

SYNCSIGNATURE: A Simple, Efficient, Parallelizable Framework for Tree Similarity Joins

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ABSTRACT

This paper introduces SYNCSIGNATURE, the first fully parallelizable algorithmic framework for tree similarity joins under edit distance. SYNCSIGNATURE makes use of implicit-synchronized signature generation schemes, which allow for an efficient and parallelizable candidate-generation procedure via hash join. Our experiments on large real-world datasets show that the proposed algorithms under the SYNCSIGNATURE framework significantly outperform the state-of-the-art algorithm in the parallel computation environment. For datasets with big trees, they also exceed the state-of-the-art algorithms by a notable margin in the centralized/single-thread computation environment. To complement and guide the experimental study, we also provide a thorough theoretical analysis for all proposed signature generation schemes.

PVLDB Reference Format:

Nikolai Karpov and Qin Zhang. SYNCSIGNATURE: A Simple, Efficient, Parallelizable Framework for Tree Similarity Joins. PVLDB, 16(2): 330 - 342, 2022.

doi:10.14778/3565816.3565833

PVLDB Artifact Availability:

The source code, data, and/or other artifacts have been made available at https://github.com/nkkarpov/syncsignature.

1 INTRODUCTION

In this paper we consider the problem of tree similarity joins under edit distance (or, tree similarity joins for short), where given a collection of trees $\mathcal{T} = \{T_1, \ldots, T_N\}$ and a distance threshold K, we want to find all similar pairs of trees (T_i, T_j) such that $\text{TED}(T_i, T_j) \leq K$, where TED denotes the tree edit distance defined as the minimum number of node insertions, deletions, and substitutions to transfer one tree to the other. We consider rooted, ordered trees with labeled nodes. A node substitution is simply a node relabeling. When we delete a node v, we relink all the children of v to v's parent. When we insert a node v as a child of a node u, we relink a contiguous segment of u's children under v.

Tree-structured data is essential in modern data representation and analysis, and is widely used in data management (e.g., XML/JSON), code analysis and natural language processing (e.g., abstract syntax trees), bioinformatics (e.g., RNA secondary structures), and many other areas. An important primitive in these settings is to identify similar items under certain distance measurement, which is critical to various data analytics such as clustering, classification, and nearest neighbor queries. The tree edit distance is the most widely used distance function for measuring the similarity between a pair of trees.

Tree similarity joins has already attracted much attention in the past two decades [1, 2, 4, 5, 13, 14, 16, 30, 37]. Most of these works adopt the *filter-then-verify* framework: We first extract a set of candidate pairs trees such that this candidate set is a superset of the final output, and then verify for each pair of trees in the candidate set whether their tree edit distance is no more than the distance threshold by an exact tree edit distance computation. Different methods vary in their ways of performing the filtering and verification steps. A popular direction is to transform trees to simpler objects such as strings [1, 2, 13] and sets [4, 5, 16, 37], and then compute some distance (e.g., Euclidean distance, L1 distance, string edit distance, etc.) on the resulting pair of strings/sets which serves as a lower bound of the tree edit distance of the original pair of trees. This lower bound can be computed much more efficiently than tree edit distance, and can be used in a pruning step to quickly filter out tree pairs that should not be in the final output. The performance of these algorithms depends on the effectiveness of the pruning steps and the time for computing the lower bound. The difficulty with this approach is that obtaining a tight lower bound via the corresponding strings or sets is difficult because much of the tree structure is lost in the transformation.

Tang et al. [30] took a partition-based approach. The idea is to build an index containing a collection of subtrees. At each step when we process a new tree T, we search in the index for the (partial) subtree rooted at *each* node of T to find other trees that share the same subtree. After that, we partition T to subtrees of balanced sizes and add all the partitions to the index. The correctness of the algorithm is guaranteed by the pigeonhole principle, that is, for any pair of trees whose tree edit distance is no more than the threshold, they share at least one common subtree, which can be guaranteed if each tree is partitioned into sufficiently many subtrees. Although various optimizations are provided to speed up the search for common subtrees, the main algorithm in [30] is an index nested loop join and the number of subtree comparisons can be massive when the number and/or size of trees is large.

The state-of-the-art algorithm by Hütter et al. [14] also uses an index nested loop join approach. In their algorithm, more structures of the trees are taken into consideration when dynamically building the index. In addition, an efficient verification scheme is presented to speed up the overall join operation.

A critical issue with the index nested loop joins is that it does not support parallel execution. Recall that in index nested loop join algorithms, we process input trees one at a time, use the index to

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Proceedings of the VLDB Endowment, Vol. 16, No. 2 ISSN 2150-8097. doi:10.14778/3565816.3565833

look for similar trees in the set of processed trees, and then update the index with the newly processed input tree. As a result, algorithms proposed [14, 30] are not fully scalable in multicore architecture or massive parallel computation environments. In this paper, we propose a simple, efficient, yet fully parallelizable algorithmic framework for tree similarity joins that scales much better in parallel computation environments.

Our Approach and Contributions. We propose an algorithmic framework named SyncSignature for tree similarity joins (Section 2). The high-level idea of our approach is to generate for each tree a set of signatures and then perform a hash join using these signatures. A crucial property of our signature-based join is that the set of signatures for each tree is generated individually, which makes the overall framework fully parallelizable. Different from the previous partition-based approach [30], our signature generation process is "implicitly synchronized" between different trees. Intuitively speaking, the process guarantees that if two trees share a large enough common subtree, then with a good probability there will be common signatures generated from this common subtree on both trees, even if the signature generation process on each tree is performed individually without any coordination. This implicit synchronization feature facilitates an effective candidate-producing procedure via hash ioin.

Under the above framework, we propose two signature generation schemes that fit tree similarity joins. Both schemes are randomized since it may *not* be possible for deterministic partition schemes to achieve effective implicit synchronization. Consequently, our algorithms may miss a small number of valid tree pairs, but with the proper choices of parameters, they can reach near-perfect accuracy in theory *and* in most datasets we studied. Through an extensive set of experiments, we show that our proposed algorithms not only exceed the state-of-the-art algorithm in the centralized model on big datasets, but that they can also be fully parallelized, giving them a significant advantage in parallel computation environments.

Other Related Work. We briefly summarize other related works.

String similarity joins. Similarity joins under edit distance has also been studied extensively on strings [3, 7, 9, 12, 15, 19, 21, 26, 34–36, 38, 39]. Among these works, the one that is most relevant to ours is [39]. To some extent, our approach can be thought of as a generalization and extension of the MinJoin algorithm used for string similarity joins in [39], but there are multiple challenges when applying the idea in MinJoin to trees. We will discuss these challenges in Section 3.1.

The computation of tree edit distance in RAM. Tree edit distance can be computed in polynomial time in the RAM model. Tai [29] proposed the first polynomial time algorithm with a running time $O(n^6)$ where *n* is the size of (the larger of) the two trees. Zhang and Shasha [40] improved Tai's algorithm to $O(n^4)$ by observing that not all possible subproblems in the dynamic programming need to be solved. Klein [18] further improved the result to $O(n^3 \log n)$ using *heavy path decomposition*. This line of research culminated in the algorithm by Demaine et al. [10] with a running time $O(n^3)$. Pawlik and Augsten also proposed an $O(n^3)$ algorithm named RTED [23], which demonstrates better practical performance. Later, the same authors proposed APTED [24] with the same running time but a

Table 1: Summary of Notations

Notation	Definition			
[<i>n</i>]	$[n] \triangleq \{1, 2, \ldots, n\}$			
K	edit distance threshold			
\mathcal{T}	set of input trees			
T _i	<i>i</i> -th tree in \mathcal{T}			
N	number of input trees, i.e., $N = \mathcal{T} $			
Σ	alphabet of labels of nodes in ${\mathcal T}$			
П	hash function $\Sigma^q \rightarrow (0, 1)$ for generating ranks			
Γ	hash function $\Sigma^* \to \mathbb{N}$ for computing fingerprints			
z	neighborhood size parameter			
τ	signature similarity parameter			
$N_r(v)$	neighborhood of v of radius r			

better space usage. For the threshold version of the tree edit distance problem where we only need to decide whether the distance is at most *K* or larger than *K*, Touzet [31] proposed an algorithm with running time $O(nK^3)$. Recently, Pawlik and Augsten [25] designed an algorithm for threshold tree edit distance with improved practical performance. All these algorithms, however, cannot be applied directly to tree similarity joins without an effective candidate generation step.

