Master's Thesis

# A Practical Scalable Shared-Memory Parallel Algorithm for Computing Minimum Spanning Trees 

Wei Zhou

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Supervisors: Prof. Dr. rer. nat. Peter Sanders Prof. Guy E. Blelloch

Department of Informatics
Karlsruhe Institute of Technology

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Wei Zhou


#### Abstract

The present thesis briefly reviews the history of the Minimum Spanning Tree (MST) Problem and a number of known algorithms to solve it. Furthermore, a new simple, elegant and practical algorithm based on Borůvka's algorithm for parallel MST computation on shared-memory machines is developed. The algorithm utilizes a parallel primitive called priority write that is easily and efficiently implementable with atomic compare-and-swap (CAS) instructions. The parallelism in the algorithm is coarse-grained and no explicit locks other than in the implementations of usual parallel primitives are needed. Experiments show that the algorithm is efficient on both synthetic and real-world graphs and is invulnerable to adversarial inputs. In our tests, it performs faster than or as fast as state-of-the-art implementations, which often perform poorly or even sequentially on particular classes of graphs. The new algorithm offers good performance even with few processors and therefore can be used as a sole universal implementation.


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## 1 Introduction

Given a connected weighted undirected graph $G=(V, E)$ where $V$ and $E$ are the set of vertices and edges, respectively, and an edge weight (or cost) function $w: E \rightarrow \mathbb{R}^{+}$, the Minimum Spanning Tree (MST) of G is defined as a connected subgraph $\mathrm{T}=\left(\mathrm{V}, \mathrm{E}_{0}\right)$ such that $\mathrm{E}_{0} \subseteq \mathrm{E}$ and the total weight of the chosen edges, $\sum_{e \in \mathrm{E}_{0}} w(e)$, is minimized. T must be a tree because fewer edges will disconnect the graph while more edges will introduce cycles in T and removing any edge on a cycle will reduce the total weight while keeping T connected. Some authors explicitly require acyclicness in the definition of MST.

The definition can be extended to unconnected G. In that case, we define the Minimum Spanning Forest (MSF) to be the set of the MSTs of all the connected components of G. We may also allow non-positive weights in the definition of MST by explicitly requiring T to be a tree with the lowest total weight. A graph with marked MSF edges is given in Figure 1.


Figure 1: A graph and its MSF. The boldface edges are in the MSF.
The Minimum Spanning Tree (Forest) Problem is to find an MST (MSF) for a given graph $\mathrm{G}=(\mathrm{V}, \mathrm{E})$. Note that there can be more than one MST (MSF) for a graph with the same minimum total weight. Any single one is a valid solution to the problem.

Minimum spanning trees have many theoretical and practical applications. The most direct and obvious one is network design, e.g. road network or electrical grid construction. The goal of the former is to connect all cities with roads and we would like to minimize the cost for the construction. MST problem also occurs as the critical part of other algorithms. Notable examples include an approximation algorithm to the NP-Hard Metric Traveling Salesman Problem (Metric TSP) ${ }^{1}$ that is no worse than twice of the optimal cost (see e.g. [1]). The MST problem is also closely related to the concept of matroids [1].

Largely due to physical limitation, the clock frequency of a CPU and its computing power cannot rise arbitrarily. In contrast, however, the data volume of modern information processing is growing faster than ever. This motivates the study of parallel computation.

[^0]In this thesis, we develop an efficient coarse-grained parallel algorithm for the MST problem for shared-memory architectures. The algorithm is applicable on multi-core computers that are ubiquitous nowadays. Experimental results indicate that, at least for the MST problem, even consumer-grade PCs can handle huge graphs efficiently.

The thesis is outlined as follows. The next section, Section 2, introduces some fundamental properties of MSTs that enable their efficient computation and then briefly presents some known algorithms, both classical and modern ones, serial and parallel ones, for MST computation. Section 3 focuses on the main result of the thesis, a new parallel algorithm for the MST problem based on Borůvka's algorithm introduced in Section 2. The new algorithm is evaluated on synthetic and real-world graphs in Section 4 together with some of the known algorithms in Section 2. Section 5 concludes the thesis with a summary and some outlook on future research.

