Joinable Parallel Balanced Binary Trees

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In this article, we show how a single function, \textit{join}, can be used to implement parallel \textit{balanced binary search trees} (BSTs) simply and efficiently. Based on \textit{join}, our approach applies to multiple balanced tree data structures, and a variety of functions for ordered sets and maps. We describe our technique as an algorithmic framework called \textit{join-based algorithms}. We show that the \textit{join} function fully captures what is needed for rebalancing trees for a variety of tree algorithms, as long as the balancing scheme satisfies certain properties, which we refer to as \textit{joinable} trees. We discuss four balancing schemes that are joinable: AVL trees, red-black trees, weight-balanced trees, and treaps. We present a variety of tree algorithms that apply to joinable trees, including \textit{insert}, \textit{delete}, \textit{union}, \textit{intersection}, \textit{difference}, \textit{split}, \textit{range}, \textit{filter}, and so on, most of them also parallel. These algorithms are generic across balancing schemes. Many algorithms are optimal in the comparison model, and we provide a general proof to show the efficiency in work for joinable trees. The algorithms are highly parallel, all with polylogarithmic span (parallel dependence). Specifically, the set-set operations \textit{union}, \textit{intersection}, and \textit{difference} have work $O(m \log(m/n + 1))$ and polylogarithmic span for input set sizes $n$ and $m \leq n$.

We implemented and tested our algorithms on the four balancing schemes. In general, all four schemes have quite similar performance, but the weight-balanced tree slightly outperforms the others. They have the same speedup characteristics, getting around 73× speedup on 72 cores (144 hyperthreads). Experimental results also show that our implementation outperforms existing parallel implementations, and our sequential version achieves close or much better performance than the sequential merging algorithm in C++ Standard Template Library (STL) on various input sizes.

CCS Concepts: • Theory of computation → Sorting and searching; Shared memory algorithms; • Computing methodologies → Shared memory algorithms;

Additional Key Words and Phrases: Balanced binary trees, searching, parallel, union

ACM Reference format:

This is an extended version for paper [14] published in Symposium on Parallel Algorithms and Architectures (SPAA) 2016. This research was supported by NSF grants CCF-2103483, CCF-1901381, CCF-1910030, and CCF-1919223. Authors’ addresses: G. Blelloch, Carnegie Mellon University, 5000 Forbes Avenue, Pittsburgh, PA 15213, USA; email: guyb@cs.cmu.edu; D. Ferizovic, Karlsruhe Institute of Technology, Kaiserstraße 12, Karlsruhe, 76131, Germany; email: dani93.f@gmail.com; Y. Sun, 900 University Ave, Riverside, CA 92521, USA; email: yihans@cs.ucr.edu.

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2329-4949/2022/04-ART7
https://doi.org/10.1145/3512769

ACM Transactions on Parallel Computing, Vol. 9, No. 2, Article 7. Publication date: April 2022.
1 INTRODUCTION

Many important and fundamental data structures used in algorithm design and programming are based on various forms of binary search trees (BSTs). Importantly, the in-order traversal of such trees provides an elegant and efficient mechanism to organize ordered data. Typically, we say a binary tree is “balanced” if the length of the path to all nodes (i.e., the depth) is asymptotically logarithmic in the number of nodes. In the sequential setting of “point” updates to the tree (i.e., single insertions or deletions), many balancing schemes for binary trees have been widely studied and are widely used, such as AVL trees [4], red-black trees [10], weight-balanced trees [52], splay trees [57], treaps [56], and many others. Some guarantee the logarithmic depth with high probability, or when amortized across node accesses.

With the recent prevalence of multi-processor (core) parallel computers, it is important to study balanced binary trees in the parallel setting. Although tree algorithms are fundamental and well studied in the sequential setting, applying them in the parallel setting is challenging. One approach is to make traditional updates such as insertions and deletions safe to apply concurrently. To work correctly, this requires various forms of locking and non-blocking synchronization [8, 9, 21, 24, 25, 30, 68]. Such concurrent updates can come with significant overhead, but more importantly only guarantee atomicity at the level of individual updates. Another approach is to update the trees in parallel by supporting “bulk” operations such as inserting a collection of keys (multi-insert), taking the union of two trees, or filtering the tree based on a predicate function. Here parallelism comes from within the operation. We refer to such algorithms as bulk parallel, or just parallel, as opposed to concurrent.

Recently, Blelloch et al. proposed an algorithmic framework [14] for such parallel bulk updates, which unifies multiple balancing schemes. Their algorithms have been applied to various applications [11, 60–62] and have been implemented in a parallel library PAM [59, 62]. Their framework bases all tree algorithms on a single primitive called join, which captures all the balancing criteria. As a result, many tree algorithms (most of them parallel) can be designed in a balancing-scheme-independent manner. Similar ideas of using join as a primitive for balanced binary trees, sequentially or in parallel, have also been considered in previous work [2, 3, 16, 64]. The original paper of Blelloch et al. about join-based algorithms shows that the framework works for four balancing schemes. However, the authors did not answer the more general question about what makes a balancing scheme fit in the join-based algorithm framework. This article extends the topic in [14] about efficient join-based parallel algorithms on binary trees, and specifically formalizes the preferred property for a balancing scheme to be joinable, which allows all join-based algorithms to be efficient on this balancing scheme.

As mentioned above, all the tree algorithms in this article are built on top of a single primitive join. For discussion purposes, we assume that a binary tree $T$ can be either empty (a nil-node), or can be a node with a left tree $T_L$, data entry $u$ (e.g., a key, or key-value), and a right tree $T_R$, denoted as node$(T_L, u, T_R)$. The function join$(T_L, k, T_R)$ for a given balancing scheme takes two balanced binary trees $T_L$ and $T_R$, and a key (or key-value) $k$ as input arguments, and returns a new valid balanced binary tree that has the same entries and the same in-order traversal as node$(T_L, k, T_R)$. The output has to be valid in satisfying the balancing criteria. We call the middle key $k$ the pivot of the join. An illustration of the join function is presented in Figure 1. In this article, we use join as the
Fig. 1. An example of the join algorithm on AVL trees. The function $\text{join}(T_L, k, T_R)$ takes two AVL trees $T_L$ and $T_R$, and key $k$ as input arguments, and returns a new valid AVL tree that has the same entries and the same in-order traversal as $\text{node}(T_L, k, T_R)$. The above figure shows one possible output of $\text{join}(T_L, k, T_R)$.

only primitive for connecting and rebalancing. Since rebalancing involves settling the balancing invariants, the join algorithm itself is specific to each balancing scheme. However, we note that in general, the join operation is oblivious to the values of the keys and depends only on the shape of the two trees.

One important observation of the join-based algorithms is that join captures all balancing criteria of each balancing scheme. As such, various tree algorithms (except join itself) can be implemented generically across balancing schemes. These algorithms need not be aware of the operated balancing scheme, but can just rely on the corresponding join algorithm for rebalancing. These join-based algorithms range from simple insertions and deletions, to more involved bulk operations. Moreover, the generality is not at the cost of efficiency in asymptotical cost or practical performance. In fact, the generic code leads to similar good performance for different balancing schemes (see Section 6). Also, all the join-based algorithms are still optimal in sequential work, and most of them can be parallelized with poly-logarithmic span (parallel dependency chain). These include some non-trivial and interesting theoretical results. For example, the join-based union algorithm, which combines two balanced binary search trees into one (keeping the ordering), costs $O(m \log(n/m + 1))$ work and $O(\log n \log m)$ span on two trees with sizes $m$ and $n \geq m$. This work bound is optimal under the comparison model [37].

To show the cost bounds of the join-based algorithms, one also must consider multiple balancing schemes. Fortunately, join also unifies the theoretical analysis. The key idea is to define a rank function for each balancing scheme, that maps every (sub-)tree to a real number, called its rank. The rank can be thought, roughly, as the abstract "height" of a tree. For example, for AVL trees the rank is simply the height of the tree, and for weight-balance trees, it is the log of the size. We then define a set of rules about rank and the join algorithm, that ensure a balancing scheme to be joinable. These rules apply to at least four existing balancing schemes. Based on these rules, we are able to show generic algorithms and analysis for all joinable trees. Our bounds for join-based algorithms then hold for any balancing scheme for which these properties hold.

We also implemented our algorithms and compare the performance with existing implementations, sequentially and in parallel. For merging two sets $S_1$ and $S_2$, we compare our union
algorithm to existing data structures that support batch updates of a batch $S_2$ on a tree of $S_1$. Our implementation has close performance when $|S_1|$ is large or close to $|S_2|$, and outperforms existing implementations by about $8\times$ when $|S_2|$ is much larger than $|S_1|$. We also compare our sequential version with STL tree and vector merging algorithms. Our implementation running on one core is about $8\times$ faster for union of two equal sized maps ($10^8$ each) than STL-tree, and over four orders of magnitude faster than STL-vector when one is of size $10^4$ and the other $10^8$. This is because our implementation is asymptotically more efficient than the STL implementation.

In the rest of the article, we will first show some preliminaries in Section 2. We will then present the useful properties that make a balancing scheme joinable in Section 3. We then present the join algorithms for four balancing schemes and show that they are all joinable in Section 4. In Section 5, we show several join-based algorithms, and present their cost bound. Finally, we show some experimental results in Section 6. Some related work and discussions are presented in Section 7.

2 PRELIMINARY

Parallel Cost Model. Our algorithms are based on nested fork-join parallelism and no other synchronization or communication among parallel tasks.\footnote{This does not preclude using our algorithms in a concurrent setting.} All analysis is using the binary-forking model [15], which is based on nested fork-join, but only allows for two branches in a fork. All our algorithms are deterministic.\footnote{Note that the bounds and the data structure themselves are not necessarily deterministic. For example, the treaps depends on random priorities. However, as long as the priorities are known (independently of the algorithms), the algorithm does not use randomization.} To analyze asymptotic costs of a parallel algorithm, we use work $W$ and span (or depth $D$), where work is the total number of operations and span is the length of the critical path. In the algorithm descriptions, the notation of $s_1 \parallel s_2$ indicates that statement $s_1$ and $s_2$ can run in parallel. Any computation with $W$ work and $D$ span will run in time $T < \frac{W}{P} + D$ assuming a PRAM (random access shared memory) [38] with $P$ processors and a greedy scheduler [18, 20, 34], or in time $T < \frac{W}{P} + D$ w.h.p. (defined below) when using a work-stealing scheduler [19]. We assume concurrent reads and exclusive writes (CREW).

Notation. We use $\langle \cdot, \cdot \rangle$ to denote a pair (similarly for tuples and sequences). We say $O(f(n))$ with high probability (w.h.p.) in $n$ to indicate $O(cf(n))$ with probability at least $1 - n^{-c}$ for $c \geq 1$. When clear from context, we drop the “in $n$.”

2.1 Balanced Binary Trees

A binary tree is either a nil-node, or a node consisting of a left binary tree $T_l$, a data entry $k$, and a right binary tree $T_r$, which is denoted as node($T_l$, $k$, $T_r$). We use the nil-node to refer to an external (empty) node with no data stored in it. The data entry can be simply a value (e.g., a key), or also hold data associated with the value. The size of a binary tree, or $|T|$, is 0 for a nil-node and $|T_l| + |T_r| + 1$ for a node($T_l$, $k$, $T_r$). The weight of a binary tree, or $w(T)$, is one more than its size (i.e., the number of nil-nodes in the tree). The height of a binary tree, or $h(T)$, is 0 for a nil-node, and $\max(h(T_l), h(T_r)) + 1$ for a node($T_l$, $k$, $T_r$). Parent, child, ancestor, and descendant are defined as usual (ancestor and descendant are inclusive of the node itself). We use $lc(T)$ and $rc(T)$ to extract the left and right child (subtree) of $T$, respectively. A node is called a leaf when both of its children are nil nodes. The left spine of a binary tree is the path of nodes from the root to a nil-node always following the left child, and the right spine the path to a leaf following the right child. The in-order values of a binary tree is the sequence of values returned by an in-order traversal of the tree. When the context is clear, we use a node $u$ to refer to the subtree $T_u$ rooted at $u$, and vice versa.
A **balancing scheme** for binary trees is an invariant (or set of invariants) that is true for every node of a tree, and is for the purpose of keeping the tree nearly balanced. In this article, we consider four balancing schemes that ensure the height of every tree of size $n$ is bounded by $O(\log n)$. When keys have a total order and the in-order values of the tree are consistent with the order, then we call it a BST. We note that the balancing schemes defined below, although typically applied to BSTs, do not require that the binary tree be a search tree.

We will then briefly introduce the balancing schemes that we use in this article. An illustration of these balancing schemes is shown in Figure 2.

**AVL Trees** [4]. AVL trees have the invariant that for every node $(T_l, e, T_r)$, the height of $T_l$ and $T_r$ differ by at most 1. This property implies that any AVL tree of size $n$ has height at most $\log_\phi (n + 1)$, where $\phi = \frac{1 + \sqrt{5}}{2}$ is the golden ratio.

**Red-Black (RB) Trees** [10]. RB trees associate a color with every node and maintain two invariants: (the red rule) no red node has a red child, and (the black rule) the number of black nodes on every path from the root down to a leaf is equal. All nil nodes are always black. Unlike some other presentations, we do not require that the root of a tree is black. Although this does not affect the correctness of our algorithms, our proof of the work bounds requires allowing a red root. We define the black height of a node $T$, denoted $h(T)$ to be the number of black nodes on a downward path from the node to a nil-node (inclusive of the node and the nil-node). Any RB tree of size $n$ has height at most $2 \log_2 (n + 1)$.

**Weight-Balanced (WB) Trees** [52]. WB trees with parameter $\alpha$ (also called BB$[\alpha]$ trees) maintain for every $T = \text{node}(T_l, e, T_r)$ the invariant $\alpha \leq \frac{w(T_l)}{w(T)} \leq 1 - \alpha$. We say two WB trees $T_1$ and $T_2$ have like weights if $\text{node}(T_1, e, T_2)$ is WB. Any $\alpha$ WB tree of size $n$ has height at most $\log \frac{1}{1 - \alpha} n$. For $\frac{\alpha}{1 - \alpha} < \alpha \leq 1 - \frac{1}{\sqrt{2}}$, insertion and deletion can be implemented on WB trees using just single and double rotations [17, 52]. We require the same condition for our implementation of join, and in particular use $\alpha = 0.29$ in experiments. We also denote $\beta = \frac{1 - \alpha}{\alpha}$, which means that either subtree cannot have a size that is more than $\beta$ times the size of the other subtree.