Similarity joins in the massive parallel computation model. We notice that parallel/distributed computation has already been studied for set and string similarity joins [6, 8, 27, 28, 32]. However, to the best of our knowledge, no fully parallelizable algorithm has been designed for tree similarity joins.

Preliminaries. We consider rooted, ordered, unweighted, and undirected trees with labeled nodes. For two nodes u, v in a tree T, the distance of u and v is defined to be the length of the unique path between u and v in T. For a node v in T, let $N_r(v)$ be the set of nodes in T that are within a distance of r from v, including v itself. We call $N_r(v)$ the neighborhood of v of radius r, or, the r-neighborhood of v.

We have listed a set of frequently used notations in this paper in Table 1.

2 THE JOIN FRAMEWORK

We begin by presenting our algorithmic framework SYNCSIGNATURE for tree similarity joins. The framework is described in Algorithm 1.

Let us briefly describe Algorithm 1 in words. Besides the input set of trees \mathcal{T} and the distance threshold *K*, Algorithm 1 takes two more parameters *z* and τ . The former determines the neighborhood size which will be used in implicitly synchronized signature generation schemes, and the later is a parameter for comparing the similarity of two set of signatures.

Our framework consists of three components.

- Signature generation. At Line 3-8, for each input tree T_i, we generate a set of signatures S_i. For each s = (s.key, s.pos) ∈ S_i, where s.key is the fingerprint of the signature and s.pos is the position of the signature which will be specified in concrete signature generation schemes, we add (i, s.pos) to the bucket in hash table D indexed by s.key.
- (2) Join. Line 9-15 is the hash join. To reduce the chance of mismatch, we include a sanity check of two signatures with the same key; the check rejects all pairs of signatures

Algorithm 1 SyncSignature(\mathcal{T}, K, z, τ)

Input: Collection of trees $\mathcal{T} = \{T_1, \ldots, T_n\}$, distance threshold *K*, neighborhood size parameter *z*, signature similarity parameter τ .

Output: $O = \{(T_i, T_j) \mid T_i, T_j \in \mathcal{T} : i \neq j, \text{TED}(T_i, T_j) \leq K\}$

1: $O \leftarrow \emptyset, I \leftarrow \emptyset$

 Initialize an empty hash table D and an empty table of counters C; generate a random hash function Π : Σ → (0, 1) where Σ is the node label universe, and a random hash function Γ : Σ* → N.

3: for each $T_i \in \mathcal{T}$ do

4: $S_i \leftarrow \text{Signature}(T_i, \Pi, z, \Gamma, \tau)$

5: for each $s \in S_i$ do

6: Add (*i*, *s.pos*) into the bucket in *D* indexed by *s.key*7: end for

7: end

8: end for

9: for each bucket of \mathcal{D} do

```
for each pair (i, pos), (j, pos') in the bucket do
10:
             if (||T_i| - |T_j|| \le K) \land (|pos - pos'| \le K) then
11:
                 Increment counter C(i, j)
12:
13:
             end if
14:
        end for
15: end for
16: for each (i, j) with C(i, j) > 0 do
17:
        if C(i, j) \ge \tau then
             I \leftarrow I \cup \{(i, j)\}
18:
        end if
19:
20: end for
21: for each (i, j) \in I do
        if LOWERBOUNDED(T_i, T_j) \leq K then
22:
             if UpperBoundED(T_i, T_j) \leq K then
23:
24:
                 O \leftarrow O \cup \{(i, j)\}
25:
             else if TED(T_i, T_i) \leq K then
26:
                 O \leftarrow O \cup \{(i, j)\}
             end if
27:
28:
        end if
29: end for
```

whose associated tree sizes differ by more than *K* or their positions in the associated trees differ by more than *K*. At Line 16-20, we collect candidate pairs of trees whose common signatures are above a predetermined threshold τ .

(3) Verification. At Line 21-29, we perform a verification step on each pair of candidates to remove false positives. Before the exact tree edit distance computation, we apply some upper and lower bounds of the tree edit distance to filter out or early accept each candidate pair.

For LOWERBOUNDED() at Line 22, one choice, as we select for our ball signature generation scheme (see Section 3.2), is to use the string edit distance of the preorder traversals of the two trees T_i, T_j , which is known to be at most K if $\text{TED}(T_i, T_j) \leq K$ (see, e.g., [13]). For the Euler-tour embedding based signature generation scheme (see Section 3.3), we will use the *half* of the string edit distance of the Euler-tour embedding of T_i, T_j , which can also be shown to be at most K if $\text{TED}(T_i, T_j) \leq K$. For UPPERBOUNDED() at Line 23, we use the LGM upper bound introduced in [14]. Both check procedures can finish in $O(\max\{|T_i|, |T_j|\}K)$ time.

We will propose a couple of signature generation schemes for SIGNATURE() at Line 4 of Algorithm 1 in Section 3.

Grouping. In the SYNCSIGNATURE framework, we use the same neighborhood size *z* for all trees in the dataset. This is *not* desirable

Algorithm 2 Tree-Similarity-Joins(\mathcal{T}, K, c)

Input: Collection of trees \mathcal{T} , distance threshold K, parameter $c \in (0, 1)$, signature similarity parameter τ

Output: $O = \{(T_i, T_j) \mid T_i, T_j \in \mathcal{T} : i \neq j, \mathsf{TED}(T_i, T_j) \leq K\}$

1: Let *max* be the smallest integer such that the size of the largest tree in \mathcal{T} is no more than (max + 1)K/c + K.

2: **for** z = 0, 1, ..., max **do**

3: $\mathcal{T}_z = \{T_i \in \mathcal{T} \mid |T_i| \in [zK/c, (z+1)K/c+K]\}$

4: SYNCSIGNATURE(*T_z*, *K*, *z*, *K*/5)
 5: end for

if the sizes of the trees in the dataset vary significantly. One idea to resolve this issue is to first group input trees according to their sizes before feeding them into Algorithm 1.

More precisely, for each z = 1, 2, ..., we create a group of trees

$$\mathcal{T}_{z} = \{T_{i} \in \mathcal{T} \mid |T_{i}| \in [zK/c, (z+1)K/c+K]\},\$$

where $0 < c \le 1$ is a constant parameter. To avoid missing any valid output pairs, every two adjacent groups overlap by an interval of length *K*. After the grouping, we call Algorithm 1 on each \mathcal{T}_z , and union the outputs of all groups at the end. The final algorithm is presented in Algorithm 2.

Note that in Algorithm 2, there is only one parameter *c*, as *z* runs over 1, 2, . . . When calling SyncSignature, we set $\tau = K/5$. We note in advance that $\tau = K/5$ satisfies the requirement $\tau = O(K)$ in Theorem 3.2 and Theorem 3.4 for our signature generation schemes. Both *z* and *c* control the neighbor size. We have $z = c\eta/K$, where η is the smallest tree size in the *z*-th group. We call *c* the (*neighborhood*) *resolution*, and will discuss how to choose a good value for *c* in Section 4.

In the rest of this paper, we always assume that the size of the input trees are all within a constant factor, since we perform the grouping operation before calling SyncSignature.