Throughout this thesis, we use $\mathrm{n}=|\mathrm{V}|$ and $\mathrm{m}=|\mathrm{E}|$ to denote the number of vertices and that of edges, respectively. Vertices are uniquely numbered and identified from 0 to $n-1$ and edges from 0 to $m-1$. An edge is represented by a tuple $e=(u, v, w)$ where $u$ and $v$ are the two endpoints and $w$ the weight. We use e.u, e.v and e.w to denote the three components of the tuple of edge $e$. We also assume that the edge weights are distinct. Ties can be broken by any deterministic property of the edges, e.g. unique edge indices. For graphs with distinct edge weights, the MST is also unique. This is easily proved by contradiction using the Cycle property described in Section 2. Unless otherwise noted, "graphs" in this thesis are assumed to be connected, undirected and weighted. The thesis focuses on MST instead of MSF. This does not incur loss of generality because all of the presented algorithms can be easily adapted for MSF computation without raising their asymptotic time complexities. For sequential algorithms, one can first perform a breadth-first search and run the MST algorithms on each connected component. Most parallel MST algorithms can handle unconnected graphs natively because of the way they work. We can otherwise always resort to a parallel algorithm to the connectivity problem ${ }^{2}$ which is intuitively an easier problem (see e.g. [2]). Since most of the algorithms discussed in this thesis can trivially be implemented with $\mathcal{O}(m+n)$ space which is asymptotically optimal ${ }^{3}$, space complexity is generally omitted throughout the thesis unless noteworthy.

[^1]
## 2 Previous Work

The Minimum Spanning Tree Problem is one of the fundamental problems of algorithmics and therefore has been extensively studied. In this section we briefly introduce some of the known algorithms for solving the MST problem. We first look at some classic serial algorithms and then later parallel ones.

MSTs have various properties that can be exploited to compute them. The following are some of the most frequently used ones. All of those properties have simple proofs by contradiction which are omitted here (see e.g. $[1,3]$ ).
Property 2.1 (Cycle property). If G contains a cycle C and there is only one edge e on C whose weight is strictly higher than edges on C , then e cannot be in an MST.
Definition 2.2 (cut, cut edge). A cut C of a graph G is a partitioning of its vertices into two disjoint sets $\mathrm{V}_{0}$ and $\mathrm{V}_{1}$. A cut edge with respect to C is an edge that connects a vertex in $\mathrm{V}_{0}$ and another in $\mathrm{V}_{1}$.
Property 2.3 (Cut property). If a cut edge e with respect to cut $\mathrm{C}=\left\{\mathrm{V}_{0}, \mathrm{~V}_{1}\right\}$ is strictly lighter than any other cut edges with respect to C , then e belongs to an MST. In the case where all edges have distinct weights, e belongs to the MST.
Property 2.4 (uniqueness). If all edges in graph G have distinct weights, then G has a unique MST.

### 2.1 Classical Algorithms

### 2.1.1 Borůvka's Algorithm

The earliest explicit formulation of the MST problem and an efficient MST algorithm is believed to be due to Borůvka in 1926 [4, 5, 6]. His algorithm has been re-discovered several times during the next decades by others, e.g. Sollin [6], and therefore bears many names, most notably "Borůvka's algorithm" and "Sollin's algorithm". We call it "Borůvka's algorithm" throughout the thesis.

Borůvka's algorithm runs in phases. It maintains the set of found connected components, denoted S , during the process. It starts with n trivial components, each containing a single vertex. We then iteratively reduce the number of components in the set by so-called Boruivka steps in each phase until there is only one connected component in the set.

A Borůvka step basically tries to find safe edges and add them to the MST by exploiting the Cut property. It does so by finding the minimum outgoing edge ${ }^{4}$ for every connected component in the set $S$. By the Cut property (by choosing the cut $C=(T, V \backslash T)$ for each component $T \in S$ ), those edges must be in the MST. After adding those edges to MST, it joins the respective connected components of the two endpoints of every found MST edge.