**Treaps** [56]. Treaps associate a uniformly random priority with every key and maintain the invariant that the priority of the key at each node is no greater than the priority of its two children. Any treap of size $n$ has height $O(\log n)$ w.h.p.. The priority can usually be decided by a hash value of the key, and thus for operations across search trees we assume equal keys have equal priorities. The notation we use for binary trees is summarized in Table 1. Some concepts in Table 1 will be introduced later in this article. In particular, we will define the rank($\cdot$) function for each balancing
The definition of some of them are postponed to Section 5.3.

scheme, which is generally based on their balancing criteria. We do so because that can help to simplify the proof of bounding the cost of the join-based algorithms on trees. We note that the rank functions are defined for the purpose of analysis. As long as the join function is implemented correctly as defined, all the join-based algorithms will behave as expected.

3 JOINABLE TREES

Here, we define the properties that make a tree joinable. These properties are used in later sections to prove bounds on each of our algorithms—i.e., if someone gives us a balancing scheme with these properties, our algorithms will be efficient without needing to know anything else about how balance is maintained. The concept of joinable trees relies on two subcomponents: a value associated with each tree node (subtree) called the rank, and a proper join algorithm. The definition of rank and the join algorithm depend on the balancing scheme. The rank of a tree node only relies on the shape of the subtree rooted at it, not the set of entries. For a tree $T$, we denote its rank as $\text{rank}(T)$.

We note that the joinable properties are mainly for the purpose of analysis. In fact, as long as the join function is implemented correctly (no need to be efficiently), all the join-based algorithms in later sections will also be correct. However, to use the general proof to show the cost bounds, these properties will help to simplify the analysis.

Definition 1 (Strongly Joinable Trees). A balancing scheme $S$ is strongly joinable, if we can assign a rank for each subtree from $S$, and there exists a join $(T_1, k, T_2)$ algorithm on two trees from $S$, such that the following rules hold:

1. **EMPTY RULE.** The rank of a nil-node is 0.
2. **MONOTONICITY RULE.** For $C = \text{join}(A, k, B)$, $\max(\text{rank}(A), \text{rank}(B)) \leq \text{rank}(C)$.
3. **SUBMODULARITY RULE.** Suppose $C = \text{node}(A, e, B)$ and $C' = \text{join}(A', e, B')$. If $0 \leq \text{rank}(A') - \text{rank}(A) \leq x$ and $0 \leq \text{rank}(B') - \text{rank}(B) \leq x$, then $0 \leq \text{rank}(C') - \text{rank}(C) \leq x$ (increasing side). In the other direction, if $0 \leq \text{rank}(A) - \text{rank}(A')$ and $0 \leq \text{rank}(B) - \text{rank}(B')$, then $0 \leq \text{rank}(C) - \text{rank}(C')$ (decreasing side).
4. **COST RULE.** $\text{join}(A, k, B)$ uses time $O(\|\text{rank}(A) - \text{rank}(B)\|)$.
5. **BALANCING RULE.** For a node $A$,

\[
\max(\text{rank}(\text{lc}(A)), \text{rank}(\text{rc}(A))) + c_l \leq \text{rank}(A) \leq \min(\text{rank}(\text{lc}(A)), \text{rank}(\text{rc}(A))) + c_u,
\]

where $c_l \leq 1$ and $c_u \geq 1$ are constants.
The last rule about balancing says that the ranks of a child and its parent cannot differ by much. This is not true for some randomization-based balancing schemes such as treaps. To generalize our results to such balancing schemes, we define a weakly joinable tree as follows.

**Definition 2 (Weakly Joinable Trees).** A balancing scheme $S$ is weakly joinable, if it satisfies the empty rule, monotonicity rule, and submodularity rule in Definition 2, and the relaxed balancing rule and weak cost rule as follows:

- **Relaxed Balancing Rule.** There exist constants $c_l, c_u, 0 < p_l \leq 1$ and $0 < p_u \leq 1$, such that for a node $A$ and any of its child $B$:
  1. $\text{rank}(B) \leq \text{rank}(A) - c_l$ happens with probability at least $p_l$.
  2. $\text{rank}(B) \geq \text{rank}(A) - c_u$ happens with probability at least $p_u$.
- **Weak Cost Rule.** $\text{join}(A, k, B)$ uses time $O(\text{rank}(A) + \text{rank}(B))$ w.h.p.

In this article, we use *joinable* to refer to weakly joinable trees. Later in this section, we will show that AVL trees, RB trees, and WB trees are strongly joinable, and treaps are weakly joinable. We also say a tree $T$ is strongly (weakly) joinable if $T$ is from a strongly (weakly) joinable balancing scheme.

Here we first present some properties of joinable trees.

**Property 1.** For a strongly joinable tree $T$, $c_l \text{h}(T) \leq \text{rank}(T) \leq c_u \text{h}(T)$.

This can be inferred directly from the balancing rule.

**Property 2 (Logarithmic Rank for Strongly Joinable Balanced Trees).** For a strongly joinable tree $T$, $\text{rank}(T) = O(\log w(T))$, where $w(T) = |T| + 1$ is $T$’s weight.

**Proof.** Let $f(r)$ be a function denoting the minimum weight that $T$ can have when $T$ has rank $r$, i.e., $f(r) = \min_{\text{rank}(T)=r} w(T)$. We now look at the two children of $T$, noted as $L$ and $R$. Without loss of generality (WLOG), assume $w(L) \geq w(R)$. In the balancing rule, we know that $\text{rank}(T) - c_u \leq \text{rank}(L) \leq \text{rank}(T) - c_l$. The total weight of $T$ is at most $2w(R)$. In other words,$$
\text{rank}(T) \leq f(r) \geq \min_{r-c_u \leq r' \leq r-c_l} f(r').
$$This means that $f(r) \geq c \cdot 2^{r/c_u}$ for some constant $c$. Thus, $\text{rank}(T)$ is at most $O(\log w(T))$. \hfill \Box

From the above lemma we have the following.

**Property 3 (Logarithmic Height for Strongly Joinable Balanced Trees).** For a strongly joinable tree $T$ with weight $n$, $h(T) = O(\text{rank}(T)) = O(\log n)$.

From the balancing rule we have the following.

**Property 4 (Similar Ranks for Balanced Strongly Joinable Trees).** For a strongly joinable tree $T = \text{node}(\text{lc}(T), k, \text{rc}(T))$, then $\text{rank}(\text{lc}(T))$ and $\text{rank}(\text{rc}(T))$ differ by at most a constant.

For weakly joinable trees, we also have a similar property for rank and height.

**Property 5 (Logarithmic Height for Weakly Joinable Balanced Trees).** For a weakly joinable tree $T$ with weight $n$, $h(T) = O(\text{rank}(T)) = O(\log n)$ w.h.p.

**Proof.** First of all, note that for a parent node $A$ and a child node $B$, $h(A) = h(B) + 1$, but $\text{rank}(A) \geq \text{rank}(B) + c_l$ happens with a constant probability. Thus, $h(T) = O(\text{rank}(T))$.

Consider a node $T$ with rank $r = \text{rank}(T)$. We follow the path and always go to the smaller subtree (with smaller weight). We call it a round every time we go down one level. After $m$ rounds we hit the leaf. Obviously, $m \leq \log n$ since every time we go to the smaller side of the subtree.
We note that with every such round, the rank increased by no more than \( c_u \) with a constant probability. Adding all the \( m \) rounds gets \( r \), which is the rank of \( T \). Based on Chernoff bound, we know that \( \Pr[r \geq (1 + \delta)c_u p_u m] \leq e^{-\frac{(1 + \delta)c_u p_u m}{3}} \). Let \( (1 + \delta)c_u p_u m \) be \( c \log n \) for sufficient large \( c \); we have

\[
\Pr[r \geq c \log n] \leq e^{-\frac{c \log m - c_u m}{3}} = n^{-c/3} \cdot e^{c_u p_u m} \leq n^{-c+c_u p_u/3}. \]

We then extend the **monotonicity rule** and prove the following theorem.

**Property 6 (Bounded Rank Increasing by Join).** For a joinable tree \( C = \text{join}(A, k, B) \), there exists a constant \( c' \leq c_u \), such that

\[
\max(\text{rank}(A), \text{rank}(B)) \leq \text{rank}(C) \leq \max(\text{rank}(A), \text{rank}(B)) + c'.
\]

**Proof.** The left half of the inequation holds because of the **monotonicity rule**.

For the right half, consider a single tree node \( v = \text{node}(\text{nil-node}, e, \text{nil-node}) \). \( \text{rank}(v) \) is at most \( c_u \) (the **balancing rule**). Consider \( v' = \text{join}(A, k, B) \), which increases the rank of both sides by at most \( \max(\text{rank}(A), \text{rank}(B)) \).

\[
\text{rank}(v') \leq \text{rank}(v) + \max(\text{rank}(A), \text{rank}(B)) \leq c_u + \max(\text{rank}(A), \text{rank}(B)).
\]

We then prove some useful definitions and theorems that will be used in later proofs of the join-based algorithms.

**Definition 3 (Layer \( i \) in Joinable Trees).** In a joinable tree, we say a node \( v \) in layer \( i \) if \( i \leq \text{rank}(v) < i + 1 \).

**Definition 4 (Rank Root).** In a joinable tree, we say a node \( v \) is a rank root, or a rank(\( i \))-root if \( v \) in layer \( i \) and \( v \)'s parent is not in layer \( i \). We use \( s_i(T) \) to denote the number of rank(\( i \))-root nodes in tree \( T \).

When the context is clear, we simply use \( s_i \).

**Property 7.** For a joinable tree, suppose a node \( T \) has rank \( r \); then the number of its descendants with rank more than \( r - 1 \) is a constant.

**Proof.** From the **balancing rule** we know that after \( 1/c_l \) generations from \( T \), the rank of the tree node must be smaller than \( r - 1 \). That is to say, there are at most \( 2^{1/c_l} \) such nodes in \( T \)'s subtree with rank no smaller than \( r - 1 \).

**Definition 5 (Rank Cluster).** For any rank root \( v \), its **rank cluster** contains all tree nodes with rank in the range \( (\text{rank}(v) - 1, \text{rank}(v)) \). We use \( d(v) \) of a rank root \( v \) to denote the size of its rank cluster.

For any \( v \), these rank clusters form a contiguous tree-structured component.

**Definition 6.** In a BST, a set of nodes \( V \) is called **ancestor-free** if and only if for any two nodes \( v_1, v_2 \) in \( V \), \( v_1 \) is not the ancestor of \( v_2 \).

Obviously, for each layer \( i \), all the rank(\( i \))-root nodes form an ancestor-free set.

The following lemma is important for proving the work bound of the join-based set-set algorithms in Section 5.3.
Lemma 1. There exists a constant \( t \) such that for a strongly joinable tree with size \( N \), \( s_i \leq \frac{N}{2^{(i/t)}} \). More precisely, \( t = 1 + \lceil c_u \rceil \).

Proof. We now organize all rank roots from layer \( kt \) to layer \((k+1)t-1\) into a group \( k \), for an integer \( k \). Assume there are \( s_k' \) such rank roots in root layer \( k \). We first prove that \( s_k' \leq s_{k-1}'/2 \), which indicates that \( s_k' \leq \frac{N}{2^k} \).

For a node \( u \) in group \( k \), we will show that its subtree contains at least two nodes in group \( k-1 \), one in each subtree. \( u \)'s rank is at least \( kt - 1 \) (because of rounding), but at most \((k+1)t-1\). For the left subtree (the right one is symmetric), we follow one path until we find a node in group \( k \) but its children \( v \) in a group smaller than \( k \), so \( \text{rank}(v) \leq k t - 1 \). We will prove that \( v \) must be in group \( k - 1 \). Because of the balancing rule, \( \text{rank}(u) - c_u \leq \text{rank}(v) \leq k t - 1 \). Considering the range of \( u \)'s rank, we have \((k-1)t \leq \text{rank}(v) \leq k t - 1 \). This means that \( v \) is in group \( k - 1 \). Therefore, every node in group \( k \) corresponds to at least two nodes in group \( k + 1 \). This proves \( s_k' \leq s_{k-1}'/2 \).

Obviously, each set of rank(i)-root is a subset of its corresponding group. This proves the above lemma.

Figure 14(b) shows an example of the layers of an AVL tree, in which we set the rank of a node to be its height. Because the rank of all AVL nodes are integers, all nodes are rank roots. Lemma 1 says that from the bottom up, every three layers, the number of nodes in an AVL shrinks by a half.\(^5\)

4 THE JOIN ALGORITHMS AND RANKS FOR EACH BALANCING SCHEME

Here we describe the join algorithms for the four balancing schemes we defined in Section 2, as well as the definition of rank for each of them. We will then prove they are joinable. For join, the pivot can be either just the data entry (such that the algorithm will create a new tree node for it), or a pre-allocated tree node in memory carrying the corresponding data entry (such that the node may be reused, allowing for in-place updates).

As mentioned in Section 1, join fully captures what is required to rebalance a tree and can be used as the only function that knows about and maintains the balance invariants. For AVL, RB, and WB trees, we show that join takes work that is proportional to the difference in the ranks of the two trees. For treaps, the work depends on the priority of the pivot. All the join algorithms are sequential so the span is equal to the work. We show in this article that the join algorithms for all balancing schemes we consider lead to optimal work for many functions on maps and sets.

At a high level, all the join algorithms try to find a “balancing point” in the larger tree, which is a node \( v \) on its left or right spine, and is balanced with the smaller tree, join at that point, and rebalance along the way back (see Figure 3). WLOG, we assume in join(T_L, k, T_R), \( T_L \) has larger or equal rank (e.g., for an AVL it means \( T_L \) has larger or equal height). In this case, we go along the right spine of \( T_L \) until we reach a node that is balanced with \( T_R \). We then connect \( v \) with \( T_R \) using \( k \), and let this new subtree replace \( v \) in \( T_L \). This guarantees that the subtree below \( v \) is balanced, but may cause imbalance in \( v \)'s ancestors. The algorithm then will try to rebalance the nodes on the way back to the root, and the rebalancing operations would depend on each specific balancing scheme. This high-level idea applies to multiple balancing schemes in this article (except for the treap join algorithm, which is slightly different because of the variant about priorities). In the following, we will elaborate each of them, and prove the cost bounds. The main result of this section is the following theorem.

Theorem 1. AVL, RB, and WB trees are strongly joinable, and treaps are weakly joinable.

---

\(^5\) Actually, considering that there is no rounding issue for AVL trees, this means that from the bottom up, every two layers, the number of nodes in an AVL tree shrinks by a half.
4.1 AVL Trees

For AVL trees, we define the rank to be the height, i.e., $\text{rank}(T) = h(T)$. Pseudocode for AVL join is given in Figure 4 and illustrated in Figure 5. Every node stores its own height so that $h(\cdot)$ takes constant time. If the two trees $T_L$ and $T_R$ differ by height at most one, join can simply create a new node$(T_l, k, T_r)$. However, if they differ by more than one, then rebalancing is required. Suppose that $h(T_l) > h(T_r) + 1$ (the other case is symmetric). The idea is to follow the right spine of $T_l$ until a node $c$ for which $h(c) \leq h(T_r) + 1$ is found (line 3). At this point, a new node$(c, k, T_r)$ is created to replace $c$ (line 4). Since either $h(c) = h(T_r)$ or $h(c) = h(T_r) + 1$, the new node satisfies the AVL invariant, and its height is one greater than $c$. The increase in height can increase the height of its ancestors, possibly invalidating the AVL invariant of those nodes. This can be fixed either with a double rotation if the invariant is invalid at the parent (line 6) or a single left rotation if the invariant is invalid higher in the tree (line 10), in both cases restoring the height for any further ancestor nodes. The algorithm will therefore require at most two rotations, as we summarized in the following lemma.