2.1 Parallel Implementation

We now describe the parallel implementation of SyncSignature framework (Algorithm 1). Let M be the number of cores/machines. First, the signature generation phase (Line 3-8) can be fully parallelized, since signature generation for each tree is performed independently. We only need to evenly distribute the set of trees to the M machines. To insert the signatures into the hash table, the M machines can first generate a hash table locally, and then merge the M hash tables.

For the join phase (Line 9-15), we try to distribute the workload to the *M* machines in the following way. Suppose there are *t* nonempty buckets in the hash table \mathcal{D} , let a_1, a_2, \ldots, a_t be the number of signatures that are hashed into the *t* buckets, and let $b_i = \binom{a_i}{2}$ be the number of signature pairs in the *i*-th bucket that we need to consider. Let $b = \sum_{i \in [t]} b_i$. We try to partition the *t* buckets to the *M* machines so that each machine handles a similar number of signature pairs. More precisely, let i_1 be the smallest index such that $b_1 + \ldots + b_{i_1} \ge b/M$; we assign the first i_1 buckets to the first machine. Let i_2 be the smallest index such that $b_{i_1+1} + \ldots + b_{i_2} \ge b/M$; we assign the $(i_1 + 1)$ -th to the i_2 -th buckets to the second machine. And so on. At the end, the *M* machines merge local counters C(i, j).

The amount of work at Line 16-20 is insignificant; we can just execute these lines sequentially using one machine.

The verification phase (Line 21-29) can be fully parallelized. We evenly distribute the set of candidate pairs in I to the M machines, and union all *M* output sets at the end. We note that the verification step in most algorithms following the filter-then-verify framework can be parallelized in the same way.

SIGNATURE GENERATION SCHEMES 3

In this section, we introduce several signature generation schemes for the SyncSignature framework. We first present a direct extension of the local hash minima partition scheme (LHM-partition) for trees; LHM-partition was originally designed for string similarity joins [39]. However, the direct extension cannot perform well in practice due to the unbalanced signature sizes. We then propose two effective signature generation schemes whose practical performance are not entirely comparable.

3.1 Local Hash Minima Partition

The direct extension of LHM-partition to tree similarity joins is as follows: For each input tree *T*, we create a random rank for each node of the tree. We next define for each node *x* its neighborhood to be all nodes in the ball of radius *r* centered at *x*. We designate a node *x* as *anchor* if *x* has the smallest rank among all nodes in the neighborhood. We then partition *T* at all of its anchor nodes, transforming the tree into a collection of subtrees. After the LHMpartition, we select all pairs of trees that share at least one common subtree as candidates of the join output, and conduct a verification step for all candidate pairs at the end.

Note that the partition on each tree is performed individually without any coordination. Intuitively, the anchors (i.e., local hash minima) provide an implicit coordination among all trees so that if two trees share a large common subtree, then with high probability this subtree will be cut at exactly the same set of anchors on both trees, generating a common partition. On the other hand, by the pigeonhole principle we know that if two trees have a small edit distance, then they must share a large common subtree, which gives the correctness of the algorithm.

However, there are several challenges of this direct extension.

- (1) Different from the string case, the neighborhood sizes of different nodes in trees can be very different - for a node with degree $\Theta(n)$, the neighborhood size can be $\Theta(n)$ even if the radius r = 1! The analysis of LHM-partition naturally requires that the probabilities for nodes being anchors are similar, which necessitates that all nodes' neighborhood sizes be close.
- (2) If we force the sizes of different nodes' neighborhoods to be similar, symmetry may break: it is possible for v to be w's neighbor while w is not v's neighbor.
- (3) A single edit operation may change a large number of partitions, and consequently make the pigeonhole principle inapplicable. While in the string case, one edit can only affect at most two anchors (thus at most three substrings).

To tackle the first problem, we transform each tree to a left-child right-sibling binary tree with the degree of each node bounded by 3. More precisely, for each node v with m ordered children v_1, \ldots, v_m , we replace edges $(v, v_1), (v, v_2), \ldots, (v, v_m)$ with edges $(v, v_1), (v_1, v_2), \ldots, (v_{m-1}, v_m).$

Algorithm 3 PARTITION-SIGNATURE $(T, \Pi, z, \Gamma, \cdot)$

Input: Input tree *T*, random hash function $\Pi : \Sigma \to (0, 1)$, neighborhood size parameter *z*, random hash function $\Gamma : \Sigma^* \to \mathbb{N}$

▷ l_v is the label of node v

Output: A set of signatures *S* of *T*

- 1: Initialize $S \leftarrow \emptyset, A \leftarrow \emptyset$
- 2: for each $v \in T$ do
- Assign rank $r_v \leftarrow \Pi(\ell_v)$ 3:
- 4: end for
- 5: Convert T into a left-child right-sibling binary tree T'
- for each $v \in T'$ do 6:
- 7. Choose the minimal radius r of neighborhood v in T' such that $|N_r(v)| \in [z, 2z)$
- 8: if $r_v = \min_{x \in N_r(v)} r_x$ then
- 9: $A \leftarrow A \cup \{v\}$
- end if 10:
- 11: end for
- 12: for each $v \in A$ do
- Cut the tree T' at node v; duplicate v with deg(v) copies, each 13: connecting to one of the adjacent nodes of v in T'
- 14: end for
- 15: Let C_1, \ldots, C_M be the set of connected components after cutting T' at nodes in A
- 16: **for** i = 1, ..., M **do**
- Let $(u_1, \ldots, u_{|C_i|})$ be the preorder traversal of C_i 17:
- $s_i.key \leftarrow \Gamma(\ell_{u_1} \cdots \ell_{u_{|C_i|}})$ 18:
- $s_i.pos \leftarrow \min_{u \in C_i} \{ \text{index of node } u \text{ in the preorder traversal of } T' \}$ 19: 20: end for
- 21: return $S = \{s_1, ..., s_M\}$

FACT 1 (SEE, E.G., [30]). If we apply the left-child right-sibling binary tree transformation on each input tree, the edit distance between any pair of trees after the transformation will be increased by at most a factor of 2.

To compensate for the factor 2 transformation distortion, we need to double the distance threshold *K* to retain the correctness of our algorithm. That is, we need to find pairs of binary trees (after the transformation) with edit distance at most 2K for verification. However, such a change will not affect the correctness of our algorithms. We will give a detailed explanation in Section 3.4.

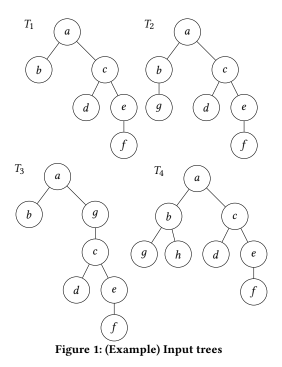
The problem of asymmetry is built into the tree structure. Fortunately, we can show that asymmetry has little effect on the algorithm's soundness if the neighborhood sizes of all nodes are close, via a careful analysis which is significantly more complicated than that for the original LHM-partition on strings (see Lemma 3.1).

The third problem can also be resolved via a more careful analysis once the node degrees of the trees are bounded by 3.

Our partition-based signature generation scheme is described in Algorithm 3. At Line 2-4, we assign each node a random rank. We then convert the tree to a left-child right-sibling binary tree (Line 5), and identify all anchors based on the random ranks (Line 6-11). Finally, we partition the tree at all anchors and create a signature for each resulting connected components/subtrees (Line 12-16).

The signature of each connected component C consists of a key and a position. The key of C is simply a hash value of the preorder traversal of nodes of C, and the position of C is the minimum index of nodes in *C* in the preorder traversal of the input tree.

The following lemma gives a good estimate of the number of anchors that Algorithm 3 produces, and will be used in the analysis in Section 3.2. The proof of this lemma is quite technical and can



be skipped during the first reading. Due to space constraints, we leave it to Appendix A.1 of the full version of this paper [17]. In the rest of this paper, when we say appendix we always mean the appendix of the full version.

