge can exist if the order in which products have been purchased varies. The Snap dataset collection contains multiple Amazon co-purchase datasets, each of them containing a single day of purchases. We choose the smallest and biggest dataset from the 2nd of March and the 1st of June 2003, we call them Amazon-0302 and Amazon-0601. We pick co-purchase datasets for evaluation because former work often concentrated on social networks and web crawl based graphs [11, 3] but [32] points out that the biggest graphs are actually graphs like the aforementioned Amazon graph containing purchase information.

To allow comparisons with former work, we run a subset of our experiments on the Twitter social circle network from [22]. This dataset includes the follower relationship of one thousand Twitter users; each of these follows 10 to 4.964 other users and relationships between these are included. The graph is originally directed but for some experiments, we add reversed edges to make the graph undirected - again for comparison with former work. We call this graph Twitter-sc-d and Twitter-sc-u for the directed respectively undirected variant.

Name	Parameters	Vertices	Edges	Example pattern
triangle	NA	3	3	$a \rightarrow b; a \rightarrow c; b \rightarrow c$
n-clique	# vertices	n	$1/2 \times n \times (n-1)$	see above
n-cycle	# vertices	n	n	$a \rightarrow b; b \rightarrow c; c \rightarrow z; z \rightarrow a$
n-s-path	# edges / selectivity	n	n-1	$a \rightarrow b; b \rightarrow c; c \rightarrow z$
house	NA	5	9	$a \rightarrow b; a \rightarrow c; a \rightarrow d; b \rightarrow c; b \rightarrow$
				$d; c \rightarrow d; c \rightarrow e; d \rightarrow e$
Filters				
distinct				$a \neq b$; $a \neq c$; $a \neq d$; $b \neq c$;
lt				$a < b; b < c; c < d; \dots$

Table 6: Summary of patterns and filters used.

5.1.3 Graph patterns

In this section, we detail the graph patterns used throughout our experiments. Most of the queries are cyclic because that has been shown to be the primary use-case for WCOJ in former research [30, 11]. WCOJ's also have been successfully applied to selective path queries in [30]; however, this result have not been reproduced by any other paper.

To most of our queries, we apply a filter to make them more realistic, e.g. a clique query does make more sense if it is combined with a smaller-than filter, which requires that the attributes are bound such that a smaller than b, smaller than c. Because otherwise, one gets the same clique in all possible orders in the output, which not only takes much more time but is also most likely not the result a user would want. We ensure that filters can be pushed down through or in the join by Spark as well as by the WCOJ to compare both algorithms on an equal basis. A complete list of all queries and filters used is shown in table 6. The less known queries are also detailed in text. Patterns and filters might be used in all possible combinations, we name the resulting query <pattern>-<filter>-<filter>, e.g. triangle-tt.

For a selective path query, we first select two sets of nodes with respect to the *selectivity* parameter. Then we search for all path of a certain length according to the *edges* parameter, e.g. 4-0.1-path finds all paths between two randomly selected, fixed sets of vertices of length 4 - the sets of nodes contain roughly 10% of all input nodes and are not guaranteed to be intersection free.

5.2 Baseline: BroadcastHashJoin vs seq

In this experiment, we compare the runtime of our sequential Leapfrog Triejoin implementation, seq, with the runtime of Spark's BroadcastHashjoin. Towards, this goal we ran all queries from table 6 on our three main datasets: ama-0302, ama-0601 and snb-sf1. The clique patterns are combined with the less-than filter, n-clique-lt, and the cycle pattern with the distinct filter, n-cycle-distinct. These seem to be the most realistic setups because cliques are fully symmetric and one wants to avoid redundant results. For cycles, the less-than filter is too restrictive because it excludes cycles for which a < b > c. We show our results in table 7 and in barcharts (fig. 5, 6 and 7). Section 5.2.2 analyzes the results.

Our experiment measures the time it takes to perform a count on the cached dataset using BroadcastHashjoin and seq. For BroadcastHashjoin, the time to run the whole query is reported. For seq, we report setup time and the time, it takes to run the join, separately. Setup time includes the sorting, materialization and copying the results of our join from a Scala Array

into the UnsafeInternalRow format expected by Spark. TODO sorting not yet (data is presorted in files) This section is focused on comparing the runtimes excluding the setup time - rational given in section 5.2.1.

5.2.1 Experiment Rationale

Question: Why do we compare against Spark's BroadcastHashjoin instead of SortMergeJoin? Answer: Because even when all data is arranged in a single partition, for simple sequential processing, Spark schedules its SortMergeJoin to use a shuffle. A shuffle writes and reads data to and from disk. Hence, SortMergeJoin is much slower than a BroadcastHashJoin. We compared the algorithms on the Amazon-0601 dataset for the triangle (8.1 seconds vs 58.9 seconds) and 5-clique pattern (32.9 seconds vs 850.9 seconds). We assume that Spark is able to optimize its broadcasts when local[1] is used to start the Spark session because then Spark uses the driver as executor.