**Lemma 2.** The join algorithm in Figure 4 on AVL trees requires at most two rotations.

This lemma directly follows the algorithm description above.

**Lemma 3.** For two AVL trees $T_l$ and $T_r$, the AVL join algorithm works correctly, runs with $O(\max(h(T_l) - h(T_r)))$ work, and returns a tree satisfying the AVL invariant with height at most $1 + \max(h(T_l), h(T_r))$.

**Proof.** Consider the case where $T_L$ has a larger height (the other case is symmetric). After connecting the small tree with tree node $c$ (as described above and in Figure 4), all tree nodes below $k$ are guaranteed to be balanced. The only tree nodes that can be unbalanced are those on the right spine of $T_L$ and above $k$. The algorithm then checks them bottom-up. If the imbalance occurs upon connection, which is as shown in Figure 4, then the double-rotation shown in the figure fixes the imbalance. Note that, after rotation, the entire subtree height restores to $h + 2$, which is the same as the subtree size in $T_l$ before join. This means that no further rebalance is needed.

Another case is that, the imbalance does not directly occur at node $p$ ($k$’s parent in the output). Some other nodes on the right spine were affected to unbalanced because its right subtree height increased by 1. Assume the lowest such node is $B$, with left child $A$ and right child $C$. Assume $h(A) = h$. This case happens only when $h(C)$ was $h + 1$, but increased to $h + 2$ after join. In other words, $h(rc(C)) = h + 1$ after join, and $h(lc(C))$ must be $h$ (both before and after join, since the shape of $C$’s left subtree is not affected by join). In that case, a single rotation on $B$ will rebalance the subtree, getting a new subtree rooted at $C$ with height $h + 2$. Note that this is also the original height of the subtree in $T_L$ rooted at $B$. As a result, no further rebalance is needed.
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Fig. 4. The join algorithm on AVL trees. joinLeftAVL is symmetric to joinRightAVL.

```plaintext
1 joinRightAVL(T_l, k, T_r) {
2     (l, k', c) = expose(T_l);
3     if h(c) ≤ h(T_r) + 1 then {
4         T' = node(c, k, T_r);
5         if h(T') ≤ h(l) + 1 then return node(l, k', T');
6         else return rotateLeft(node(l, k', rotateRight(T')));
7     } else {
8         T' = joinRightAVL(c, k, T_r);
9         T'' = node(l, k', T');
10        if h(T') ≤ h(l) + 1 then
11           return T'';
12        else return rotateLeft(T''); }
13 }
14 join(T_l, k, T_r) {
15     if h(T_l) > h(T_r) + 1 then return joinRightAVL(T_l, k, T_r);
16     else if h(T_r) > h(T_l) + 1 then return joinLeftAVL(T_l, k, T_r);
17     else return node(T_l, k, T_r); }
```

Fig. 5. An example for join on AVL trees. An example for join on AVL trees (h(T_l) > h(T_r) + 1). We first follow the right spine of T_l until a subtree of height at most h(T_r) + 1 is found (i.e., T_2 rooted at c). Then, a new node(c, k, T_r) is created, replacing c (Step 1). If h(T_1) = h and h(T_2) = h + 1, the node p will no longer satisfy the AVL invariant. A double rotation (Step 2) restores both balance and its original height.

Since the algorithm only visits nodes on the path from the root to c, and only requires at most two rotations (Lemma 2), it does work proportional to the path length. The path length is no more than the difference in height of the two trees since the height of each consecutive node along the right spine of T_l differs by at least one. Along with the case when h(T_r) > h(T_l) + 1, which is symmetric, this gives the stated work bounds. The resulting tree satisfies the AVL invariants since rotations are used to restore the invariant. The height of any node can increase by at most one, so the height of the whole tree can increase by at most one. □

**Theorem 2.** AVL trees are strongly joinable.

**Proof.** The AVL invariant and the definition of AVL trees’ rank ensure the **empty rule** and **balancing rule** (c_l = 1, c_u = 2). Lemma 3 ensures the **cost rule** and **monotonicity rule**. We prove the **submodularity rule** as follows.

We start with the increasing side.

First note that the ranks of AVL trees are always integers. For C = node(A, k, B) and C' = join(A', k, B'), WLOG suppose rank(A) ≥ rank(B).

When 0 ≤ rank(A') − rank(A) ≤ x and 0 ≤ rank(B') − rank(B) ≤ x, the larger rank of A' and B' is within rank(A) + x. Then, the rank of C' is in the range [rank(A), rank(A) + x + 1] (Lemma 4). On
The other hand, $C = \text{node}(A, k, B)$, $\text{rank}(C)$ is no smaller than $\text{rank}(A) + 1$ (based on AVL invariants). Thus, $0 \leq \text{rank}(C) - \text{rank}(C') \leq x$ except for when $\text{rank}(C') = \text{rank}(A)$ and $\text{rank}(C) = \text{rank}(A) + 1$. We will show that this is impossible.

First of all, $\text{rank}(C') = \text{rank}(A)$ means that $\text{rank}(A') = \text{rank}(A)$. Also, $\text{rank}(B')$ must be at most $\text{rank}(A) - 2$, otherwise the $\text{join}$ will directly connect $A'$ and $B'$, and $\text{rank}(C')$ will be at least $\text{rank}(A) + 1$. Considering $\text{rank}(B') \geq \text{rank}(B)$, this means that $A$ and $B$ cannot be balanced, which leads to a contradiction.

This proves the increasing side of submodularity rule. We then prove the decreasing side. Both $\text{rank}(A)$ and $\text{rank}(B)$ are at most $\text{rank}(C) - 1$. Based on Lemma 3, $\text{joining}$ them back gets $C'$ with rank at most $\text{rank}(C)$.

\[ \square \]

### 4.2 RB Trees

In RB trees, $\text{rank}(T) = 2(\hat{h}(T) - 1)$ if $T$ is black and $\text{rank}(T) = 2\hat{h}(T) - 1$ if $T$ is red. Tarjan describes how to implement the $\text{join}$ function for RB trees [64]. Here, we describe a variant that does not assume the roots are black (this is to bound the increase in rank in some $\text{join}$-based algorithms). The pseudocode is given in Figure 6. We store at every node its black height $\hat{h}(\cdot)$. One special issue for the RB trees is to decide the color of the pivot. Therefore, we introduce a special parameter
flag in the join function of RB trees, which indicates if we want to assign a specific color to the pivot in certain circumstances. This will be used in other join-based algorithms in Section 5.2. In particular, we will set the color for the middle node (the new node to hold the pivot) to be red by default. However, when flag is true, and when the two input trees are already balanced, we will set the color for the middle node to black. This is to avoid decreasing in rank when we re-join two black subtrees with their black parent. In the join-based algorithms, we set this flag to be true only when the pivot we use is from a double-black node, i.e., when the node and all its children are black.

In the algorithm, the first case is when \( \hat{h}(T_v) = \hat{h}(T_l) \). As mentioned above, if flag is true, we will still use it as a black node and directly concatenate the two input trees. This increases the rank of the input trees by at most 2. Otherwise, if both root\((T_v) \) and root\((T_l) \) are black, we create red node\((T_1, k, T_r) \). When either root\((T_1) \) or root\((T_l) \) is red, we create black node\((T_1, k, T_r) \).

The second case is when \( \hat{h}(T_v) < \hat{h}(T_l) = \hat{h} \) (the third case is symmetric). Similarly to AVL trees, join follows the right spine of \( T_1 \) until it finds a black node \( c \) for which \( \hat{h}(c) = \hat{h}(T_v) \). It then creates a new red node\((c, k, T_r) \) to replace \( c \). Since both \( c \) and \( T_r \) have the same height, the only invariant that can be violated is the red rule on the root of \( T_r \), the new node, and its parent, which can all be red. In the worst case, we may have three red nodes in a row. This is fixed by a single rotation: if a black node \( v \) has rc\((v) \) and rc\((rc(v)) \) both red, we turn rc\((rc(v)) \) black and perform a single left rotation on \( v \), turning the new node black, and then perform a single left rotation on \( v \). The update is illustrated in Figure 7. The double- or triple-red issue is resolved after the rotation. The rotation, however, makes the new subtree root red, which can again violate the red rule between the root of the rotated tree and its parent. This imbalance can propagate to the root. If the original root of \( T_l \) is red, the algorithm may end up with a red root with a red child, in which case the root will be turned black, turning \( T_1 \) rank from \( 2\hat{h} - 1 \) to \( 2\hat{h} \). If the original root of \( T_l \) is black, the algorithm may end up with a red root with two black children, turning the rank of \( T_1 \) from \( 2\hat{h} - 2 \) to \( 2\hat{h} - 1 \). In both cases, the rank of the result tree is at most \( 1 + \text{rank}(T_l) \).

We note that the rank of the output can increase the larger rank of the input trees by 2 only when the pivot was a double-black node, and the two input trees are balanced and both have black roots. This additional condition is to guarantee the submodularity rule for RB trees.

Lemma 4. For two RB trees \( T_l \) and \( T_r \), the RB join algorithm works correctly, runs with \( O(|\text{rank}(T_l)| - \text{rank}(T_r)) \) work, and returns a tree satisfying the RB invariants and with rank at most \( 2 + \max(\text{rank}(T_l), \text{rank}(T_r)) \).

Proof. The base case where \( h(T_l) = h(T_r) \) is straightforward. For symmetry, here we only prove the case when \( h(T_l) > h(T_r) \). We prove the proposition by induction.

We first show the correctness. As shown in Figure 7, after appending \( T_r \) to \( T_l \), if \( p \) is black, the rebalance has been done, and the height of each node stays unchanged. Thus, the RB tree is still valid. Otherwise, \( p \) is red, and \( p \)'s parent \( g \) must be black. By applying a left rotation on \( p \) and \( g \), we get a balanced RB tree rooted at \( p \), except the root \( p \) is red. If \( p \) is the root of the whole tree, we change \( p \)'s color to black, and the height of the whole tree increases by 1. The RB tree is still valid. Otherwise, if the current parent of \( p \) (originally \( g \)'s parent) is black, the rebalance is done here. Otherwise, a similar rebalance is required on \( p \) and its current parent. Thus, finally we will either

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6In particular, in some of our algorithms, we may need to get \( t' = \text{join}(lc(t), t, \text{rc}(t)) \), which ideally should just get \( t' \) the same as the original (sub-)tree \( t \). However, consider when all tree nodes are black. If we call such join operation on all nodes in \( t \), setting pivot to be red will cause the rank of \( t' \) to be only a half of the original tree \( t \). This will affect the analysis of some join-based algorithms.

ACM Transactions on Parallel Computing, Vol. 9, No. 2, Article 7. Publication date: April 2022.
Fig. 7. An example of join on red-black trees. An example of join on red-black trees ($\hat{h} = \hat{h}(T_i) > \hat{h}(T_r)$). We follow the right spine of $T_i$ until we find a black node with the same black height as $T_r$ (i.e., c). Then a new red node(c, k, $T_r$) is created, replacing c (Step 1). The only invariant that can be violated is when either c’s previous parent $p$ or $T_r$’s root $d$ is red. If so, a left rotation is performed at some black node. Step 2 shows the rebalance when $p$ is red. The black height of the rotated subtree (now rooted at $p$) is the same as before ($h + 1$), but the parent of $p$ might be red, requiring another rotation. If the red-rule violation propagates to the root, the root is either colored red, or rotated left (Step 3).

(1) The rotation does not propagate to the root, and thus the rank of the tree remains as max($\hat{h}(T_i), \hat{h}(T_r)$).
(2) (Step 3 Case 1) The original root color is red, and thus a double-red issue occurs at the root and its right child. In this case, the root is colored black. The black height of the tree increases by 1, but since the original root is red, the rank increases by only 1.
(3) (Step 3 Case 2) The original root color is black, but the double-red issue occurs at the root’s child and grandchild. In this case, another rotation is applied as shown in Figure 7. The black height remains, but the root changed from black to red, increasing the rank by 1. □
THEOREM 3. RB trees are joinable.

PROOF. The RB invariant and the definition of RB trees’ rank ensures the empty rule and balancing rule ($c_f = 1, c_u = 2$). Lemma 4 ensures the cost rule and monotonicity rule. We prove the submodularity rule as follows.

We start with the increasing side.

First note that the ranks of RB trees are always integers. For $C = node(A, k, B)$ and $C' = join(A', e, B')$. WLOG suppose $rank(A) \geq rank(B)$.

When $0 \leq rank(A') - rank(A) \leq x$ and $0 \leq rank(B') - rank(B) \leq x$, the larger rank of $A'$ and $B'$ is within $rank(A) + x$. Then the rank of $C'$ is in the range $[rank(A), rank(A) + x + 2]$ (Lemma 4).

On the other hand, $C = node(A, k, B); C$ is either $rank(A) + 1$ or $rank(A) + 2$ (because all ranks are integers). Thus, $0 \leq rank(C') - rank(C) \leq x$ except for the following cases.

1. $rank(C') = rank(A) = r$.
2. $rank(C') = r + 1$, but $rank(C) = r + 2$.
3. $rank(C') = r + x + 2$, but $rank(C) = r + 1$.

We first show that case (1) is impossible.

First of all, $rank(A') \geq rank(A) = rank(C')$. On the other hand, $rank(C') \geq rank(A')$ because $C' = join(A', e, B')$. Therefore, $rank(A') = rank(C') = rank(A)$.

Also, $rank(B)$ is at least $r + 1$ because $A$ and $B$ are balanced. Considering $rank(B') \geq rank(B)$, $rank(B')$ is at least $r + 1$, but at most $rank(C') = r$.

If $A'$ and $B'$ are balanced, the rank of $C'$ is at least $rank(A) + 1$, which leads to a contradiction.

If $A'$ and $B'$ are not balanced, their ranks differ by only 1. This is only possible when one of them has black height $h$ with a black root, and the other one has black height $h - 1$ with a red root. However, the join algorithm results in double-red on the root’s child and its grandchild. After fixing it the rank also increases by 1, which leads to a contradiction.

We then show that case 2 is impossible.

First of all, $rank(B)$ is at least $rank(A) - 1$ because $A$ and $B$ are balanced. Considering $rank(B') \geq rank(B)$, $rank(B')$ is at least $rank(A') - 1$.