LEMMA 3.1. Let $n \triangleq |T|$ be the size of the tree. If $z = o(n^{1/3})$, then the number of anchors generated by Algorithm 3 falls in the range (n/(3z), 2n/z) with probability 1 - o(1).

An Example. Let us use an example to illustrate how PARTITION-SIGNATURE works.

Figure 1 describes the set of four input trees. We set the distance threshold K = 1, two parameters z = 2 and $\tau = 1$, and assume that the random hash function Π gives the following order of labels on the trees:

 $\Pi(b) < \Pi(d) < \Pi(g) < \Pi(a) < \Pi(c) < \Pi(e) < \Pi(f) < \Pi(h).$

Given the random ordering Π , the set of anchors in T_1 is $\{b, d\}$. Since z = 2, the neighborhood associated with each node is simply the node itself and all of its adjacent nodes. Cutting T_1 at $\{b, d\}$ and duplicating the anchor nodes, we get three signatures $\{(b, 2), (abcdef, 1), (d, 4)\}$, where for the illustration purpose the fingerprint of each ball is simply represented as a concatenation of the labels of nodes in the ball according to the preorder traversal without applying the hash function Γ . We perform the same operation on other trees; Table 2 summarizes the tree signatures. Table 3 shows the hash table after inserting all signatures.

A Major Issue with PARTITION-SIGNATURE when working with the SyncSignature framework. The main issue with Algorithm 3 is that it may produce subtrees with very different sizes, and many of them are small. Particularly, since at least have of the nodes are leaves, many of them would become singletons after the partition under the random ranking scheme (see, e.g., nodes *b*, *d* in Table 2). This phenomenon is very different from the string/path

Table 2: Signatures produced by PARTITION-SIGNATURE

tree	ball signatures			
T_1	(b, 2), (abcdef, 1), (d, 4)			
T ₂	(bg, 2), (abcdef, 1), (d, 5)			
<i>T</i> ₃	(b, 2), (abg, 1), (gcdef, 3), (d, 5)			
T_4	(bgh, 2), (abcdef, 1), (d, 5)			

Table 3: Hash table for PARTITION-SIGNATURE

key	bucket contents			
b	$(T_1, 2), (T_3, 2)$			
d	$(T_1, 4), (T_2, 5), (T_3, 5), (T_4, 5)$			
abg	$(T_3, 1)$			
bgh	$(T_4, 2)$			
gcdef	$(T_3, 3)$			
abcdef	$(T_1, 1), (T_2, 1), (T_4, 1)$			

setting, since a path only has two leaves/end-nodes. Small partitions are more likely to get matched. Consequently, the join framework may produce many false positives and set a significant burden to verification step, making the algorithm impractical on large datasets. In our example, all 6 pairs of trees will be candidate pairs for verification.

In the next two subsections, we propose two variants of Algorithm 3 that can generate subtrees/signatures of similar sizes for a much better performance.

3.2 Signatures Using Neighborhoods

The first idea to resolve the issue mentioned above is to simply take the neighborhoods of all anchors as the subtree signatures. We call this scheme BALL-SIGNATURE, described in Algorithm 4. Compared with Algorithm 3, the main difference is that in Algorithm 4 we directly take the neighborhoods of anchors as signatures at Line 13. To further speed up the signature generation process, we first sort all nodes of the tree according to their ranks in the increasing order, and then try to find anchors starting from the node with the lowest rank. We end the process when 5τ anchors have been identified, or all anchors have been found. We note that 5τ is just used to facilitate the theoretical analysis; any larger constant than 5 will also work.

An Example. Let us again use the four input trees in Figure 1 to illustrate Algorithm 1 equipped with BALL-SIGNATURE. We again set K = 1, z = 2, and $\tau = 1$. Assume the random hash function Π gives the following order of labels on the trees:

$$\Pi(b) < \Pi(c) < \Pi(g) < \Pi(a) < \Pi(d) < \Pi(e) < \Pi(f) < \Pi(h).$$

Given this random ordering, the set of anchors in T_1 is $\{b, c\}$. The set of signatures of T_1 is $\{(ab, 1), (acde, 1)\}$, where for the illustration purpose the fingerprint of each ball is again represented as a concatenation of the labels of nodes in the ball according to the preorder traversal without applying the hash function Γ . We perform the same operations on other trees; Table 4 summarizes the tree signatures.

We next add all signatures to the hash table \mathcal{D} ; the results are described in Table 5.

Algorithm 1 checks at Line 11 the following pairs:

 $\{(T_1, 1), (T_3, 1)\}, \{(T_1, 1), (T_2, 1)\}, \{(T_1, 1), (T_4, 1)\}, \{(T_2, 1), (T_4, 1)\}.$

Input: Input tree *T*, random hash function $\Pi : \Sigma \to (0, 1)$, neighborhood size parameter *z*, random hash function $\Gamma : \Sigma^* \to \mathbb{N}$, signature similarity parameter τ

Output: A set of signatures *S* of *T*

1: Initialize $S \leftarrow \emptyset$

2: for each $v \in T$ do

- 3: Assign rank $r_v \leftarrow \Pi(\ell_v)$
- 4: end for
- 5: Sort nodes in T according to their rank in the increasing order, getting $\{v_1, \ldots, v_{|T|}\}$
- 6: Convert T into a left-child right-sibling binary tree T'
- 7: **for** $v = v_1, ..., v_{|T|}$ **do**

8: Choose the minimal radius r of neighborhood v in T' such that $|N_r(v)| \in [z, 2z)$

if $r_v = \min_{x \in N_r(v)} r_x$ then 9:

Let $(u_1, \ldots, u_{|N_r(v)|})$ be the preorder traversal of $N_r(v)$ 10:

11:

 $s.key \leftarrow \Gamma(\ell_{u_1} \cdots \ell_{u_{|N_r(v)|}})$ s.pos $\leftarrow \min_{u \in N_r(v)} \{\text{index of node } u \text{ in the preorder traversal of } T'\}$ 12:

 $S \leftarrow S \cup \{s\}$ 13:

- end if 14:
- if $|S| = 5\tau$ then 15:
- return S 16:
- end if 17:
- 18: end for
- 19: return S

Table 4: Sets of signatures produced by BALL-SIGNATURE

tree	ball signatures
T_1	(<i>ab</i> , 1), (<i>acde</i> , 1)
T_2	(<i>abg</i> , 1), (<i>acde</i> , 1)
T_3	(ab, 1), (gcde, 3)
T_4	(<i>abgh</i> , 1), (<i>acde</i> , 1)

Table 5: Hash table for BALL-SIGNATURE

key	bucket contents			
ab	$(T_1, 1), (T_3, 1)$			
acde	$(T_1, 1), (T_2, 1), (T_4, 1)$			
abg	$(T_2, 1)$			
gcde	$(T_3, 3)$			
abgh	$(T_4, 1)$			

All pairs except $\{(T_1, 1), (T_4, 1)\}$ pass the check. The final set of candidates for verification is $\{(T_1, T_3), (T_1, T_2), (T_2, T_4)\}$. The three pairs will pass the verification step and appear in the final output because each has an edit distance of 1.

Analysis of Algorithm 4. In BALL-SIGNATURE, the neighborhoods of anchors may overlap. Compared with disjoint signatures generated from a partition scheme, overlapping signatures introduces a new challenge: a single edit may affect many signatures. For example, consider a star tree T with one central node and 100 leaf nodes. Let K = 1, z = 2, and $\tau = 1$. Under a random hash ranking, about half of the leaf nodes will be anchors, whose neighborhoods will contain the central node and itself. Consequently, a single edit on the central node (getting another tree T') will affect about half of the signatures. This could cause the similarity check at Line 17 of Algorithm 1 to fail even that $\text{TED}(T, T') \leq 1$.