It is easy to prove that no more than $\mathcal{O}(\log \mathfrak{n})^{5}$ Borůvka steps are needed. That is because the number of vertices in the smallest connected component at least doubles after each step and a connected component cannot contain more than $n$ vertices. The total time complexity of Borůvka's algorithm is therefore $\mathcal{O}\left(\log n \cdot T_{\text {borstep }}(m, n)\right)$, where $T_{\text {borstep }}(m, n)$ depends on the

[^2]concrete implementation of Borůvka steps. Borůvka steps can be implemented in $\mathcal{O}(\mathfrak{m})$ time so that the total running time of Borůvka's algorithm is $\mathcal{O}(m \log n)$. One such implementation is described below.

The implementation assumes an edge-list representation ${ }^{6}$ of graphs and connected components are implicitly defined by recording the representative of the connected component of every vertex. Let $R[u], u \in V$, denote the representative of the connected component in which vertex $u$ lies. The implementation works in three steps: find-min, grafting and then shortcutting.

Find-Min Step. The find-min step enumerates all edges that connect two different connected components and updates the current lightest outgoing edge for both connected components. By using $R$ values to check whether both endpoints are in the same component, it is easy to see find-min works in $\mathcal{O}(\mathrm{m})$ time.

Grafting Step. After finding the minimum edges, they are added to the MST. The grafting step merges the two connected components joined by any such minimum edge $e=(u, v, w)$ by "grafting" one component to the other, i.e. setting $R[i] \leftarrow \mathfrak{j}$ where $\mathfrak{i}=R[u]$ and $\mathfrak{j}=R[v]$ are the representatives of the components of $u$ and $v$, respectively. Note that after this step, the meaning of $R$ changes slightly so that the true representative of the component containing a non-representative vertex $\mathfrak{u}$ is found by following the path defined by those $R$ values, i.e. by iteratively setting $u \leftarrow R[u]$ until it stops changing. However, this may never stop when there is a cycle. This issue is discussed later. This grafting step takes $\mathcal{O}(n)$ time because every connected component has no more than one minimum outgoing edge and we only do an assignment for each such edge.

Shortcutting Step. The final shortcutting step then fully "shortcuts" all such paths by finding the true representative of $\mathfrak{i}$ 's component and then setting the $R$ values for all vertices along this path to this representative. This step takes $\mathcal{O}(n)$ time because each transition from $\mathfrak{i}$ to $R[i]$ will only be processed at most twice - once while trying to find the true representative and once while shortcutting along the path - and there are only $\mathcal{O}(n)$ such transitions.

The total running time of this Borůvka step implementation is therefore $\mathcal{O}(m)$, implying an $\mathcal{O}(m \log \mathfrak{n})$-time implementation of Borůvka's algorithm. Algorithm 1 details this implementation of Borůvka step and Borůvka's algorithm is given in Algorithm 2.

As mentioned before, there is a caveat in the grafting step: an edge could be chosen as the minimum edge by the components of both its endpoints. It is thus important not to add an edge to the MST twice in line 8 of Algorithm 1 on the one hand. On the other hand, careful analysis reveals that the relation " $\rightarrow$ " where " $A \rightarrow B$ " means the minimum edge for connected component $\mathcal{A}$ goes to component B , forms a pseudo-forest, meaning there is no cycle except for roots, where every root also points back via the same minimum edge, forming cycles of length 2 (see Figure 2). This situation results in an infinite loop in the shortcutting step. This can be solved by breaking the symmetry during the grafting: before setting $R[i] \leftarrow \mathfrak{j}$ we check if the minimum edge for the current ( $i$ 's) component is also the minimum for $j$ 's and, if so, we only graft $\mathfrak{i}$ to $\mathfrak{j}$ when $\mathfrak{i}>\boldsymbol{j}$. This decision is arbitrary as long as it is deterministic - we may also choose to graft when $\mathfrak{i}<\mathfrak{j}$. The solution is reflected in line 13 .

There are other popular implementations of Borůvka's algorithm assuming different graph representations. The edge-list representation is given in detail because it is particularly suitable for the new algorithm in this thesis and enables a more elegant parallelization. A asymptotically faster variant of Borůvka's algorithm by Yao [7] is given in Section 2.2.1.