If $rank(C) = r + 2$, $C$ must be a double-black node. This means that $rank(A) = rank(B) = r$, and they are both black. $rank(A') \geq rank(A) = rank(C') - 1$. On the other hand, $rank(C') \geq rank(A')$ because $C' = join(A', k, B')$. Therefore, $rank(A') = rank(C')$ or $rank(A') = rank(C') - 1 = rank(A)$.

1. If $rank(A') = rank(C')$, from the same statement of case (1), we can show this is impossible.

   This is because the join algorithm will always lead to the result where $C'$ has rank larger than $A'$.

2. If $rank(A') = rank(C') - 1 = rank(A)$, $A'$ must be black since $rank(A)$ is even. In this case, $rank(B')$ is $r$, or $r + 1$.

   If $rank(A') = rank(B') = r$, $B'$ is also black. In this case, the algorithm will result in $C' = r + 2$. This leads to a contradiction.

   If $rank(B') = r + 1$, $B'$ is red but is still balanced with $A'$. In this case, the algorithm also results in $C'$ with rank $r + 2$. This also leads to a contradiction.

Finally, we prove that case (3) is impossible.

$rank(C') = r + x + 2$ means that $A'$, $B'$, and $C'$ are all black. Also, $A'$ and $B'$ must both have rank $r + x$ and black roots. This means that $C'$ (and also $C$) is a double-black node. Thus, $rank(C)$ must be $r + 2$, which leads to a contradiction.

This proves the increasing side of the submodularity rule. Next, we look at the decreasing side. There are two cases.
1. \textbf{joinRightWB} \((T_l, k, T_r)\) \\
2. \hspace{1em} \( (l, k’, c) = \text{expose}(T_l) \); \\
3. \hspace{1em} \textbf{if} (balance(|T_l|, |T_r|)) \textbf{then return} node\( (T_l, k, T_r) \); \textbf{else} \{ \\
4. \hspace{2em} \( T’ = \text{joinRightWB}(c, k, T_r) \); \\
5. \hspace{2em} \( (l_1, k_1, r_1) = \text{expose}(T’) \); \\
6. \hspace{2em} \textbf{if} \ like(l, T’[\|]) \textbf{then return} node\( (l, k’, T’) \); \\
7. \hspace{2em} \textbf{else if} (\|like(l, |l|)) \textbf{and} (\|like(|l| + |l_1|, r_1)) \textbf{then} \\
8. \hspace{3em} \textbf{return} \text{rotateLeft}(\text{node}(l, k’, \text{rotateRight}(T’))); \\
9. \hspace{2em} \textbf{else return} \text{rotateLeft}(\text{node}(l, k’, \text{rotateRight}(T’))); \\
\}

\textbf{joinLeftWB} \((T_l, k, T_r)\) \\
\hspace{1em} \textbf{if} heavy\( (T_l, T_r) \) \textbf{then return} \text{joinRightWB}(T_l, k, T_r); \\
\hspace{1em} \textbf{else if} heavy\( (T_r, T_l) \) \textbf{then return} \text{joinLeftWB}(T_l, k, T_r); \\
\hspace{1em} \textbf{else return} node\( (T_l, k, T_r) \); \\
\}

Fig. 8. The join algorithm on weight-balanced trees – joinLeftWB is symmetric to joinRightWB.

(1) Both \( rank(A’) < rank(A) \) and \( rank(B’) < rank(B) \) hold. They have to decrease by at least one because the rank of an RB tree is always an integer. First of all, we know that \( rank(C’) \leq \max(rank(A’), rank(B’)) + 2 \).

If \( rank(C’) = \max(rank(A’), rank(B’)) + 2 \), \( C’ \) must be a double-black node. \( rank(C) \) is \( rank(A) + 2 \) (also \( rank(B) + 2 \)). Also, \( A, B, C, A’, B’, \) and \( C’ \) are all black. Then, \( rank(A’) \) and \( rank(B’) \) can be at most \( rank(A) - 2 \) and \( rank(B) - 2 \), respectively. joining \( A’ \) and \( B’ \) increase the maximum rank by at most 2. Therefore, \( rank(C’) \) is no more than \( rank(C) \) holds.

If \( rank(C’) \leq \max(rank(A’), rank(B’)) + 1 \), joining them back results in an output tree of rank at most \( \max(rank(A’), rank(B’)) + 1 \). This proves \( rank(C’) \leq rank(C) \).

(2) Either \( rank(A’) = rank(A) \) or \( rank(B’) = rank(B) \). WLOG assume \( rank(A’) = rank(A) \) and \( rank(B’) \leq rank(B) \). There are three cases.

(a) \( A \) is black with rank \( 2h \), and \( C \) is red with rank \( 2h + 1 \). \( rank(B’) \leq rank(B) = 2h \).

(b) \( A \) is black with rank \( 2h \), and \( C \) is black with rank \( 2h + 2 \). \( rank(B’) \leq rank(B) \leq 2h + 1 \). When \( rank(B’) \leq 2h \), \( rank(C’) \leq rank(A’) + 2 = rank(C) \). When \( rank(B’) = 2h + 1 \), \( B \) is red and \( C’ \) is not a double-black node. Therefore, we also get \( rank(C’) \leq rank(B’) + 1 = rank(C) \).

(c) \( A \) is red with rank \( 2h + 1 \), and \( C \) is black with rank \( 2h + 2 \). \( rank(B’) \leq rank(B) \leq 2h + 1 \). Therefore, \( rank(C’) \leq rank(A’) + 1 = rank(C) \).

In summary, the \textit{submodularity rule} holds for RB trees. RB trees are strongly joinable. \hspace{1em} \Box

### 4.3 WB Trees

For WB trees, \( rank(T) = \log_2(w(T)) - 1 \). We store the weight of each subtree at every node. The algorithm for joining two WB trees is similar to that of AVL trees and RB trees. The pseudocode is shown in Figure 8. The \textit{like} function in the code returns true if the two input tree sizes are balanced based on the factor of \( \alpha \), and false otherwise. If \( T_l \) and \( T_r \) have like weights, the algorithm returns a new node\( (T_l, k, T_r) \).

Suppose \( |T_l| \leq |T_r| \); the algorithm follows the right branch of \( T_l \) until it reaches a node \( c \) with like weight to \( T_r \). It then creates a new node\( (c, r, T_r) \) replacing \( c \). The new node will have weight greater than \( c \) and therefore could imbalance the weight of \( c \)’s ancestors. This can be fixed with a single or double rotation (as shown in Figure 9) at each node assuming \( \alpha \) is within the bounds given in Section 2.
Fig. 9. An illustration of single and double rotations possibly needed to rebalance weight-balanced trees. In the figure, the subtree rooted at \( u \) has become heavier due to joining in \( T_L \) and its parent \( v \) now violates the balance invariant.

**Lemma 5.** For two \( \alpha \) WB trees \( T_L \) and \( T_R \) and \( \alpha \leq 1 - \frac{1}{\sqrt{2}} \approx 0.29 \), the WB join algorithm works correctly, runs with \( O(\lceil \log(w(T_L)/w(T_R)) \rceil) \) work, and returns a tree satisfying the \( \alpha \) WB invariant and with rank at most \( 1 + \max(\text{rank}(T_L), \text{rank}(T_R)) \).

The proof can be found in [13].

Notice that this upper bound is the same as the restriction on \( \alpha \) to yield a valid WB tree when inserting a single node. Then, we can induce that when the rebalance process reaches the root, the new WB tree is valid. The proof is intuitively similar as the proof stated in [17, 52], which proved that when \( 2^{-11} \leq \alpha \leq 1 - \frac{1}{\sqrt{2}} \), the rotation will rebalance the tree after one single insertion. In fact, in our join algorithm, the “inserted” subtree must be along the left or right spine, which actually makes the analysis easier.

**Theorem 4.** WB trees are strongly joinable when \( \alpha \leq 1 - \frac{1}{\sqrt{2}} \approx 0.29 \).

**Proof.** The WB invariant and the definition of WB’s rank ensures the empty rule and balancing rule \((c_l = \log_{1-\alpha} 2, c_u = \log_{\alpha} 2)\). Lemma 5 ensures the cost rule and the monotonicity rule.

For the submodularity rule, note that \( \text{rank}(C) = \log_2(w(A) + w(B)) \), and \( \text{rank}(C') = \log_2(w(A') + w(B')) \). When either \( A \) or \( B \) changes its weight by more than \( \log_2 x \), obviously the total weight of \( C \) will not increase or decrease by a factor of \( \log_2 x \).

\[ \square \]

### 4.4 Treaps

For treaps \( \text{rank}(T) = \log_2(w(T)) - 1 \). The treap join algorithm (as in Figure 10) first picks the key with the highest priority among \( k, k(T_L), \) and \( k(T_R) \) as the root. If \( k \) is the root, then we can return node\((T_L, k, T_R)\). Otherwise, WLOG, assume \( k(T_L) \) has a higher priority. In this case, \( k(T_L) \) will be the root of the result, \( lc(T_L) \) will be the left tree, and \( rc(T_L), k, \) and \( T_R \) will form the right tree. Thus, join recursively calls itself on \( rc(T_L), k, \) and \( T_R \) and uses the result as \( k(T_L) \)'s right child. When \( k(T_R) \) has a higher priority, the case is symmetric. The cost of join is therefore the depth of the key \( k \) in the resulting tree (each recursive call pushes it down one level). In treaps, the shape of the result tree, and hence the depth of \( k \), depend only on the keys and priorities and not the history. Specifically,
joinTreap(T₁, k, Tᵣ) { 
if prior(k, k₁) and prior(k, k₂) then return node(T₁, k, Tᵣ) else { 
(l₁, k₁, r₁) = expose(T₁); 
(l₂, k₂, r₂) = expose(Tᵣ); 
if prior(k₁, k₂) then return node(l₁, k₁, joinTreap(r₁, k, Tᵣ)); 
else return node(joinTreap(l₁, k, l₂), k₂, r₂); }
}

Fig. 10. The join algorithm on treaps. prior(k₁, k₂) decides if the node k₁ has a higher priority than k₂.

if a key has the t-th highest priority among the keys, then its expected depth in a treap is O(log t) (also whp). If it is the highest priority, for example, then it remains at the root.

Lemma 6. For two treaps T₁ and Tᵣ, if the priority of the pivot k is the t-th highest among all keys in T₁ ∪ {k} ∪ Tᵣ, the treap join algorithm works correctly, runs with O(log t + 1) work in expectation and whp, and returns a tree satisfying the treap invariant with rank at most 1 + max(rank(T₁), rank(Tᵣ)).

Theorem 5. Treaps are weakly joinable.

Proof. The empty rule and monotonicity rule hold from the definition of rank. The submodularity rule holds for the same reason as the WB tree.

For the relaxed balancing rule, note that the weight of the tree shrinks by a factor of 1/3 to 2/3 with probability 1/3. This means that going down from a parent to a child, the rank of a node decreases by a constant between log₂ 3 and log₂ 3/2 with probability 1/3. This proves the relaxed balancing rule.

For weak cost rule, note that the cost of join is at most the height of the larger input tree, which is O(rank(T)) whp. □

We also present Lemma 7, which is useful in the proofs in a later section. Recall that we use d(v) to denote the size of a rank root v’s rank cluster. In treaps, each rank cluster is a chain. This is because if two children of one node v are both in layer i, the weight of v is more than 2i+1, meaning that v should be layer i + 1. This means that for a treap node v, d(v) is the number of generations to take from v to reach a node u with rank at most rank(v) − 1. The lemma below means that the expected size of a rank cluster E[d(v)] is also a constant.

Lemma 7. If v is a rank root in a treap, d(v) is less than a constant in expectation.

Proof. Consider a rank(k)-root v that has weight N ∈ [2ᵏ, 2ᵏ₊₁). The probability that d(v) ≥ 2 is equal to the probability that one of its grandchildren has weight at least 2ᵏ. This probability P is

\[
P = \frac{1}{2^k} \sum_{i=2^k+1}^{N} \frac{i - 2^k}{i}
\]

\[
\leq \frac{1}{2^k} \sum_{i=2^k+1}^{2^k₊₁} \frac{i - 2^k}{i}
\]

\[
\approx 1 - \ln 2.
\]

We denote 1 − ln 2 as pₑ. Similarly, the probability that d(v) ≥ 4 should be less than pₑ², and the probability shrinks geometrically as d(v) increase. Thus, the expected value of d(v) is a constant. □

Finally, we prove that Lemma 1 also holds for treaps.
Joinable Parallel Balanced Binary Trees

5 ALGORITHMS USING JOIN

The join function, as a subroutine, has been used and studied by many researchers and programmers to implement more general set operations. In this section, we describe algorithms for various functions that use just join for rebalancing. The algorithms are generic across balancing schemes. The pseudocodes for the algorithms in this section are shown in Figure 13. Beyond join, the only access to the trees we make use of is through expose, which returns the left child, root entry, and the right child. This can be done by simply reading the root. The set-set operations, including union, intersection, and difference, are efficient in work, which means the work of the algorithm is asymptotically the same as the best known sequential algorithm. In particular, the cost of the join-based algorithms are optimal in the number of comparisons performed (see more details below). The pseudocode for all the algorithms introduced in this section is presented in Figures 11, 13, and 15. We note that for RB trees, for every invocation of join, we need to give an extra flag to indicate if the pivot of join is a double-black node (defined in Section 4.2). For simplicity, this is not shown in the code given. We note that this is used only to ensure the desired theoretical bound, and removing this condition does not affect the correctness.

As mentioned, all the join-based algorithms use join as a black box. As long as the join algorithm is implemented correctly as defined, the join-based algorithms can behave as expected. Also, the algorithms themselves do not require the rank function. However, to prove the efficiency in work and span, we need more careful design and use of rank functions.

We present a summary of the cost of all join-based algorithms discussed in the article in Table 2.

5.1 Two Helper Functions: split and join2

We start with presenting two helper functions split and join2. For a BST $T$ and key $k$, split$(T, k)$ returns a triple $(T_L, b, T_R)$, where $T_L$ ($T_R$) is a tree containing all keys in $T$ that are smaller (larger) than $k$, and $b$ is a flag indicating whether $k \in T$. For two binary trees $T_L$ and $T_R$, not necessarily BSTs, join2$(T_L, T_R)$ returns a binary tree for which the in-order values are the concatenation of the in-order values of the binary trees $T_L$ and $T_R$ (the same as join but without the middle key). For BSTs, all keys in $T_L$ have to be less than keys in $T_R$.
Table 2. The Core join-based Algorithms and Their Asymptotic Costs – The Cost is Given Under the Assumption That All Parameter Functions Take Constant Time to Return

<table>
<thead>
<tr>
<th>Function</th>
<th>Work</th>
<th>Span</th>
</tr>
</thead>
<tbody>
<tr>
<td>insert, delete, update, find, first, last, range, split, join2, previous, next, rank, select, up_to, down_to</td>
<td>$O(\log n)$</td>
<td>$O(\log n)$</td>
</tr>
<tr>
<td>union, intersection, difference</td>
<td>$O\left(m \log \left(\frac{n}{m} + 1\right)\right)$</td>
<td>$O(\log n \log m)$</td>
</tr>
<tr>
<td>map, reduce, map_reduce, to_array</td>
<td>$O(n)$</td>
<td>$O(\log n)$</td>
</tr>
<tr>
<td>build, filter</td>
<td>$O(n)$</td>
<td>$O(\log^2 n)$</td>
</tr>
</tbody>
</table>

For functions with two input trees (union, intersection and difference), $n$ is the size of the larger input, and $m$ of the smaller.