It appears to be impossible to prove that SyncSignature equipped with BALL-SIGNATURE will not miss any valid pairs under an adversarial set of K edits. However, if the K edits occur at random locations, we can still show that our join algorithm succeeds with a good probability.

THEOREM 3.2. For binary trees T_1, T_2 with $|T_1| = n$ and $\text{TED}(T_1, T_2) \leq$ K, let S_1 and S_2 be the sets of signatures outputted by Algorithm 4 on T_1 and T_2 respectively. For any $z = o(\min\{n/K, n^{1/3}\})$ and $\tau = O(K)$, if the (at most) K edits take place at random nodes, then the probability that S_1 and S_2 shares at least τ common signatures is at least 1 - o(1). The running time of Algorithm 4 is $O(nz \log n)$.

Proof: By Lemma 3.1, we know that with probability 1 - o(1), the number of anchors generated by Algorithm 4 is at least n/(3z), which is at least 5τ for $\tau = O(K)$ given z = o(n/K). Therefore, with probability 1 - o(1), Algorithm 4 returns S containing exactly 5τ signatures (early termination at Line 15-16).

We next consider the number of signatures that will be affected by K edits. Observe that on a binary tree, one edit can affect at most O(1) nodes. For each signature s_i (corresponding to a subtree of size at most 2z), let Y_i be the indicator variable for the event that at least one of the K edits affects s_i . If the K edits are randomly selected, then we have

$$\Pr[Y_i = 1] \le K \cdot O(1) \cdot 2z/n = O(Kz/n).$$

Let $Y = \sum_{i=1}^{5\tau} Y_i$. Given z = o(n/K), we have

$$\mathbf{E}[Y] \le 5 \cdot O(\tau K z/n) = o(\tau).$$

By Markov's inequality we have $\Pr[Y \le \tau] \ge 1 - o(1)$, and consequently $\Pr[5\tau - Y \ge \tau] \ge 1 - o(1)$. That means, for a pair of trees T_1, T_2 with $\mathsf{TED}(T_1, T_2) \leq K$, they will share at least τ common signatures with probability 1 - o(1).

For the running time, for each node we spent O(z) time to construct a ball neighborhood and $O(z \log n)$ for computing the signature. The time cost of other steps is of lower-order term. Thus the total running time is $O(zn \log n)$.

Signatures Using Euler-Tour Embedding 3.3

The second idea to resolve the issue of unbalanced signature sizes is to first embed trees to strings, use LHM-partition on strings, and then map substrings back to subtrees to produce signatures. The signature generation scheme based on Euler-tour embedding is described in Algorithm 5. At Line 1 we convert the tree to a string which is the Euler-tour of the tree. Next, we use LHM-partition to partition the string to substrings (Line 2-8). Finally, for each substring, we retrieve the corresponding subtree (to be explained in the next paragraph) and generate a signature of the subtree.

The Euler-tour translates a tree *T* of size *n* to a string of length 2n. Each element in the string consists of the label of the corresponding tree node and a sign. Each node of the tree T appears twice in the string with different signs ('+' and '-'). The details are described in Algorithm 6. Each substring in the Euler-tour can be inversely mapped to a subtree in T as follows: we first identify the corresponding node in T for each letter in the substring, getting a set F. We then remove duplicated nodes in F (recall that each node in *T* corresponds to 2 letters in the Euler-tour). The set of nodes in F must be connected due to the construction of the Euler-tour. The

Algorithm 5 EULER-SIGNATURE $(T, \Pi, z, \Gamma, \cdot)$

Input: Input tree *T*, distance threshold *K*, random hash function $\Pi : \Sigma \rightarrow (0, 1)$, neighborhood size parameter *z*, random hash function $\Gamma : \Sigma^* \rightarrow \mathbb{N}$

Output: A set of signatures *S* of *T*

- 1: $E \leftarrow \text{Euler-Tour}(\text{root of } T, T)$
- 2: For *i*-th element y_i in *E*, assign a rank $r_i \leftarrow \Pi(y_i)$

3: Let I be the subset of $\{z + 1, \ldots, |E| - z\}$ such that $r_i =$ $\min_{j \in \{i-z,\dots,i+z\}} \{r_j\}; \text{ denote } I = (i_1,\dots,i_k) \text{ with } i_1 < \dots < i_k$ 4: Initialize $P \leftarrow \{y_1 \dots y_{i_1} - 1\}$ 5: **for** j = 1, ..., k - 1 **do** $P \leftarrow P \cup \{y_{i_j} \ldots y_{i_{j+1}} - 1\}$ 6: 7: end for 8: $P \leftarrow P \cup \{y_{i_k} \dots y_{|E|}\}$ 9: for each $p \in P$ do Let Q be the corresponding subtree of p in T10: Let $(u_1, \ldots, u_{|Q|})$ be the preorder traversal of Q11: $s.key \leftarrow \Gamma(\ell_{u_1} \cdots \ell_{u_{|Q|}})$ 12: $s.pos \leftarrow \lfloor (\text{index of the first element of } p \text{ in } E)/2 \rfloor$ 13: $S \leftarrow S \cup \{s\}$ 14: 15: end for

16: return S

Algorithm 6 EULER-TOUR(*v*, *T*)

Input: Input tree *T* and a vertex *v* in *T* **Output:** Euler-tour of the subtree rooted at *v* 1: **if** *v* is leaf **then** 2: **return** $\ell_v^+ \ell_v^-$ 3: **else** 4: Let v_1, \ldots, v_k be the children of *v* 5: **return** $\ell_v^+ \circ \text{EULER-TOUR}(v_1, T) \circ \ldots \circ \text{EULER-TOUR}(v_k, T) \circ \ell_v^-$ 6: **end if**

Table 6: Euler-tours of T_1 , T_2 , T_3 , and T_4 .

tree	Euler-tours
T_1	$a^{+}b^{+}b^{-}c^{+}d^{+}d^{-}e^{+}f^{+}f^{-}e^{-}c^{-}a^{-}$
T_2	$a^{+}b^{+}g^{+}g^{-}b^{-}c^{+}d^{+}d^{-}e^{+}f^{+}f^{-}e^{-}c^{-}a^{-}$
<i>T</i> ₃	$a^{+}b^{+}b^{-}g^{+}c^{+}d^{+}d^{-}e^{+}f^{+}f^{-}e^{-}c^{-}g^{-}a^{-}$
T_4	$a^{+}b^{+}g^{+}g^{-}h^{+}h^{-}b^{-}c^{+}d^{+}d^{-}e^{+}f^{+}f^{-}e^{-}c^{-}a^{-}$

subtree of the inverse mapping is simply the induced subtree in T of the nodes in F.