[^3]```
Algorithm 1: Borůvka Step
    Data: Graph \(G=(\mathrm{V}, \mathrm{E})\)
                            Array \(R[0 . . n-1]\), where \(R[u]\) is the representative of the connected component in
            which vertex \(u\) lies
    Result: Updated array of representatives
    begin
        best \([i] \leftarrow\) sentinel, \(i \in \mathrm{~V} \quad / *\) assuming sentinel is an edge of weight \(\infty\) */
        /* find-min step */
        foreach \(e \in E\) where \(R[u] \neq R[v]\) do
            if e.w \(<\operatorname{best}[R[e . u]] . w\) then
                best \([\mathrm{R}[\mathrm{e} . \mathrm{u}]] \leftarrow e\)
            if \(e . w<\operatorname{best}[\mathrm{R}[e . v]] . w\) then
                best \([R[e . v]] \leftarrow e\)
        /* best \([i]\) is now the lightest edge leaving \(\mathfrak{i}\) 's component if \(i\) is a
            representative and there are still valid edges connecting i's
            component and another; otherwise best \([i]=\) sentinel. */
        Add those minimum edges to the MST (e.g. by marking these edges).
        /* grafting step */
        foreach \(i \in V\) do
            if best \([i] \neq\) sentinel then \(\quad / * i\) is a representative */
                \(u \leftarrow\) the endpoint (of edge best \([i]\) ) in the other component
                \(j \leftarrow R[u]\)
                if best \([i]=\operatorname{best}[j]\) and \(i<j\) then
                Do nothing /* break symmetry; R[i] stays i */
                else
                \(\mathrm{R}[\mathrm{i}] \leftarrow \mathrm{j} \quad / * \operatorname{graft} \mathrm{i}\) to j */
        /* shortcutting step */
        foreach \(i \in V\) do
            \(r \leftarrow i\)
            while \(r \neq R[r]\) do
                \(r \leftarrow R[r]\)
            /* \(r\) is now the representative of \(i\) 's component */
            \(j \leftarrow i\)
            while \(R[j] \neq r\) do
                \((R[j], j) \leftarrow(r, R[j]) \quad / *\) simultaneous assignments */
```