Although both sequential, these two functions, along with the join function, are essential to parallelize other algorithms. Intuitively, when processing a tree in parallel, we recurse on two subcomponents of the tree in parallel by splitting the tree by some key. After the recursions return, we combine the results of the left and right parts, with or without the middle key, using join or join2. Because of the balance of the tree, this framework usually gives high parallelism with low span (e.g., poly-logarithmic).

**Split.** As mentioned above, split$(T, k)$ splits a BST $T$ by a key $k$ into $T_L$ and $T_R$, along with a bit $b$ indicating if $k \in T$. Intuitively, the split algorithm first searches for $k$ in $T$, splitting the tree along the path into three parts: keys to the left of the path, $k$ itself (if it exists), and keys to the right. Then by applying join, the algorithm merges all the subtrees on the left side (using keys on the path as intermediate nodes) from bottom to top to form $T_L$, and merges the right parts to form $T_R$. Writing the code in a recursive manner, this algorithm first determines if $k$ falls in the left (right) subtree, or is exactly the root. If it is the root, then the algorithm directly returns the left and the right subtrees as the two return trees and true as the bit $b$. Otherwise, WLOG, suppose $k$ falls in the left subtree. The algorithm further split the left subtree into $T_{L'}$ and $T_{R'}$ with the return bit $b'$. Then, the return bit $b = b'$, the $T_L$ in the final result will be $T_{L'}$, and $T_r$ means to join $T_{R'}$ with the original right subtree by the original root. Figure 12 gives an example.

The cost of the algorithm is proportional to the rank of the tree. Intuitively (and informally), this is because the cost of split is the sum of the cost of a sequence of join algorithms, which is $O(r(T_{L}) - r(T_{R}))$. For the chain of trees involved, the middle terms cancel out, and the total cost is then $O(r(T))$. We summarize and prove the cost of the split algorithm in the following theorem.

**Theorem 7.** The work of split$(T, k)$ is $O(h(T))$ for all strongly joinable trees and treaps. The two resulting trees $T_L$ and $T_R$ will have rank at most rank$(T) + c_0$ for some constant $c_0$.
Joinable Parallel Balanced Binary Trees

PROOF. We only consider the work of joining all subtrees on the left side. The other side is symmetric. Suppose we have \( l \) subtrees on the left side, denoted from bottom to top as \( T_1, T_2, \ldots, T_l \). As stated above, we consecutively join \( T_1 \) and \( T_2 \) returning \( T'_2 \), then join \( T'_2 \) with \( T_3 \) returning \( T'_3 \) and so forth, until all trees are merged. The overall work of \textit{split} is the sum of the cost of \( l - 1 \) \textit{join} functions. One observation is that, the pivot of the \textit{join} of \( T'_{l-1} \) and \( T_l \), denoted as \( e_l \), used to be \( T'_l \)’s parent. Meanwhile, \( T'_{l-1} \) is a subset of \( T'_l \)’s original sibling, denoted as \( X_i \). We use \( T(e_l) = \text{node}(X_i, e_l, T_l) \) to denote the original subtree rooted at \( e_l \) in the input.

We first prove by induction that computing \( T'_l = \text{join}(T'_{l-1}, e_l, T_l) \) gets \( T'_l \) with rank no more than \( \text{rank}(X_{l+1}) \).

From the induction hypothesis, we know that \( \text{rank}(T'_{l-1}) \leq r(X_i) \). Considering \( T'_l = \text{join}(T'_{l-1}, e_l, T_l) \) and \( T(e_l) = \text{node}(X_i, e_l, T_l) \), from the \textbf{submodularity rule}, we can get \( \text{rank}(T'_l) \leq \text{rank}(T(e_l)) \). Considering \( T(e_l) \) is a subtree in \( X_{l+1} \), we have

\[
\text{rank}(T'_l) \leq \text{rank}(T(e_l)) \leq \text{rank}(X_{l+1}).
\]

We next prove the cost. The cost of the \( i \)-th \textit{join} is \( W_i \leq c \cdot \text{rank}(T_i) - \text{rank}(T'_{i-1}) \). Note that \( \text{rank}(T'_{l-1}) \leq \text{rank}(X_i) \leq \text{rank}(T_i) + c_0 - c_1 \). Also, note that \( T'_l \) is achieved by joining \( T_i \) and another tree. Therefore, \( \text{rank}(T_i) - \text{rank}(T'_{i-1}) \leq \text{rank}(T'_l) - \text{rank}(T'_{i-1}) \).

This means that either \( W_i = c \cdot \text{rank}(T'_l) - \text{rank}(T'_{i-1}) \), or \( W_i \) is a constant no more than \( c_2 = c \cdot (c_0 - c_1) \). Therefore, \( W_i \leq c \cdot \text{rank}(T'_l) - \text{rank}(T'_{i-1}) + 2c_2 \).

\[
\sum_{i=1}^{h(T)} W_i \leq \sum_{i=1}^{h(T)} c \cdot \text{rank}(T'_l) - \text{rank}(T'_{i-1}) + 2c_2 \\
\leq 2c \cdot c_2 \cdot h(T) + \text{rank}(T'_l) \\
\leq O(h(T)) + \text{rank}(T(e_h(T))) \leq O(h(T)).
\]
For treaps, each join uses the key with the highest priority since the key is always on an upper level. Hence, by Lemma 6, the complexity of each join is $O(1)$ and the work of split is at most $O(h(T))$. Obviously for treaps we have $\text{rank}(T_l)$ and $\text{rank}(T_r)$ at most $\text{rank}(T)$. 

**Join2.** As stated above, the join2 function is defined similar to join without the middle entry. The join2 algorithm first choose one of the input trees, and extract its last (if it is $T_l$) or first (if it is $T_r$) element $k$. The two cases take the same asymptotical cost. The extracting process is similar to the split algorithm. The algorithm then uses $k$ as the pivot to join the two trees. In the code shown in Figure 11, the split_last algorithm first finds the last element $k$ (by following the right spine) in $T_l$ and on the way back to root, joins the subtrees along the path. We denote the result of dropping $k$ in $T_L$ as $T'$. Then, $\text{join} (T', k, T_r)$ is the result of join2. Unlike join, the work of join2 is proportional to the rank of both trees since both split and join take at most logarithmic work.

**Theorem 8.** The work of $T = \text{join2}(T_l, T_r)$ is $O(\text{rank}(T_l) + \text{rank}(T_r))$ for all joinable trees. Furthermore, $\text{rank}(T) \leq \max(\text{rank}(T_l), \text{rank}(T_r)) + c'$, where $c'$ is defined in Theorem 6.

**Proof.** The cost bound holds because split_last and join both take work asymptotically no more than the larger tree rank. We next prove the range of rank($T$). First of all, splitting the last from $T_l$ only decreases its rank (Theorem 7). Therefore, based on Theorem 6, $T = \text{join} (T', k, T_r)$ has rank no more than $\max(\text{rank}(T_r), \text{rank}(T_l)) + c'$. This proves the theorem. 

### 5.2 Set Functions Using join

In this section, we will present join-based algorithms on BSTs for set-set functions, including union, intersection, and difference. Many other set operations, such as symmetric difference, can be implemented by a combination of union, intersection, and difference with no extra asymptotical work. We will start with presenting some background of these algorithms, and then explain in detail about the join-based algorithms. Finally, we show the proof of their cost bound.

**Background.** The parallel set functions are particularly useful when using parallel machines since they can support parallel bulk updates. As mentioned, although supporting efficient algorithms for basic tree operations (e.g., insertion and deletion) are rather straightforward, it is more difficult to implement efficient bulk operations. This is more challenging considering parallelism and dealing with multiple balancing schemes. For example, combining two ordered sets of size $n$ and $m \leq n$ in the format of two arrays would take work $O(m + n)$ using the standard merging algorithm in the merge sort algorithm. In this case, even inserting a single element into a set of size $n$ will have linear cost. Another simple implementation is to store both sets as balanced trees, and insert the elements in the smaller tree into the larger one, costing $O(m \log n)$ work. It overcomes the issue of redundant scanning and copying, because many subtrees in the larger tree remain untouched. However, this results in $O(n \log n)$ work for combining two ordered sets of the same size $n$, while it is easy to make it $O(n)$ by arrays. This is because the algorithm fails to make use of the ordering in the smaller tree.

The lower bound for comparison-based algorithms for union, intersection, and difference for inputs of size $n$ and $m \leq n$, and returning an ordered structure,\(^7\) is $\log_2 \binom{m+n}{n} = \Theta(m \log(n / m + 1))$, which is the number of possible ways $n$ keys can be interleaved with $m$ keys [37]. Brown and Tarjan first matched these bounds, asymptotically, using a sequential algorithm based on RB trees [22]. Although designed for merging, the algorithm can be adapted for union, intersection, and difference with the same bounds. The bound is interesting since it shows that implementing insertion with

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\(^7\)By “ordered structure,” we mean any data structure that can output elements in sorted order without any further comparisons (e.g., a sorted array, or a binary search tree).
union, or deletion with difference, is asymptotically efficient (\(O(\log n)\) time), as is taking the union of two equal sized sets (\(O(n)\) time). However, the Brown and Tarjan algorithm is complicated, only works on RB trees, and is completely sequential.

Adams later described very elegant algorithms for union, intersection, and difference, as well as other functions based on join [2, 3]. Adams’ algorithms were proposed in an international competition for the Standard ML community, which is about implementations on “set of integers.” Prizes were awarded in two categories: fastest algorithm, and most elegant yet still efficient program. Adams won the elegance award, while his algorithm is almost as fast as the fastest program for very large sets, and was faster for smaller sets. Because of the elegance of the algorithm, at least three libraries use Adams’ algorithms for their implementation of ordered sets and tables (Haskell [48] and MIT/GNU Scheme, and SML). The idea of Adams’ algorithm enlightens our parallel join-based set algorithms and the implementation in the PAM library, for which the sequential version on WB trees is exactly the same as Adams’ algorithm.

Although only considering WB trees, Adams’ algorithms actually show that in principle all balance criteria for search trees can be captured by the single function join. As long as a valid join algorithm on a certain balancing scheme is provided, the correctness of the join-based set operations can be guaranteed on the corresponding balancing scheme.

Surprisingly, however, there have been almost no results on bounding the work (time) of Adams’ algorithms, in general or on specific tree types. Adams informally argues that his algorithms take \(O(n + m)\) work for WB tree, but that is a very loose bound. Blelloch and Reid-Miller later show that similar algorithms for treaps [16] are optimal for work (i.e., \(\Theta(m \log(\frac{n}{m} + 1))\)), and are also parallel. Their algorithms, however, are specific for treaps. The problem with bounding the work of Adams’ algorithms, is that just bounding the time of split, join, and join2 with logarithmic costs is not sufficient.\(^8\) One needs additional properties of the trees. As a result, there is no tight bound even for Adams’ original algorithms on WB trees, not to mention other balancing schemes.

Our work gives the first work-optimal bounds for the join-based algorithms for all four balancing schemes in Section 2. We show that with the join algorithms in Section 4, we achieve asymptotically optimal bounds on work for the set operations. These bounds hold when either input tree is larger (this was surprising to us). Furthermore, the algorithms have \(O(\log n \log m)\) span, and hence are highly parallel.

**Algorithms.** \(\text{union}(T_1, T_2)\) takes two BSTs and returns a BST that contains the union of all keys. The algorithm uses a classic divide-and-conquer strategy, which is parallel. At each level of recursion, \(T_1\) is split by \(k(T_2)\), breaking \(T_1\) into three parts: one with all keys smaller than \(k(T_2)\) (denoted as \(L_1\)), one in the middle either of only one key equal to \(k(T_2)\) (when \(k(T_2) \in T_1\)) or empty (when \(k(T_2) \not\in T_1\)), and the third one with all keys larger than \(k(T_2)\) (denoted as \(R_1\)). Then two recursive calls to \(\text{union}\) are made in parallel. One unions \(\text{lc}(T_2)\) with \(L_1\), returning \(T_1\), and the other one unions \(\text{rc}(T_2)\) with \(R_1\), returning \(T_r\). Finally, the algorithm returns \(\text{join}(T_1, k(T_2), T_r)\), which is valid since \(k(T_2)\) is greater than all keys in \(T_1\) are less than all keys in \(T_r\).

The functions \(\text{intersection}(T_1, T_2)\) and \(\text{difference}(T_1, T_2)\) take the intersection and difference of the keys in their sets, respectively. The algorithms are similar to \(\text{union}\) in that they use one tree to split the other. However, the method for joining and the base cases are different. For \(\text{intersection}\), \(\text{join2}\) is used instead of \(\text{join}\) if the root of the first is not found in the second. Accordingly, the base case for the \(\text{intersection}\) algorithm is to return an empty set when either set is empty. For \(\text{difference}\),

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\(^8\)Bounding the cost of \(\text{join}, \text{split}, \) and \(\text{join2}\) by the logarithm of the smaller tree is probably sufficient, but implementing a data structure with such bounds is very much more complicated.
Table 3. Descriptions of Notations Used in Section 5.3

<table>
<thead>
<tr>
<th>Notation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T_p$</td>
<td>The pivot tree</td>
</tr>
<tr>
<td>$T_d$</td>
<td>The decomposed tree</td>
</tr>
<tr>
<td>$n$</td>
<td>$\max(</td>
</tr>
<tr>
<td>$m$</td>
<td>$\min(</td>
</tr>
<tr>
<td>$T_p(v), v \in T_p$</td>
<td>The subtree rooted at $v$ in $T_p$</td>
</tr>
<tr>
<td>$T_d(v), v \in T_p$</td>
<td>The tree from $T_d$ that $v$ splits $^9$</td>
</tr>
<tr>
<td>$s_i$</td>
<td>The number of nodes in layer $i$ (Definition 4)</td>
</tr>
<tr>
<td>$v_{kj}$</td>
<td>The $j$-th node on layer $k$ in $T_p$</td>
</tr>
<tr>
<td>$d(v)$</td>
<td>The size of the rank cluster of a rank root $v$ (Definition 5)</td>
</tr>
</tbody>
</table>

$join2$ is used anyway because $k(T_2)$ should be excluded in the result tree. The base cases are also different in the obvious way.