An Example. We will again use the input trees in Figure 1 as an example to illustrate Algorithm 1 equipped with EULER-SIGNATURE. We again set the distance threshold K = 1, and two parameters z = 2, $\tau = 1$. We assume that the random hash function Π gives the following order of labels:

$$\begin{split} \Pi(h^{-}) &< \Pi(g^{-}) < \Pi(b^{-}) < \Pi(f^{+}) < \Pi(f^{-}) < \Pi(a^{+}) \\ &< \Pi(a^{-}) < \Pi(b^{+}) < \Pi(c^{+}) < \Pi(c^{-}) < \Pi(d^{+}) < \Pi(d^{-}) \\ &< \Pi(e^{+}) < \Pi(e^{-}) < \Pi(g^{+}) < \Pi(h^{+}). \end{split}$$

The algorithm first generates the Euler-tours of the four trees, presented in Table 6. We then partition each Euler-tour string and generate signatures for each tree; see Table 7. The corresponding sets of signatures of the partitions are presented in Table 8; for simplicity, instead of mapping back to the subtrees and using the

Table 7: Partitions of Euler-tours.

tree	partitions of Euler-tours
<i>T</i> ₁	$a^{+}b^{+} \mid b^{-}c^{+}d^{+}d^{-}e^{+} \mid f^{+}f^{-}e^{-}c^{-}a^{-}$
<i>T</i> ₂	$a^{+}b^{+}g^{+} \mid g^{-}b^{-}c^{+}d^{+}d^{-}e^{+} \mid f^{+}f^{-}e^{-}c^{-}a^{-}$
T ₃	$a^{+}b^{+} \mid b^{-}g^{+}c^{+}d^{+}d^{-}e^{+} \mid f^{+}f^{-}e^{-}c^{-}g^{-}a^{-}$
T_4	$a^{+}b^{+}g^{+}g^{-}h^{+} \mid h^{-}b^{-}c^{+}d^{+}d^{-}e^{+} \mid f^{+}f^{-}e^{-}c^{-}a^{-}$

Table 8: Set of signatures produced by EULER-SIGNATURE.

tree	Euler signatures
T_1	(ab, 0), (bcde, 1), (acef, 4)
T_2	(abg, 0), (bgcde, 2), (acef, 5)
T_3	(ab, 0), (bgcde, 1), (agcef, 4)
T_4	(abgh, 0), (bhcde, 3), (acef, 6)

Table 9: Hash table for EULER-SIGNATURE.

key	bucket contents			
ab	$(T_1, 0), (T_3, 0)$			
bcde	$(T_1, 1)$			
acef	$(T_1, 4), (T_2, 5), (T_4, 6)$			
abg	$(T_2, 0)$			
bgcde	$(T_2, 2), (T_3, 1)$			
agcef	$(T_3, 4)$			
abgh	$(T_4, 0)$			
bhcde	(<i>T</i> ₄ , 3)			

random hash function Γ, we just utilize the substrings as signature keys. The corresponding hash table is presented in Table 9. Next, Algorithm 1 checks at Line 11 the following five pairs:

 $\{(T_1, 0), (T_3, 0)\}, \{(T_1, 4), (T_2, 5)\}, \{(T_1, 4), (T_4, 6)\},\$

 $\{(T_2, 5), (T_4, 6)\}, \{(T_2, 2), (T_3, 1)\}.$

All pairs except { $(T_1, 4), (T_4, 6)$ } pass the check. The verification step will further reject the candidate pair (T_2, T_3) whose edit distance is greater than K = 1. The final output is again { $(T_1, T_3), (T_1, T_2), (T_2, T_4)$ }. **Analysis of Algorithm 5.** The following lemma states that Eulertour embedding will not increase the edit distance by more than a factor of 2. Due to space constraints, the proof can be found in Appendix A.2.

LEMMA 3.3. Let E_1 and E_2 be the Euler-tour embedding of trees T_1 and T_2 , respectively. We have $ED(E_1, E_2) \leq 2 \cdot TED(T_1, T_2)$.

Now we present the main theorem for Algorithm 5.

THEOREM 3.4. For binary trees T_1, T_2 with $|T_1| = n$ and $\text{TED}(T_1, T_2) \leq K$, let S_1 and S_2 be the sets of signatures outputted by Algorithm 5 on T_1 and T_2 respectively. For any z = o(n/K) and $\tau = O(K)$, the probability that S_1 and S_2 shares at least τ common signatures is at least 1 - o(1). The running time of Algorithm 5 is $O(n \log n)$.

Proof: We make use of the following lemma which states that the number of partitions created by LHM-partition is tightly concentrated to its mean.

LEMMA 3.5 ([39]). Given an input Euler-tour E and a neighborhood size parameter z, for any c > 0, the size of set I(|I| = X) created by Algorithm 5 satisfies $\Pr\left[|X - \mathbb{E}[X]| > \sqrt{c\mathbb{E}[X]}\right] < 1/c$, where $\mathbb{E}[X] = \frac{|E| - 2z}{2z+1}$.

By Lemma 3.3, we know that after the embedding, the two resulting strings E_1 and E_2 has (string) edit distance at most 2K.

We now consider a particular edit operation O on E_1 . It is easy to see that the number of anchors that O can affect is bounded by 2. Consequently, the number of partitions O can affect is bounded by 3. Given that there are at most 2K edit operations, the total number of partitions of E_1 that may be affected by edit operations is bounded by 6K. By Lemma 3.5, if z = o(n/K), setting $c = n^{0.1} = \omega(1)$ and recalling that $|E_1| = 2n$ and $\tau = O(K)$, we have with probability 1 - o(1), there are at least

$$(1 - o(1)) \cdot \frac{|E_1| - 2z}{2z + 1} - 6K \ge 0.9 \cdot \frac{|E_1|}{2z} - 6K \ge 1$$

unaffected partitions after the 2K edits.

For the running time, the Euler tour can be constructed in O(n) time. The set of partitions *P* can be constructed in O(n) time by a linear scan. Same as before, the fingerprint computation costs $O(n \log n)$ time. The time cost of other steps is of lower-order term. Thus the total running time is $O(n \log n)$.

3.4 Plugging Signature Generation Schemes to the SyncSignatureFramework

Our signature generation schemes BALL-SIGNATURE and EULER-SIGNATURE are both Monte Carlo randomized algorithms, which has a small error probability. More precisely, Theorem 3.2 (for BALL-SIGNATURE) and Theorem 3.4 (for EULER-SIGNATURE) state that for each pair of binary trees whose edit distance is at most K, the pair will be include in the final output with probability at least $1 - \delta$ for a $\delta = o(1)$. The standard *parallel repetition* method can further reduce the error δ all the way to zero. More precisely, we repeat the signature generation step and the hash join step for log N times (N is the number of input trees), each time using independent random ranks. We then union the $\log N$ sets of candidate pairs for verification. Now, each pair of trees with edit distance at most K will be included in the output with probability at least $1 - \delta^{\log N} \geq$ $1 - 1/N^{100}$. By a union bound on $O(N^2)$ pairs of trees, all pairs of trees with edit distance at most K will be included in the output with probability at least 1 - 1/N. On the other hand, the verification phase in the SyncSignature framework guarantees that there is no false positive in the output.

By Fact 1, the binary tree transformation may increase the edit distance between a pair of trees by at most a factor of 2, to compensate which we need to find pairs of binary trees (after the transformation) with edit distance at most 2K for verification. However, this change will *not* affect the correctness of Theorem 3.2 (for BALL-SIGNATURE) and Theorem 3.4 (for EULER-SIGNATURE) – the statements and the proofs of the theorems still hold if we replace K with 2K everywhere. We note that the final verification step is performed on the *original* input trees with distance threshold K.

We have the following theorems for SyncSignature (Algorithm 1) equipped with our signature generation schemes. Recall that after grouping, we can assume that the sizes of all input trees are within a constant factor. Let n is the size of the smallest tree in the input.

THEOREM 3.6. For any $z = o(\min\{n/K, n^{1/3}\})$ and $\tau = O(K)$, equipped with BALL-SIGNATURE (with log N parallel repetition), Algorithm 1 solves the tree similarity joins exactly with probability at least 1 - 1/N.

THEOREM 3.7. For any z = o(n/K) and $\tau = O(K)$, equipped with EULER-SIGNATURE (with log N parallel repetition), Algorithm 1 solves the tree similarity joins exactly with probability at least 1 - 1/N.

Performing parallel repetition will naturally increase the running time. In our experiments, we do *not* use parallel repetition, thus our algorithms may have false negatives (in theory, δ fraction of false negatives in expectation). Our experiments in Section 4 demonstrate that on the datasets we have tested, our algorithms achieve almost perfect accuracy in most cases without parallel repetition. In the worst case, our algorithms have a false negative rate of 1.3% (Figure 2 and Figure 6). We also show in Section 4.1 that three parallel repetitions decrease the false negative rate to 0%.