Question: Why do we exclude setup times from the WCOJ times?

Answer: Because our final implementation **dist** is meant to cache the readily sorted and formatted edge tables and reuse it for multiple queries. We anticipate that this is necessary to benefit from WCOJ's in general. Furthermore, we optimize the setup times in later implementation. Hence, the current setup code is much slower than the one we expect to use for later implementations.

Question: Why is the time to copy results into UnsafeInternalRow format for WCOJ counted as setup time?

Answer: It is time solely spent for integration with Spark and not Leapfrog Triejoin specific. Furthermore, it could be avoided by working directly on the UnsafeInternalRow format within our seq implementation. However, this would require us to work with unmanaged memory (the UnsafeInternalRow interface is slower than working on Arrays) and we deem this as an unnecessary engineering overhead.

Question: Is Spark's code generation a huge advantage for the BroadcastHashjoin? Answer: Yes, we ran Spark without code generation for comparision on the Amazon-0302 dataset for triangle and 5-clique: with code generation Spark takes 3.1 and 4.2 seconds without 14 and 16.

5.2.2 Analysis

For now, we settle to simply point out the most important observations and postpone deeper analysis, e.g. influence of dataset size and characteristics, to experiments run against our later implementations, i.e. seq-graph-pattern and dist.

We are able to beat Spark's BroadcastHashjoin on all datasets and queries except 5-clique-lt on Amazon-0602. Generally, we see that for n-clique patterns the speedup over Spark decreases for bigger n. This is due to the fact that many binary joins in a n-clique are actually semi-joins which to do not increase but decrease the size of intermediary results, e.g. for 5-clique on Amazon-0302 only 3 out of 9 joins lead to a bigger intermediary result.

The cycle query results are highly interesting because we see an increasing speedup for higher n on Amazon-0602 but a decreasing speedup on Amazon-0302. Unfortunately, we are (TODO currently) not able to provide n-cycle results for the SNB-sf1 dataset, due to the fact that BroadcastHashjoin's take more than 22 hours for the 6-cycle which blocked our experiments. 5-cycle runs at the moment.

The House and 5-clique pattern seem to be quite similiar - the House is a 5-clique with two missing edges. However, as the count of their results indicates these two edges lead to dramatically

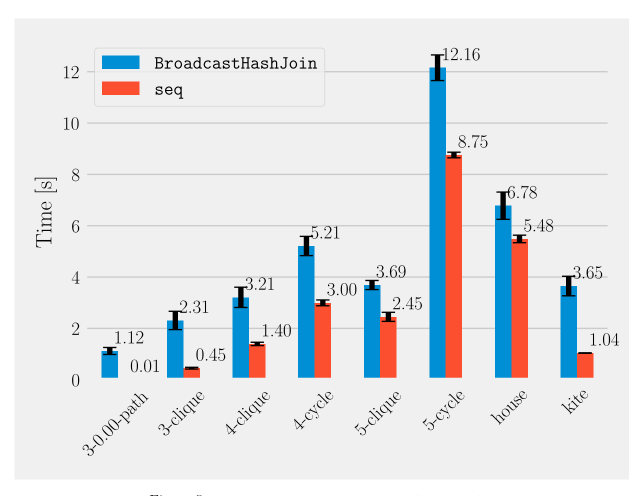


Figure 5: seq vs BroadcastHashJoin on Amazon-0302

different outcomes. Hence, their different timing and speedup behaviour.

The Kite pattern produces consistently the second highest speedup after the 3-clique. Most likely due to the fact that a Kite is two triangles back-to-back.

The path query shows very different behaviour on the Amazon and the SNB datasets. This might be due to the different selectivity; it is extremely high on the co-purchase datasets and rather low on the social network benchmark. This different in selectivity is not surprising given that the SNB network fulfills the small world property, while the Amazon dataset relates products purchased together which naturally leads to multiple loosely connected, denser components. We will run the path queries with a different selectivity on the two input vertice sets to confirm this hypothesis.