The cost of the algorithms described above can be summarized in the following theorem.

**Theorem 9.** For all strongly joinable trees (and treaps), the work and span of the algorithm (as shown in Figure 13) of union, intersection, or difference on two balanced BSTs of sizes $m$ and $n$ ($n \geq m$) is $O(m \log(\frac{n}{m} + 1))$ (in expectation for treaps) and $O(\log n \log m)$, respectively (w.h.p. for treaps).

The work bound for these algorithms is optimal in the comparison-based model. In particular, considering all possible interleavings, the minimum number of comparisons required to distinguish them is $\log \left( \binom{m+n}{n} \right) = \Theta(m \log(\frac{n}{m} + 1))$ [37]. A generic proof of Theorem 9 working for four balancing schemes will be shown in the next section. The span of these algorithms can be reduced to $O(\log m)$ for WB trees even on the binary-forking model [15] by doing a more complicated divide-and-conquer strategy.

### 5.3 The Proof of Theorem 9

We now prove Theorem 9, for all the joinable trees and all three set algorithms (union, intersection, difference) from Figure 13.

For this purpose, we make two observations. The first is that all the work for the algorithms can be accounted for within a constant factor by considering just the work done by the *splits* and the *joins* (or $join2$s), which we refer to as *split work* and *join work*, respectively. This is because the work done between each split and join is constant. The second observation is that the split work is identical among the three set algorithms. This is because the control flow of the three algorithms is the same on the way down the recursion when doing *splits*—the algorithms only differ in what they do at the base case and on the way up the recursion when they join.

Given these two observations, we prove the bounds on work by first showing that the join work is asymptotically at most as large as the split work (by showing that this is true at every node of the recursion for all three algorithms), and then showing that the split work for union (and hence, the others) satisfies our claimed bounds.

We start with some notation, which is summarized in Table 3. In the three algorithms, the first tree ($T_1$) is split by the keys in the second tree ($T_2$). We therefore call the first tree the *decomposed tree* and the second the *pivot tree*, denoted as $T_d$ and $T_p$, respectively. The tree that is returned is denoted as $T$. Since our proof works for either tree being larger, we use $m = \min(|T_p|, |T_d|)$ and $n = \max(|T_p|, |T_d|)$. We denote the subtree rooted at $v \in T_p$ as $T_p(v)$, and the tree of keys from $T_d$ $^9$The nodes in $T_d(v)$ form a subset of $T_d$, but not necessarily a subtree. See details later.
that \( v \) splits as \( T_d(v) \) (i.e., split \((v, T_d(v))\)) is called at some point in the algorithm. For \( v \in T_P \), we refer to \( |T_d(v)| \) as its splitting size.

Figure 14(a) illustrates the pivot tree with the splitting size annotated on each node. Since \textit{split} has logarithmic work, we have

\[
\text{split work} = O\left(\sum_{v \in T_P} \left(\log |T_d(v)| + 1\right)\right),
\]

which we analyze in Theorem 11. We first, however, show that the join work is bounded by the split work. We use the following Lemma.

**Lemma 9.** For \( T = \text{union}(T_P, T_d) \) on strongly joinable trees, then \( \max(\text{rank}(T_P), \text{rank}(T_d)) \leq \text{rank}(T) \leq \text{rank}(T_P) + \text{rank}(T_d) \).

**Proof.** We prove it by induction on the tree size. For small trees, this conclusion obviously holds. Note that \( T_d \) will be split up into two trees \( T_I \) and \( T_r \), with rank at most \( r = \text{rank}(T_d) \) (Theorem 7). \( lc(T_p) \) and \( rc(T_p) \) will take \text{union} with either \( T_I \) or \( T_r \), i.e., \( L = \text{union}(lc(T_P), T_I) \) and \( R = \text{union}(rc(T_P), T_I) \). Because of the induction hypothesis, \( 0 \leq \text{rank}(L) \leq \text{rank}(lc(T_P)) + \text{rank}(T_I) \leq \text{rank}(lc(T_P)) + r \), and similarly \( 0 \leq \text{rank}(R) \leq \text{rank}(lc(T_P)) + r \). From the submodularity rule, joining them increases the rank of \( T_P \) by at least 0 and at most \( \text{rank}(T_d) \).

\( \square \)

**Theorem 10.** For each function call to \text{union}, intersection, or difference on strongly joinable trees and treaps \( T_P(v) \) and \( T_d(v) \), the work to do the join (or join2) is asymptotically no more than the work to do the split.

**Proof.** For \text{intersection} or \text{difference}, the cost of \text{join} (or \text{join2}) is \( O(\log(|T|)) \), where \( T \) is the result tree. Notice that \text{difference} returns the keys in \( T_d \setminus T_P \). Thus, for both \text{intersection} and \text{difference} we have \( T \subseteq T_d \). The join work is \( O(\log(|T_d|)) \), which is no more than \( O(\log(|T_d|)) \) (the split work).

For \text{union}, if \( \text{rank}(T_P) \leq \text{rank}(T_d) \), the join will cost \( O(\text{rank}(T_d)) \), which is no more than the split work.

Consider \( \text{rank}(T_P) > \text{rank}(T_d) \) for strongly joinable trees. The rank of \( lc(T_P) \) and \( rc(T_P) \), which are used in the recursive calls, is at least \( \text{rank}(T_P) - c_u \). Using Lemma 9, the rank of the two trees returned by the two recursive calls will be at least \( \text{rank}(T_P) - c_u \) and at most \( \text{rank}(T_P) + \text{rank}(T_d) \), and differ by at most \( O(\text{rank}(T_d)) = O(\log |T_d|) \). Thus, the join cost is \( O(\log |T_d|) \), which is asymptotically no more than the split work.

Consider \( \text{rank}(T_P) > \text{rank}(T_d) \) for treaps. If \( \text{rank}(T_P) > \text{rank}(T_d) \), then \( |T_P| \geq |T_d| \). The root of \( T_P \) has the highest priority among all \( |T_P| \) keys, so on expectation it takes at most the \( \frac{|T_P| + |T_d|}{|T_P|} \leq 2 \) th place among all the \( |T_d| + |T_P| \) keys. From Lemma 6, we know that the cost on expectation is \( E[\log t] + 1 \leq \log E[t] + 1 \leq \log 2 + 1 \), which is a constant.

\( \square \)

This implies the total join work is asymptotically bounded by the split work.

We now analyze the split work. We do this by layering the pivot tree starting at the leaves and going to the root and such that nodes in a layer are not ancestors of each other. We use the definition in Section 3. We define layers based on the ranks and denote the number of \text{rank}(i)-root nodes as \( s_i \). Lemma 1 shows that \( s_i \) shrinks geometrically for joinable trees, which helps us prove our bound on the split work. Figure 14(b) shows an example of the layers of an AVL tree on the two input trees of the \text{join}-based set functions.

**Lemma 10.** For any ancestor-free set \( V \subseteq T_P \), \( \sum_{v \in V} |T_d(v)| \leq |T_d| \).

\(^{10}\text{When } |T_d(v)| = 0, \text{ we set } log|T_d(v)| \text{ as } 0.\)
Fig. 14. An illustration of splitting tree and layers. The tree in (a) is $T_p$, and the dashed circles are the exterior nodes. The numbers on the nodes are the sizes of the tree from $T_d$ to be split by this node, i.e., the “splitting size” $|T_d(v)|$. (b) is an illustration of layers on an AVL tree.

The proof of this lemma is straightforward.

Not all nodes are rank roots. However, Property 7 shows that each rank cluster attached to a rank root contains only a constant number of nodes, and they are all the rank root’s descendants.

By applying Lemma 1 and Property 7, we prove the split work. In the following proof, we denote $v_{kj}$ as the $j$-th node in layer $k$.

**Theorem 11.** The split work in union, intersection, and difference on two joinable trees of size $m$ and $n$ is $O(m \log(n/m + 1))$.

**Proof.** The total work of split is the sum of the log of all the splitting sizes on the pivot tree $O(\sum_{v \in T_p} \log(|T_d(v)| + 1))$. Denote $l$ as the number of layers in the tree. Also, notice that in the pivot tree, in each layer there are at most $|T_d|$ nodes with $|T_d(v_{kj})| > 0$. Since those nodes with splitting sizes of 0 will not cost any work, we can assume $s_j \leq |T_d|$. We calculate the dominant term $\sum_{v \in T_p} \log(|T_d(v)| + 1)$ in the complexity by summing the work across layers. We first only consider all rank roots.

$$
\sum_{k=0}^{l} \sum_{j=1}^{s_k} \log \left( |T_d(v_{kj})| + 1 \right) \leq \sum_{k=0}^{l} s_k \log \left( \frac{\sum_{j=1}^{s_k} |T_d(v_{kj})| + 1}{s_k} \right) \\
= \sum_{k=0}^{l} s_k \log \left( \frac{|T_d| + 1}{s_k} \right).
$$
We split it into two cases. If \( |T_d| \geq |T_p| \), \( \frac{|T_d|}{s_k} \) always dominates 1. We have

\[
\sum_{k=0}^{l} s_k \log \left( \frac{|T_d|}{s_k} + 1 \right) = \sum_{k=0}^{l} s_k \log \left( \frac{n}{s_k} + 1 \right) \tag{4}
\]

\[
\leq \sum_{k=0}^{l} \frac{m}{2^{\lfloor k/c \rfloor}} \log \left( \frac{n}{m/2^{\lfloor k/c \rfloor}} + 1 \right) \tag{5}
\]

\[
\leq c \sum_{k=0}^{\lfloor l/c \rfloor} \frac{m}{2^k} \log \frac{n}{m/2^k} + c \sum_{k=0}^{\lfloor l/c \rfloor} \frac{m}{2^k} \log 2^k
\]

\[
= O \left( m \log \frac{n}{m} \right) + O(m)
\]

\[
= O \left( m \log \left( \frac{n}{m} + 1 \right) \right). \tag{6}
\]

If \( |T_d| < |T_p| \), \( \frac{|T_d|}{s_k} \) can be less than 1 when \( k \) is smaller, thus the sum should be divided into two parts. Also note that we only sum over the nodes with splitting size larger than 0. Even though there could be more than \( |T_d| \) nodes in one layer in \( T_p \), only \( |T_d| \) of them should count. Thus, we assume \( s_k \leq |T_d| \), and we have

\[
\sum_{k=0}^{l} s_k \log \left( \frac{|T_d|}{s_k} + 1 \right) = \sum_{k=0}^{l} s_k \log \left( \frac{m}{s_k} + 1 \right) \tag{7}
\]

\[
\leq 2^{\log_2 \frac{n}{m}} \sum_{k=0}^{\log_2 \frac{n}{m}} |T_d| \log (1 + 1)
\]

\[
+ \sum_{k=\lceil c \log_2 \frac{n}{m} \rceil}^{l} \frac{n}{2^{\lfloor k/c \rfloor}} \log \left( \frac{m}{n/2^{\lfloor k/c \rfloor}} + 1 \right) \tag{8}
\]

\[
\leq O \left( m \log \frac{n}{m} \right) + c \sum_{k'=0}^{\left\lfloor \frac{l}{c} \right\rfloor} \frac{m}{2^{k'}} \log 2^{k'}
\]

\[
= O \left( m \log \frac{n}{m} \right) + O(m)
\]

\[
= O \left( m \log \left( \frac{n}{m} + 1 \right) \right). \tag{9}
\]

From (4) to (5) and (7) to (8), we apply Lemmas 1 and 8 and the fact that \( f(x) = x \log \left( \frac{x}{e} + 1 \right) \) is monotonically increasing when \( x \leq n \).

The above cost does not consider the nodes that are not rank roots. Recall that we use \( d(v) \) to denote the rank cluster of a rank root \( v \). Applying Property 7, the total cost (in expectation for
treaps) is less than

\[
\sum_{k=0}^{l} \sum_{j=1}^{x_k} d(v_{kj}) \log((T_d(v_{kj}) + 1)
\]

\[
= d(v_{kj}) \times 2 \sum_{k=0}^{l} \sum_{j=1}^{x_k} \log((T_d(v_{kj}) + 1)
\]

\[
= O \left( m \log \left( \frac{n}{m} + 1 \right) \right).
\]

Therefore, the split work is \(O(m \log (\frac{n}{m} + 1))\) in all cases discussed in the theorem. \(\square\)

**Theorem 12.** The total work of union, intersection, or difference of all four balancing schemes on two trees of size \(m\) and \(n\) \((m \geq n)\) is \(O(m \log (\frac{n}{m} + 1))\).

This directly follows Theorems 10 and 11.

**Theorem 13.** The span of union and intersection or difference on all four balancing schemes is \(O(\log n \log m)\). Here, \(n\) and \(m\) are the sizes of the two trees.

Proof. For the span of these algorithms, we denote \(D(h_1, h_2)\) as the span on union, intersection, or difference on two trees of height \(h_1\) and \(h_2\). According to Theorem 10, the work (span) of split and join are both \(O(\log |T_d|) = O(h(T_d))\). We have

\[
D(h(T_p), h(T_d)) \leq D(h(T_p) - 1, h(T_d)) + 2h(T_d)
\]

Thus, \(D(h(T_p), h(T_d)) \leq 2h(T_p)h(T_d) = O(\log n \log m)\). \(\square\)

Combining Theorems 12 and 13, we come to Theorem 9.

### 5.4 Other Tree Algorithms Using join

**Insert and Delete.** Instead of the classic implementations of insert and delete, which are specific to the balancing scheme, we define versions based purely on join, and hence independent of the balancing scheme.

We present the pseudocode in Figure 15 to insert an entry \(e\) into a tree \(T\). The base case is when \(t\) is empty, and the algorithm creates a new node for \(e\). Otherwise, this algorithm compares \(k\) with the key at the root and recursively inserts \(e\) into the left or right subtree. After that, the two subtrees are joined again using the root node. Because of the correctness of the join algorithm, even if there is imbalance, join will resolve the issue.

The delete algorithm is similar to insert, except when the key to be deleted is found at the root, where delete uses join2 to connect the two subtrees instead. Both the insert and the delete algorithms run in \(O(\log n)\) work (and span since sequential).

One might expect that abstracting insertion or deletion using join instead of specializing for a particular balance criteria has significant overhead. Our experiments show this is not the case—and even though we maintain the reference counter for persistence, we are only 17% slower sequentially than the highly optimized C++ STL library (see Section 6).

**Theorem 14.** For a joinable tree \(T\) with weight \(n = |T|\), the join-based insertion algorithm does \(O(\log n)\) work. The rank of the output tree is at most \(c_u + \text{rank}(T)\).