We would also like to comment that a small accuracy loss due to randomization is universal in big data analytics, for example, in practically all data stream algorithms [22], locality sensitive hashing for nearest neighbor search [11], large graph analysis [33], etc. For the problem of tree similarity joins, first, a small number of false negatives may not affect the downstream applications. For example, if the subsequent task is clustering and each similar pair forms an edge, a few (random) missing edges in the input graph may not have a notable impact on the quality of the clustering output. To support this statement, we have conducted some experiments on clustering (delayed to Appendix B.4 in the full version due to space constraints). Our experiments show that the false negatives of our join schemes (without parallel repetition) will only cause at most 0.02% percent of nodes to be misclustered. Second, noise in the input data may obscure the algorithm's small accuracy loss. For example, in biological applications, the data reads obtained using the most advanced SMRT technology can have up to 12-18% errors.

4 EXPERIMENTS

In this section, we present our experimental studies of the SYNCSIG-NATURE framework and the two signature generation schemes.

The Setup. We abbreviate Algorithm 2 using Algorithm 4 for the signature generation as BJoin, and that using Algorithm 5 as EJoin. We set the neighborhood resolution c, the only parameter of our algorithms, to be 0.3 by default. We will discuss the influence of c on the running time and accuracy of the algorithms shortly.

We compare BJoin and EJoin with the previous best algorithm for the tree similarity join [14] (denoted by TJoin). The implementation of TJoin is available online.¹ We note that the partition-based algorithm proposed in [30], as well as algorithms in earlier works [13, 20, 37], have a considerably longer running time; these algorithms cannot finish in 10 hours in all of our experiments, and do not fit our figures if we want to provide a high resolution comparison between BJoin, EJoin, and TJoin. We thus choose to leave other algorithms out of the figures, and refer readers to [14] for a comparison.

The theoretical analysis of our algorithms necessitates a tree size of at least $\Omega(K)$. In our experiments, we test for K up to 40. Thus, for trees of sizes smaller than 100, we will use TJoin to find similar pairs. We note that a similar action was taken for the partition-based join algorithm Tang; see the discussion in the experimental section of [14].

¹The implementation can be obtained from https://frosch.cosy.sbg.ac.at/mpawlik/tedexperiments.

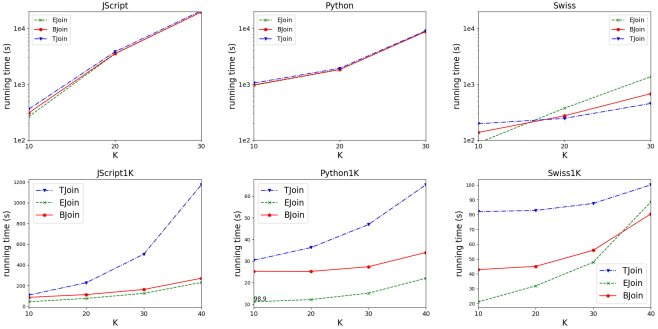


Figure 2: Running time comparisons in the single-thread setting. The number labels associated with points represent the accuracy; points without number labels have accuracy at least 99%

name	#trees	min. size	max. size	avg. size
Swiss1K	122, 772	1,000	48, 286	1,902
Swiss	565, 254	105	48, 286	917
Python1K	35, 754	1,000	46, 481	3,016
Python	148, 270	1	46, 481	948
JScript1K	39, 110	1,000	1, 716, 813	9,006
JScript	142, 373	4	1, 716, 813	2,619

Table 10: Statistics of datasets.

We use three large real-world datasets: (1) Swiss² : protein sequence data. (2) Python³: abstract syntax trees of Python files. (3) JScript⁴: abstract syntax trees of JavaScript files. To better appreciate the advantage of our algorithms on large trees, we also created datasets Swiss1K, Python1K, and JScript1K, which are obtained from Swiss, Python, and JScript after filtering out trees of sizes smaller than 1000. The statistics of the datasets we use in our experiments are summarized in Table 10.

This section consists of four parts. In the first part, we compare BJoin and EJoin with TJoin in the single-thread (sequential) computation setting. In the second part, we show the influence of the tree size and the parameter *c* on the accuracy and running time of BJoin and EJoin. After that, we compare all algorithms in the multi-thread (parallel) computation setting. Finally, we briefly compare BJoin and EJoin. We also test how false negatives of our algorithms affect a downstream application *clustering*, but due to space constraints, we leave this part to Appendix B.4. All algorithms are implemented in C++. All experiments are conducted on PowerEdge R740 server equipped with 2× Intel Xeon Gold 6248R 3.0GHz (24-core/48-thread per CPU) and 256GB RAM. All results are medians of 11 runs.

4.1 Comparisons in the Single-Thread Setting

We first compare BJoin and EJoin with the previous best algorithm TJoin [14] in the single-thread setting. The results are described in Figure 2.

The performance of three algorithms on full datasets (JScript, Python, and Swiss) are very similar. This is due to the fact that the number of output pairs is very large when we include all small trees. Consequently, the verification of the output pairs takes up the majority of the running time; In all plots for JScript and Python, the verification takes more than 98% of the total time. This phenomena is why we would like to test on truncated datasets to demonstrate that BJoin and EJoin outperform TJoin on large trees.

Due to the decreased output sizes, all algorithms on truncated datasets run substantially faster than that on full datasets. On all truncated datasets (JScript1K, Python1K, and Swiss1K), both BJoin and EJoin perform much better than TJoin, typically by a factor 2-4. On JScript1K and Python1K, the ratios increase when *K* increases, but the situation is different on Swiss1K, where the running times of BJoin and EJoin grow faster when *K* increases. This discrepancy could be related to the amount of output pairs produced by various datasets using different distance thresholds. Overall, BJoin and EJoin have better performance than TJoin even in the single-thread setting.

The accuracy of BJoin and EJoin is almost perfect (at least 99.9%) on the JScript, JScript1K, Swiss, and Swiss1K datasets, and is at least 98.9% on Python and Python1K.

 $^{^{2}} https://www.uniprot.org/downloads$

³https://files.sri.inf.ethz.ch/data/py150.tar.gz

⁴https://files.sri.inf.ethz.ch/data/js_dataset.tar.gz

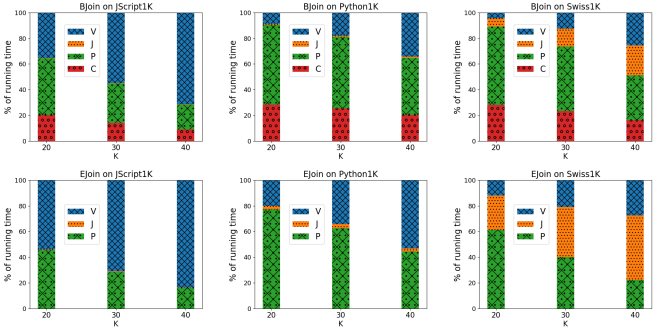


Figure 3: Running time percentages of binary-tree conversion (C), partition (P), join (J) and verification (V) of BJoin and EJOin

Running Time Distributions. In Figure 3, we present the running time of *binary-tree conversion, partition, join*, and *verification* of BJoin and EJoin on the three datasets. We can see that the verification step becomes more expensive when the distance threshold *K* increases, which is because the number of candidate pairs increases fast when *K* grows. On JScript and Python, the running time of the join step is negligible, while on Swiss, the join step makes a significant contribution when *K* is large. This may be because Swiss (biological data) is a different type of dataset compared with JScript and Python (abstract syntax trees). The binary-tree conversion step in BJoin takes a non-trivial percentage of the running time, which decreases percentage-wise as *K* increases. Note that EJoin does not have a binary-tree conversion step.