Finally, we observe that all three datasets lead to quite different results which are most likely not comparable to each other without deeper research in the characteristics of the datasets themselves. In particular, it becomes clear that co-purchase datasets and social network datasets must have very different characteristics. Although, SNB-sf1 is much smaller than Amazon-0601, queries on it take a similar or even much more time, e.g. 5-clique-lt takes 14.21 seconds on the bigger dataset and 12.65 seconds smaller, even though, the result set is much smaller on SNB-sf1; 4-cycles-distinct takes roughly 8 times longer on the small dataset and has a much bigger result set. In general, we see a higher speedup on SNB-sf1

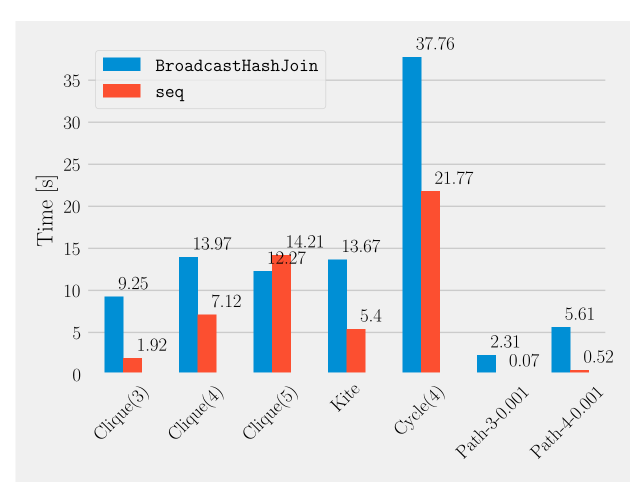


Figure 6: seq vs BroadcastHashJoin on Amazon-0601

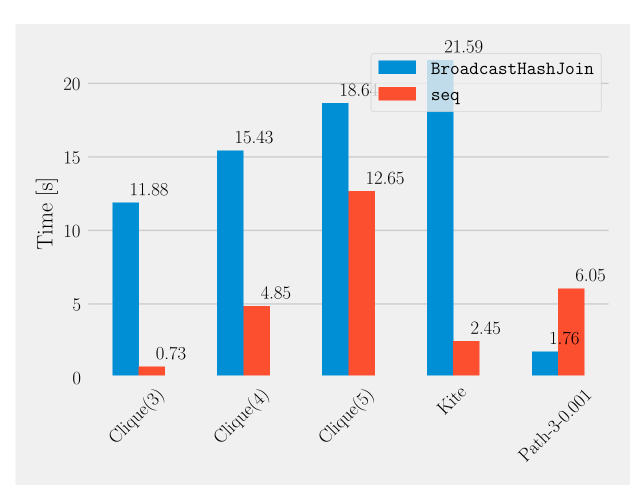


Figure 7: seq vs BroadcastHashJoin on SNB-sf1

Query	# Result	BroadcastHashJoin	seq	setup	Speedup
Clique(3)	188,705	3.08	0.49	2.51	6.3
Clique(4)	47,824	3.85	1.25	4.55	3.1
Clique(5)	7,064	4.22	2.32	7.60	1.8
House	2,941,664	7.87	5.66	6.18	1.4
Kite	$220,\!637$	3.82	1.04	6.73	3.7
Cycle(4)	3,076,324	4.90	2.75	3.40	1.8
Cycle(5)	6,389,425	13.15	8.28	4.87	1.6
Cycle(6)	11,682,732	38.00	27.62	6.79	1.4
Path-3-0.001	2,092	1.36	0.01	0.89	136.0
Path-4-0.001	8,829	3.11	0.03	1.58	103.7

Query	# Result	BroadcastHashJoin	seq	setup	Speedup
Clique(3)	1,137,579	9.25	1.92	6.23	4.8
Clique(4)	911,548	13.97	7.12	11.74	2.0
Clique(5)	585,171	12.27	14.21	19.23	0.9
House	177,745,510	66.53	57.37	23.45	1.2
Kite	3,672,375	13.67	5.40	15.93	2.5
Cycle(4)	$45,\!175,\!984$	37.76	21.77	14.12	1.7
Cycle(5)	263,436,965	227.97	115.60	24.96	2.0
Cycle(6)	1,484,438,088	2020.04	872.62	83.12	2.3
Path-3-0.001	98,929	2.31	0.07	2.30	33.0
Path-4-0.001	922,888	5.61	0.52	4.25	10.8

Query	# Result	BroadcastHashJoin	seq	setup	Speedup
Clique(3)	540,225	11.88	0.73	1.09	16.3
Clique(4)	260,786	15.43	4.85	2.33	3.2
Clique(5)	$40,\!137$	18.64	12.65	3.08	1.5
House	100,206,468	300.13	85.18	6.58	3.5
Kite	1,780,235	21.59	2.45	2.20	8.8
Cycle(4)	232,636,376	636.77	162.89	10.68	3.9
Path-3-0.001	65,290,479	1.76	6.05	3.28	0.3
Path-4-0.001	7,235,522,926	111.73	655.52	293.27	0.2

Table 7: Runtimes for BroadcastHashJoin and seq. The speedup is calculated between join times and excludes setup. From top to bottom for dataset: ama-0302, ama-0601 and snb-sf1. All times in seconds.

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