Proof. We prove this by induction. This is obviously true for the base case. Consider \(T = \text{node}(L, t, R)\), assume \(e\) goes to the left subtree \(L\), and the new left subtree is \(L'\). Then the output
Fig. 15. Pseudocode of some join-based functions. They are all independent of balancing schemes. The syntax $S_1||S_2$ means that the two statements $S_1$ and $S_2$ can run in parallel based on fork-join parallelism.
tree is $T' = join(L', t, R)$. Because of the induction hypothesis $\text{rank}(L') \leq \text{rank}(L) + c_u$. Because of the submodularity rule, $\text{rank}(T') \leq \text{rank}(T) + c_u$. □

Theorem 15. For a joinable tree $T$ with weight $n = |T|$, the join-based deletion algorithm does $O(\log n)$ work. The rank of the output tree is at most $\text{rank}(T)$.

Proof. We first prove this by induction. This is obviously true for the base case. Consider $T = \text{node}(L, t, R)$, assume $k$ falls in the right subtree $R$, and the new left subtree is $R'$. Then the output tree is $T' = join(L', t, R')$. From the induction hypothesis, $\text{rank}(R') \leq \text{rank}(R)$. Based on the decreasing side of the submodularity rule, $\text{rank}(T') \leq \text{rank}(T)$. This proves that the output tree of delete is at most the rank of the input.

In delete, there is at most one invocation of join2, which takes time no more than $\log n$. For $T' = join(L, t, R')$, we next prove the cost. Consider two cases.

1. The key $k \neq k(R)$. WLOG assume $k$ falls in the right subtree of $R$ (the other case is symmetric). $R' = join(lc(R), R, t, rc(R), k)$. Then

$$\text{rank}(R') \geq \text{rank}(lc(R)) \geq \text{rank}(R) - c_u \geq \text{rank}(L) - 2c_u + c_l.$$ 

Meanwhile, $\text{rank}(R') \leq \text{rank}(R)$ as we have proved above. Therefore,

$$\text{rank}(R') \leq \text{rank}(R) \leq \text{rank}(L) + c_u - c_l.$$ 

In summary, $\text{rank}(L)$ and $\text{rank}(R')$ differ by a constant. The cost of a single join is a constant.

2. The key $k = k(R)$. $R' = join2(lc(R), rc(R))$. The cost of this join is at most $O(\log n)$, but this only happens once.

In summary, the total cost of all join is $h(T) + O(\log n) = O(\log n)$, and the cost of the join2 is at most $O(\log n)$. This proves the above theorem. □

Build. A balanced binary tree can be created from a sorted array of key-value pairs using a balanced divide-and-conquer over the input array and combining with join. To construct a balanced binary tree from an arbitrary array, we first sort the array by the keys, then remove the duplicates. All entries with the same key are consecutive after sorting, so the algorithm first applies a parallel sorting and then follows by a parallel packing. The algorithm then extracts the median in the de-duplicated array, and recursively constructs the left/right subtree from the left/right part of the array, respectively. Finally, the algorithm uses join to connect the median and the two subtrees. The work is then $O(W_{\text{Sort}}(n) + W_{\text{Remove}}(n) + n)$ and the span is $O(S_{\text{Sort}}(n) + S_{\text{Remove}}(n) + \log n)$. For work-efficient sort and remove-duplicates algorithms with $O(\log n)$ span this gives the bounds in Table 2.

Bulk Updates. We use multi_insert and multi_delete to commit a batch of write operations. The function multi_insert($T, A, m$) takes as input a P-Tree root $t$, and the head pointer of an array $A$ with its length $m$.

We present the pseudocode of multi_insert in Figure 15. This algorithm first sorts $A$ by keys, and then removes duplicates in a similar way as in build. We then use a divide-and-conquer algorithm multi_insert_s to insert the sorted array into the tree. The base case is when either the array $A$ or $T$ is empty. Otherwise, the algorithm uses a binary search to locate $t$’s key in the array, getting the corresponding index $b$ in $A$. $d$ is a bit denoting if $k$ appears in $A$. Then, the algorithm recursively multi-inserts $A$’s left part (up to $A[b]$) into the left subtree, and $A$’s right part into the right subtree. The two recursive calls can run in parallel. The algorithm finally concatenates the two results by the root of $T$. A similar divide-and-conquer algorithm can be used for multi_delete, using join2 instead of join when necessary.
Decoupling sorting from inserting has several benefits. First, parallel sorting is well studied and there exist highly optimized sorting algorithms that can be used. This simplifies the problem. Second, after sorting, all entries in $A$ to be merged with a certain subtree in $T$ become consecutive. This enables the divide-and-conquer approach which provides good parallelism, and also gives better locality.

The total work and span of inserting or deletion of a sorted array of length $m$ into a tree of size $n \geq m$ is $O(m \log (\frac{m}{n} + 1))$ and $O(\log m \log n)$, respectively [14]. The analysis is similar to the union algorithm.

**Range.** $\text{range}(T, k_L, k_R)$ extracts a subset of tuples from $T$ in a key range $[k_L, k_R]$, and outputs them in a new P-Tree. The cost of the $\text{range}$ function is $O(\log n)$. The pure $\text{range}$ algorithm copies nodes on two paths, one to each end of the range, and uses them as pivots to $\text{join}$ the subtrees back. When the extracted range is large, this pure $\text{range}$ algorithm is much more efficient (logarithmic time) than visiting the whole range and copying it.

**Filter.** The $\text{filter}(T, \phi)$ function returns a tree with all tuples in $T$ satisfying a predicate $\phi$. This algorithm filters the two subtrees recursively, in parallel, and then determines if the root satisfies $\phi$. If so, the algorithm uses the root as the pivot to $\text{join}$ the two recursive results. Otherwise, it calls $\text{join2}$. The work of $\text{filter}$ is $O(n)$ and the depth is $O(\log^2 n)$ where $n$ is the tree size.

**Map and Reduce.** The function $\text{map\_reduce}(T, f_m, \langle f_r, I \rangle)$ on a tree $t$ (with data type $E$ for the tuples) takes three arguments and returns a value of type $V^\prime$. $f_m : E \mapsto V^\prime$ is the a map function that converts each stored tuple to a value of type $V^\prime$. $\langle f_r, I \rangle$ is a monoid where $f_r : V^\prime \times V^\prime \mapsto V^\prime$ is an associative reduce function on $V^\prime$, and $I \in V^\prime$ is the identity of $f_r$. The algorithm will recursively call the function on its two subtrees in parallel, and reduce the results by $f_r$ afterwards.

## 6 EXPERIMENTS

To evaluate the performance of our algorithms, we performed several experiments across the four balancing schemes using different set functions, while varying the core count and tree sizes. We also compare the performance of our implementation to other existing libraries and algorithms.

In this article, we mainly show the performance for the set operations. For other $\text{join}$-based algorithms, some performance evaluations are available in [11, 60–62].

**Implementation Details.** We implemented all the algorithms in the PAM (Parallel Augmented Map) library [59, 62]. PAM is a C++ library supporting normal ordered maps (key-value store) and augmented maps (as defined in [62]) based on balanced binary trees. All four discussed balancing schemes are supported in PAM. The algorithms in PAM are based on the $\text{join}$-based algorithms, which matches the versions described in this article. Our code is available online [59]. We refer to the trees implemented in PAM as P-Trees [61].

As a general-purpose library, PAM also supports persistence (and thus multi-versioning) based on path-copying, and reference count garbage collection. A persistent (or purely functional) update will not modify the input tree, but generate a new version with the update as the output. To perform an update operation using path-copying, the algorithm will copy all tree nodes on the affected path, and the new root pointer represents the output tree. We refer the readers to [61, 62] for more details. This means that a tree node can be shared among multiple versions, and thus we use a reference counter for each node to aid memory management. PAM also allows for in-place updates. In our experiments, since the two implementations that we are comparing with are not persistent, we test the in-place version of the algorithms in PAM.

Each tree node in P-tree contains the key, the value, two child pointers ($2 \times 8 = 16$ bytes), the balance information (4 bytes if needed; see details below), the size of the subtree (4 bytes), and a
reference count (4 bytes, used for garbage collection; see details above). The balancing information is only needed for AVL trees (the height) and RB trees (the black height and the color). For both cases, we use a 4-byte integer to store them (for RB tree we encoded the color as a bit in the black height). For treaps, the priority of each node is generated by a user-specified hash function on the key. For WB trees, the weight can be directly computed by one plus the size. This means that for a key-value pair with 8-byte key and 8-byte value (which is the setting we used in our experiments), each node is 40 bytes (for WB trees and treaps) or 44 bytes (for AVL and RB trees). As discussed below, experiments show that WB trees generally show the best performance among all the balancing schemes we tested. We believe the reason is exactly because WB tree does not store extra balancing information, and generally has lower height than treaps. Therefore, the default interface of PAM is using WB trees.

We note that some metadata stored in the tree nodes, such as the size and the reference counter, are not necessary for implementing set operations. However, they are useful in supporting more complete functionalities in the ordered map interface. For example, the size of each subtree is useful in answering queries such as selecting the k-th element, or answering the rank of an entry in the ordered map. The reference counter is useful in memory management for persistent updates. This information is not maintained in the other implementations that we compare to. In Section 7, we will present some discussions about more space-efficient search tree structures.

**Experiment Setups and Baseline Algorithms.** For all the experiments, we use a 72-core Dell R930 with 4x Intel(R) Xeon(R) E7-8867 v4 (18 cores, 2.4 GHz, and 45 MB L3 cache), and 1 Tbyte memory. Each core is two-way hyperthreaded giving 144 hyperthreads. Our code was compiled using g++ 5.4.1. In the semantics, we only use the binary fork-join, and parallel for loop. We compile with -O2 because this gives us more stable results. We use numactl -i all in all experiments with more than one thread. It evenly spreads the memory pages across the processors in a round-robin fashion.

In all our experiments, we use keys and values of the 64-bit integer data type. Throughout this section, \( n_1 \) and \( n_2 \) represent the sizes of the two input sets. We generate multiple sets varying in size up to \( 10^8 \). For most of the experiments (except for Figure 17), the keys are 64-bit integers drawn from a uniform distribution. As mentioned, some experiments using different input distributions, and real-world data distributions (e.g., Yahoo! Cloud Serving Benchmark (YCSB) [28] using Zipfian distribution), have been shown in some other papers [14, 61]. We note that uniform distribution also means that the two input trees are fully interleaving with each other, which is the worst case for our algorithms. Because our algorithms are comparison-based, the actual key distribution does not affect the running time, but how the two sets interleaving with each other matters. To better understand this, in Figure 17 we will also show when the two trees have different overlapping ratio in key ranges. Our experiments show that when the two sets partially overlap with each other, our algorithms can exhibit better performance.

We test our algorithm by comparing it to other available implementations. This includes the sequential version of the set functions defined in the C++ Standard Template Library (STL) [50], which we use for std::vector, and STL’s std::set (implemented by RB tree). To see how well our algorithm performs in a parallel setting, we compare it to parallel WBB-trees [31] and the MCSTL library [33], both supporting array-tree union. For array-tree union, we use the tree to denote the first set (size \( n_1 \)) and the array as the second set (size \( n_2 \)). For MCSTL and WBB-trees, the entry stored in each tree node is just a key. For STL-map, STL-set, and our implementation based on PAM, the entries are key-value pairs.

We note that more experiments on the join-based algorithms are also available in some other papers, e.g., on different input distributions [14], comparing with concurrent data structures.
Joinable Parallel Balanced Binary Trees, and comparing with existing geometric libraries [60]. The PAM library has also been tested on real-world database benchmark and micro-benchmarks, including YCSB [28] and TPC benchmarks [1] (see more details in [61]).

**Comparing Different Balancing Schemes and Set Operations.** To compare the four balancing schemes, we choose union as the representative operation. Other operations give similar results. We compare the schemes across varying input sizes. The two input sets are of the same size varying from $10^5$ to $10^8$. Figure 16(a) shows the runtime of union for varying tree sizes and all four balancing schemes on 72 cores (144 threads). The overall trend and running times are similar across the balancing schemes. Generally speaking, WB trees have the best performance among the four tested balancing schemes. This is because the other three balancing schemes also need to store an extra field as the balancing information (e.g., the height for AVL trees). For WB trees, the balancing criteria is the size of each subtree, which is maintained anyway for answering other queries (e.g., quickly selecting the $i$-th element in the set). It is perhaps not surprising that all balancing schemes achieve similar performance because the dominant cost is in cache misses along the paths in the tree, and all schemes keep the trees reasonably balanced. Treaps have the worst performance among the four, likely due to the larger height of the tree. The difference in running times between treaps (slowest) and WB trees (fastest) can be up to 70%. However, for AVL and RB trees, the difference from WB trees is usually within 30%. For this reason, the PAM library uses the WB trees as the default underlying balancing scheme.

We will use the WB tree as the representative tree in the following experiments. We first show tests to compare different set operations. The two input sets are of the same size varying from $10^5$ to $10^8$. To make intersection and difference have reasonable outputs (instead of outputting an empty set for most of the time), we select at least a half of the elements in the second set from the first set. Figure 16(b) compares time for the union, intersection, and difference functions. The three functions have very similar performance. For intersection and difference, the time also includes possible cost in collecting nodes that are not in the output. For this reason, intersection and difference are slightly more expensive than union. We note that by using a persistent version of these algorithms, one can avoid the extra cost of garbage collection. This is out of the scope of this article, so we omit the details.

**Comparing to Existing Parallel Implementations.** To see how well our algorithm performs in the parallel setting, we compare it to parallel WBB-trees [31] and the MCSTL library [33]. WBB-trees, as well as the MCSTL, offer an interface for bulk insertions and deletions, which takes a tree and a sorted array, and inserts (or deletes) the elements in the array into (from) the tree. Our version is symmetric in taking both input sets as trees. If the size of the array is smaller than the tree, array-tree unions have an inherent advantage over tree-tree unions since accessing an array is much more cache efficient than accessing an tree. As shown in Figure 16(c), we set the first set size ($n_1$) as $10^8$ and vary the other set size ($n_2$) from $10^5$ to $10^8$. In this case, the smaller set will be represented as the input array in WBB-trees and MCSTL. In this case, WBB-trees have better performance than our implementation when $n_2$ is small. This is probably because WBB-tree itself has a more cache-aware layout (eight keys per cache line as opposed to one), leading to a better cache utilization compared to both the MCSTL and our implementation. When $n_2$ gets larger, our implementation achieves similar performance to WBB-tree, and is much faster than MCSTL.

In Figure 16(d), we fix the second set size $n_2$ as $10^8$, which is the array in the input of WBB-tree and MCSTL. We vary the first set size ($n_1$) from $10^5$ to $10^8$. In this case, the running time of both WBB-tree and MCSTL becomes (almost) flat. In other words, lowering the second set size does not reduce the running time for either WBB-tree or MCSTL, and the running time is proportional.
Fig. 16. Experimental results on join-based algorithms.