Parallel Repetition. Figure 4 shows the effect of parallel repetition on BJoin and EJoin. We find that repeating the signature generation and join steps twice increases the accuracy of both BJoin and EJoin to at least 99.7%, and repeating the steps three times already boosts the accuracy of both algorithms to 100%.

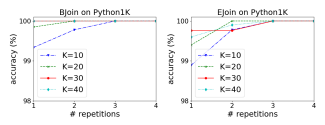


Figure 4: Accuracy Improvement using Parallel Repetition

4.2 Influence of Tree Size and Resolution c

Influence of the Tree Size. Our next set of experiments study the influence of the tree sizes on the performance of the three algorithms. Generally speaking, our algorithms have better and more stable performance when the tree sizes are large (say, at least 50*K* where *K* is the distance threshold). Due to space constraints, we leave this part to Appendix B.1.

Influence of Resolution c. Figure 5 presents the influence of neighborhood resolution c on the accuracy of BJoin and EJoin. We utilize truncated datasets rather than full datasets because the verification on the output pairs consumes the majority of the processing time on full datasets, making the influence of c harder to discern.

The experimental results are consistent with the theoretical prediction: the smaller the value of c (therefore z), the less likely we are to overlook a similar pair of trees. In Figure 5, we can see that both BJoin and EJoin achieve nearly perfect accuracy on JScript1K and Swiss1K datasets. While on Python1K, BJoin slightly outperforms EJoin, and both are above 98%.

Due to space constraints, we leave the experimental results on the influence of resolution c on the running time of the algorithms to Appendix B.2. Generally speaking, the value of c does *not* have a significant impact on the running time of BJoin and EJoin.

4.3 Comparisons in the Multi-Thread Setting

As described in Section 2, the main advantage of our SyncSignature framework is that it is fully parallelizable. We have implemented the multi-threading version of BJoin and EJoin and compare them with TJoin. To be fair, we also parallelize the verification step of TJoin. Recall that TJoin uses an index nested loop join which cannot be parallelized.

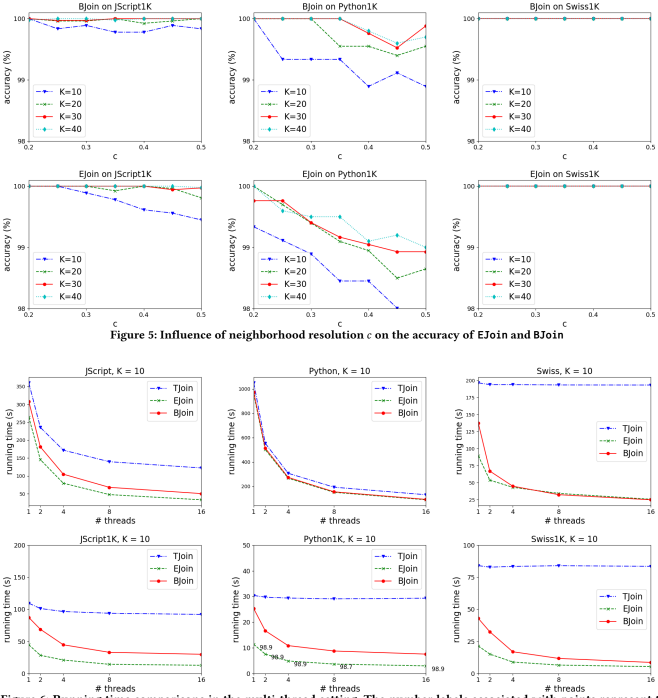


Figure 6: Running time comparisons in the multi-thread setting. The number labels associated with points represent the accuracy; points without number labels represent accuracy of at least 99%

The results on parallel executions of the algorithms are presented in Figure 6. We have delayed some results (for a different threshold distance K) in Figure 12 in Appendix B.3 due to space constraints. The *x*-axis of these figures stands for the number of threads we use in the parallel implementation. We observe that the running times of EJoin and BJoin decrease quickly when the number of threads increases. The trend slows when the number of threads becomes large, which is because the time spent on synchronizing/aggregating results from different threads becomes non-trivial.

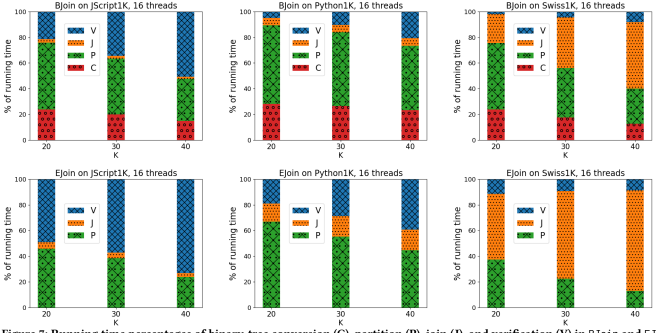


Figure 7: Running time percentages of binary-tree conversion (C), partition (P), join (J), and verification (V) in BJoin and EJoin using 16 threads

On Python, since the verification step takes up the majority of the time in all algorithms, the performance of the three algorithms are close. The situation is different on JScript, on which BJoin and EJoin outperform TJoin by a notable margin. On Swiss, the advantage of BJoin and EJoin is more significant.

On truncated datasets JScript1K, Python1K, and Swiss1K, the advantage of BJoin and EJoin over TJoin becomes significant. For K = 10, using 16 threads, BJoin and EJoin outperform TJoin by ratios of approximately 3 and 7 on JScript1K, ratios of 4 and 10 on Python1K, and ratios of 10 and 15 on Swiss1K. We note that the curves for TJoin on Swiss is quite flat, which is due to the fact that the join step in TJoin dominates the running time, and this step cannot be parallelized.

Figure 7 present the running time of *binary-tree conversion*, *partition*, *join*, and *verification* of BJoin and EJoin on the three datasets in the multi-thread setting. The running time distribution in the multi-thread setting is similar to that in the single-thread setting (see Figure 3), except that the time percentage of join is larger in the parallel setting, which is because the hash tables generated by different threads are merged sequentially in our implementation.

4.4 A Brief Comparison of BJoin and EJoin

Finally, we give a brief comparison between BJoin and EJoin. From what we have observed, the practical performance of the two algorithms are not entirely comparable. Generally speaking, EJoin performs slightly better than BJoin in terms of the running time, while the accuracy of BJoin is slightly better than that of EJoin.

There are two reasons why the running time of BJoin could be larger than that of EJoin. First, in BJoin we need to perform an additional binary-tree conversion step that is not needed in EJoin. This conversion step has a non-trivial contribution to the total running time, as seen in Figure 3. Second, we know from Theorem 3.2 and Theorem 3.4 that that the running time of *BALL-SIGNATURE* is $O(nz \log n)$, whereas that of *EULER-SIGNATURE* is only $O(n \log n)$. The actual difference of the two signature generation schemes is not as large as *z* because the constants hidden in the big-O notation are different. We also observe that sometimes BJoin performs better than EJoin when *K* is large, which is because $z \propto 1/K$ and $O(nz \log n)$ becomes smaller when *K* increases.

Regarding the accuracy, it could be due to some structural information loss in EJoin when finding the anchors in the Euler-tour of each tree. Fortunately, the accuracy loss is not significant and can be offset by two or three parallel repetitions as previously discussed.

5 CONCLUSION

In this paper, we propose a synchronized signature based algorithmic framework for solving tree similarity joins under edit distance, together with two concrete signature generation schemes. Our synchronized signature based algorithms are simple, efficient, yet parallelizable. They exceed the previous best algorithms for tree similarity joins in terms of running time in the centralized/singlethread environment at a small loss of accuracy, and have a substantially larger lead in the parallel/multi-thread context.

ACKNOWLEDGMENTS

This work was supported in part by the NSF CCF-1844234.

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