(a) Union, 144 threads  
Various balancing schemes  
\(n_1 = n_2\) shown on x-axis

(b) WB Trees, 144 threads  
Various set operations  
\(n_1 = n_2\) shown on x-axis

(c) WB Trees, union, 144 threads  
Comparing to MCSTL and WBB-trees  
\(n_1 = 10^8\), varying \(n_2\) as shown on x-axis

(d) WB Trees, union, 144 threads  
Comparing to MCSTL and WBB-trees  
\(n_2 = 10^8\), varying \(n_1\) as shown on x-axis

(e) WB Trees, union, \(n_1 = n_2 = 10^8\)  
Comparing to MCSTL and WBB-trees  
Varying number of threads as shown on x-axis

(f) WB Trees, union, 1 thread (sequential)  
Comparing to STL-map and STL-set  
\(n_1 = 10^8\), varying \(n_2\) as shown on x-axis

Scalability Tests. We also compare the parallelism of these implementations. In Figure 16(e), we show their performance across 72 cores (144 threads). The inputs are both of size \(10^8\), and generated from a uniform distribution of integers. The trend shows that all three tested implementations to \(n_2\). In our symmetric version, the order of the two input sizes does not make a difference. The curves of P-Trees in Figure 16(c) and (d) show a similar trend.
have good speedup up to 72 threads. Our version gets some extra speedup when hyperthreading is involved. With small number of threads, WBB-trees are slightly faster than our code, but when it comes to all 144 threads, P-Trees are slightly faster than WBB-trees. This indicates that our algorithm achieves better parallelism. On 144 threads, P-Trees get a 73-fold self-speedup, while MCSTL and WBB-trees have self-speedup numbers as 17 and 20, respectively. Because the sequential version of WBB-trees is about 3.5× faster than our implementation, the parallel performances using all 144 threads of P-Trees and WBB-trees are close to each other.

To conclude, in terms of parallel performance, our code and WBB-trees are always much better than MCSTL because of MCSTL’s slow sequential performance (indicating that it may have more expensive work). WBB-trees achieve a slightly better performance than ours when fewer threads are used, but when using all available threads, WBB-trees achieve similar performance to ours, indicating that our implementation explores better parallelism. Also, when the second input set size is small, neither MCSTL nor WBB-trees can take the advantage, while the running time of our implementation is always proportional to the small set size.

Comparing to Sequential Implementations. The STL supports set_union, set_intersection, and set_difference on any container class, including sets based on RB trees, and sorted vectors (arrays). Since the STL does not offer any parallel version of these functions, we could only use it for sequential experiments. For two inputs of size \( n_1 \) and \( n_2 \), set_union takes \( O(n_1 + n_2) \) time on std::vectors by using the standard merging algorithm which uses two pointers moving from left to right on the two inputs, comparing the current values, and writing the lesser to the end of the output. This is also the merge algorithm in a standard merge sort. For std::set that uses RB trees, we can insert elements from the smaller set into the larger, leading to a time of \( O(\min(n_1, n_2) \log(n_1 + n_2)) \). These two sequential algorithms are noted as STL-tree and STL-vector in our tests.

Figure 16(f) gives a comparison of times for union. For equal lengths, our implementation is about a factor of 7× faster than STL-tree, and about 5× slower than STL-vector. This is not surprising since we are asymptotically faster than STL-tree. Meanwhile, for STL-vector, the theoretical cost \( O(n_1 + n_2) \) is asymptotically the same as ours \( O(n_2 \log(n_1/n_2 + 1)) \) when \( n_1 = n_2 \). However, STL-vector’s array-based implementation just reads and writes the values, one by one, from flat arrays, and therefore has much less overhead and much fewer cache misses than ours. We note that using trees as the representation of ordered sets has the benefit of allowing for fast insertions and deletions, which is not supported by array-based implementations. For taking the union of smaller and larger inputs, our union is orders of magnitude faster than either STL version. This is because their theoretical work bound \( O(n_1 + n_2) \) and \( O(n_2 \log(n_1 + n_2)) \) is worse than our \( O(n_2 \log(n_1/n_2 + 1)) \), which is optimal in the comparison model.

Testing Different Key Range Overlapping. In all previous experiments, we use two sets with fully interleaved key ranges. As mentioned, this is actually the worst case for our algorithms. We also tested two sets of different key range overlapping, and show the running time in Figure 17(a) and (b). In Figure 17(a), both sets have a key range of \( 10^9 \), but they have partially overlapped key range. The first set will always have keys in range \([0, 10^5]\), and the second set will have keys in key range \([i \times 10^8, i \times 10^8 + i \times 10^5]\) for test \( i \). We vary \( i \) from 0 to 10, which means the key range overlapping of the two sets varies from 0% (fully non-overlapping) to 100% (fully interleaving). As shown in Figure 17, as the key range overlapping ratio decreases, the total running time also decreases linearly. This is not surprising because when the overlapping range is small, it means that in the divide-and-conquer algorithm, the two subproblems usually have unbalanced size, and some of the subproblems quickly reach the base case. When the two key ranges do not overlap with
each other, the union algorithm degenerates to a join2 algorithm, which has work \( O(\log^2 n) \). This matches our experimental result, where when the overlapping is 0%, the running time is smaller than 1 ms.

We also tested when the key range of a set is a subset of the other key range. In particular, in Figure 17(b), we always set the key range of the first set to be \([0, 10^9]\), and the other key range is set to be \([0, 2i \times 10^8]\) for \(i = 1\) to \(5\). This means the key range of \(T_2\) is always contained in \(T_1\)'s key range, and the overlapping changes from 20% to 100%. Similar to Figure 17(a), we see that when the overlapping ratio decreases, the performance also improves in an almost linear manner. This is also because in many subproblems, one of the tree sizes hits 0 quickly. As a result, the total work performed by the algorithm is much smaller than normal (i.e., when they fully interleave with each other).

7 RELATED WORK

Join-Based Algorithms. Tarjan first studied the join and split functions in [64]. Tarjan showed how to efficiently implement the functions on RB and splay trees but did not give any applications for the two functions. Adams [2, 3] applied join and split on WB trees to some bulk functions and showed an elegant way to implement union, intersection, and difference. The method was then implemented in some languages and libraries such as MIT/GNU Scheme in Haskell [48]. Adams’ paper was reported buggy [36, 58] in parameter choosing and also in that the join (which is called concat3 in his paper) function does not rebalance the tree, but the framework of set functions implementation is still used and studied today. Adams did not consider parallelism in the algorithm, but the divide-and-conquer scheme is inherently parallel.

join and split appear in the LEDA library [49] for sorted sequences, and the CGAL library for ordered maps [67]. None of this work considered parallel algorithms based on the functions, or how to build an interface out of just join. Frias and Singler [33] use join and split on RB trees for an implementation of the MCSTL, a multi-core version of the C++ STL. Their algorithms are lower level based on partitioning across processors, and are for bulk insertion and deletion. The functions join, split and join2 are also studied and used in various previous work for trees to support multiple applications [12, 27, 56].

Set Operations. Merging two ordered sets has been well studied in the sequential setting. Hwang and Lin [37] describe an algorithm to merge two arrays, which costs optimal work. Their algorithm
works for arrays, and since writing back both array costs $O(m + n)$ work, the algorithm only returns the cross pointers between two arrays. Brown and Tarjan [22] considered input data to be arranged in a BST, which allows the merged result to be explicitly given by a new BST in time $O(m \log (n + 1))$. Their algorithm works on AVL and 2-3 trees. None of these algorithms considered parallelism. Katajainen et al. [42] studied the space efficiency on merging two sets in parallel. Their focus is not on reducing time complexity. Furthermore, the above-mentioned works are not based on join and are much more complicated than our algorithm.

There is also previous work studying parallel set operations on two ordered sets, but each previous algorithm only works on one type of balance tree. Paul et al. studied bulk insertion and deletion on 2-3 trees in the PRAM model [55]. Park and Park showed similar results for RB trees [54]. These algorithms are not based on join and are not work-efficient, requiring $O(m \log n)$ work. Katajainen [41] claimed an algorithm with $O(m \log (n + 1))$ work and $O(\log n)$ span using 2-3 trees, but it appears to contain some bugs in the analysis [16]. Blelloch and Reid-Miller described a similar algorithm as Adams’ (as well as ours) on treaps with optimal work (in expectation) and $O(\log n)$ span (with high probability) on an EREW PRAM with scan operations. This implies $O(\log n \log m)$ span on a plain EREW PRAM, and $O(\log n \log^* m)$ span on a plain CRCW PRAM. The pipelining that is used is quite complicated. Akhremtsev and Sanders [7] describe an algorithm for array-tree join based on $(a, b)$-trees with optimal work and $O(\log n)$ span on a CRCW PRAM. Our focus in this article is in showing that very simple algorithms are work efficient, have polylogarithmic span, and generic for multiple balancing schemes, and less with optimizing the span. We note that their algorithms use multi-way search tree instead of binary, which potentially would have better I/O efficiency than the binary trees. We present some discussions about this at the end of this section. There has also been work about batched updates on certain balancing schemes [5, 6].

**Set Algorithms in the Concurrent and Distributed Setting.** Many researchers have considered concurrent implementations of balanced search trees (e.g., [21, 43, 44, 51]). None of these are work efficient for union since it is necessary to insert one tree into the other requiring at least $\Omega(m \log n)$ work. Researchers have also studied distributed memory implementations of maps and sets, including a distributed version of STL as part of the HPC++ effort [39], and the STAPL library [63]. The emphasis of this work is on how the maps and sets are partitioned across the memories.

**Discussion about Other Balancing Schemes.** This article shows that the join-based algorithm framework applies to four balancing schemes. The authors believe that similar ideas apply to other balancing schemes. For example, a recent paper [35] proposed the WAVL tree, which uses a similar notion of rank as our work, but only allows integer ranks. Their methodology applies to balanced binary trees where the height of the two siblings can differ by at most 2. They also showed the equivalence of WAVL tree to an RB tree. As a result, our join-based framework also applies to WAVL trees. This is another recent paper on zip trees [65], which is a randomized balanced tree structure. The join algorithms for treaps and all join-based algorithms work correctly on zip trees. However, zip trees does not directly fit in the definition of weakly joinable trees, and thus the bounds and analysis does not directly apply to zip trees.

**Discussion about Concurrent Tree Structures.** There is previous work focusing on supporting concurrent operations on BSTs [23, 26, 45–47, 53, 66]. Some of this previous work aims at making such operations lock-free or wait-free [9, 21, 24, 32, 68]. The supported operations usually include insertion, deletion, update, lookup, range queries, and sometimes more complicated queries. Instead of focusing on supporting concurrent operations, P-Trees provide bulk functions (e.g., union and multi_insert) to commit a batch of operations in parallel. P-Trees and the PAM library are also safe for concurrency, but concurrency is supported by making P-Trees persistent by path-copying...
This requires all concurrent threads to work on a snapshot of the tree. To allow for serializable concurrent operations on P-Trees, one can batch all update operations and commit them using the parallel bulk functions. Since this is not the main focus of this article, we omit the details, and more information can be found in \[11, 61\].

**Discussion about I/O- and Space-Efficient Search Tree Structures.** One concern of using binary trees is the space overhead and I/O efficiency. In particular, each data entry is stored in a tree node, which needs to maintain the pointers, size, reference count, and so on. For this reason, there are also researchers attempting to design algorithms for B-tree or B-tree-like data structures.

Akhremtsev and Sanders \[7\] described an algorithm for array-tree set algorithms based on \((a, b)\)-trees. Their algorithms are also based on the (multi-)split-join framework, and are work efficient and highly parallelized. Their data structure uses an array of pointers to the nodes on the left or right spine of the input tree with each rank. This provides fast access to spine nodes of a given rank, and enables theoretical efficiency. In their paper \[7\], their implementation was compared to a previous version of the PAM library, and achieved better performance. Their experiments combine a bulk (an array) to a tree structure, while our union algorithm combines two trees. As mentioned above, our implementation maintains extra metadata such as reference count and size, and uses binary tree. The advantage of their algorithm is likely due to the better space efficiency (less metadata in each tree node) and I/O efficiency by using \((a, b)\)-trees (multiple keys maintained in one tree node). We choose to store extra information and use binary structure because we want to support more functionalities. Because of the extra space used, our implementation supports persistence (multi-versioning) with garbage collection, and more operations such as selecting the \(k\)-th element (see more details in Section 6). Also, using the binary structure enables cheaper path-copying (for persistence), which would be expensive for a regular B-tree or \((a, b)\)-tree (see details below). This is important for some applications such as databases with multi-versioning \[61\]. Although these functionalities are not specifically necessary for implementing set operations, we do so in order to support the general-purpose ordered map interface.

P-trees have also been compared with concurrent B-tree-like structures, and have been shown to achieve competitive or better performance by batching the updates and using a divide-and-conquer algorithm similar to union \[61\].

It is also possible to extend our join-based algorithms to B-tree-like data structures. One concern of using B-trees is that the large internal nodes in B-trees are expensive to copy, and therefore not well suited for supporting persistence. In 2019, Dhulipala et al. \[29\] extend the join-based algorithms to C-trees based on treaps. Instead of supporting a full ordered map interface, C-trees only focused on graph streaming. While C-trees are binary trees, they store multiple keys (in their case, graph edges) in the same tree node to allow for better cache locality and compression. Each node is represented by a randomly sampled head key \(k\), and a subsequent chunk, which contains all elements between \(k\) and the next head. Each chunk can further be compressed. A C-tree is then represented as a prefix, which is the chunk before the first head, and a tree, where each node contains a head and its subsequent chunk. When a persistent update occurs, only the head of the relevant internal nodes need to be copied. C-trees are shown to be practical for graph streaming. The C-tree provides useful insights for extending our work to B-tree-like data structures. To extend our general ordered map library to B-tree or its relatives, and to enable both good theoretical bounds and practical performance is an interesting future direction.

### 8 CONCLUSION

In this article, we study parallel balanced binary trees. We proposed the join-based algorithms, along with its analysis framework. We show conditions to make trees joinable, such that the
join-based (parallel) algorithms work correctly and efficiently. In particular, we show for the first time that a very simple “classroom-ready” set of algorithms is indeed work optimal when used with four different balancing schemes—AVL, RB, WB trees and treaps—and also highly parallel. The only tree-specific algorithm that is necessary is the join algorithm. For analysis, our approach defines the notion of rank (differently for different balancing schemes) and shows invariants on the rank. The definition of rank, along with the join algorithm for a specific balancing scheme, ensures that the join-based algorithms are work optimal with polylogarithmic span.

We also test the performance of our algorithm. Our experiments show that our sequential algorithm is about $7 \times$ faster for union on two maps of size $10^8$ compared to the STL RB tree implementation. In parallel, our code outperforms or is competitive to two baseline algorithms (MCSTL and WBB-tree) on different input sizes. Our code also achieves $73 \times$ speedup on 72 cores with hyperthreading.

ACKNOWLEDGMENTS

We thank the reviewers for their comments and suggestions.

REFERENCES


Received September 2020; revised October 2021; accepted December 2021.