```
Algorithm 2: Borůvka's Algorithm
    Input: Graph \(G=(\mathrm{V}, \mathrm{E})\)
    Output: The MST of G
    begin
        \(\mathrm{R}[\mathrm{i}] \leftarrow \mathfrak{i}, \mathfrak{i} \in \mathrm{V} \quad / *\) initialization \(* /\)
        \(c c \leftarrow n \quad / *\) number of connected components implicitly defined by \(R\) */
        while cc \(>1\) do
            Invoke Borůvka step (Algorithm 1)
            \(\mathrm{cc} \leftarrow|\{i \in \mathrm{~V} \mid \mathfrak{i}=\mathrm{R}[\mathrm{i}]\}| \quad / *\) every component has a representative */
        return MST edges found during Borůvka steps
```



Figure 2: A graph and the pseudo-forest formed during a Borůvka step. The dashed arrows define the pseudo-forest. Note that three 2-cycles are formed. All of them must be broken to prevent infinite loops in the shortcutting step.

### 2.1.2 Prim's Algorithm

Prim's algorithm [8] (independently by Dijkstra [9] at around the same time), a three-decade later re-discovery of Jarník's algorithm [6], is another classical algorithm on the MST problem. It, together with Kruskal's algorithm discussed later, has received better coverage in standard texts on algorithms than Borůvka's algorithm, perhaps because the most recent re-discovery of Borůvka's algorithm by Sollin (that aroused renewed interest therein [6]) appeared a couple of years later than these two.

The algorithm works by choosing an arbitrary vertex as the starting MST and then growing the tree gradually by finding the nearest vertex to the current tree which has not been added to it. The correctness of the algorithm follows from the Cut property.

The running time of Prim's algorithm depends on how fast the nearest vertex can be found. For a dense graph $\left(m=\Theta\left(n^{2}\right)\right)$, it suffices to maintain the minimum distance between each vertex and the MST in an array D (initialized with oss). This way, the next vertex can be found by a loop in time $\mathcal{O}(n)$. After adding the vertex, the minimum distances stored in array $D$ are updated for its neighbors, which again takes $\mathcal{O}(n)$ time. The algorithm terminates after $n-1$ steps and thus has a time complexity $\mathcal{O}\left(n^{2}\right)$, which is asymptotically optimal.

The running time can be improved for sparse graphs with priority queues. A priority queue is an abstract data type that supports insert, delete, find-min, delete-min and decrease-key operations. find-min finds the smallest element in the priority queue and delete-min removes it. decrease-key takes a pointer to a present element in the priority queue and decreases its key by a given (non-negative) difference. General delete operation can be implemented by a decrease-key with a sufficiently large difference followed by a delete-min.

If we maintain the vertices in a priority queue with their respective minimum distances as the keys, finding the nearest vertex reduces to finding (and deleting) the minimum in the priority queue. After adding the minimum edge and the nearest vertex $u$ to the MST, we update the distances for u's neighbors as before or add a neighbor if it has never been present in the priority queue. Because we only need to update vertex $v$ 's distance if the weight between $u$ and $v$ is smaller than the current distance of $v$, the update can be implemented by a decrease-key operation. Thus we need ( $n-1$ ) delete-min operations, ( $n-1$ ) inserts and $\mathcal{O}(m)$ decrease-keys. The running time of Prim's algorithm then fully depends on the concrete priority queue implementation. For binary heaps (see e.g. [1]), all three operations have complexity $\mathcal{O}(\log n)$ where $\mathfrak{n}$ is the number of elements in the heap, giving an $\mathcal{O}((m+n) \log \mathfrak{n})$ implementation of Prim's algorithm. Advanced priority queues like Fibonacci heaps [10] and thin heaps [11] offer constant amortized time complexity for all standard operations except for deletions which take $\mathcal{O}(\log n)$ time, enabling an $\mathcal{O}(m+n \log n)$ time implementation of Prim's algorithm, though the constant factor hidden in the complexity limits their usefulness [12, 13]. On the contrary, a data structure named pairing heap [14] proves to be fast also in practice [12, 13], though its decrease-key operation is now known to be $\Omega(\log \log \mathfrak{n})$ [15], i.e. not optimal after a decade of being conjectured so. A modified version of pairing heaps achieving this lower bound is also known [16].

An interesting aspect of Prim's algorithm with binary heaps is its expected time complexity when the edge weights are random. [17] proves that the expected running time in this case is $\mathcal{O}(m+n \log n \log (1+m / n))=\mathcal{O}(m+n \log n \log \log n)^{7}$ even when an adversary gets to choose

[^4]the graph topology as long as the weights are random. This justifies the efficiency of this simple implementation demonstrated in practice [12, 13].

### 2.1.3 Kruskal's Algorithm

Another standard algorithm for computing MSTs in sequential settings is Kruskal's algorithm [18]. It maintains the set of connected components starting with $n$ trivial ones just as in Borůvka's algorithm. However, it does not locally choose the minimum edge for each component, but globally finds the minimum edge connecting two different components and then joins them in one step. The procedure is executed until the number of connected components is reduced to one. The most prevailing way to implement the searching procedure is to first sort all edges by their weights in ascending order and inspect every edge in the sorted order to see whether both endpoints are in the same component. If they are, the edge is skipped; otherwise both components are joined and the edge is marked to be in the MST.

If we abstract the needed operations from the algorithm, we in fact need the so called disjoint set or union-find abstract data type that supports

- make-set $(x)$ : making a singleton set containing $x$,
- find $(x)$ : finding the representative of the set in which $x$ lies, and
- unite $(x, y)$ : joining the sets containing two given elements $x$ and $y$.

The most suitable union-find data structure for Kruskal's algorithm is similar to the R array used in Borůvka's algorithm. Each set is represented as a tree that is implicitly defined by those $R$ values where $R[i]$ is the parent of $i$ in its tree. The representative of $i$ 's set is found by following $R$ until we reach the top (where $R[i]=i$ ). Merging is done by grafting the root of one tree to that of the other.

This representation looks simple but inefficient. However, it can be proven that, when two techniques called path compression and union-by-size are employed, the total time for executing any sequence of operations containing $m$ finds and $n$ make-sets (thus at most ( $n-1$ ) unites) is $\mathcal{O}(n+m \alpha(m+n, n))$, where $\alpha(m, n)$ is a very slow-growing function called the inverse Ackermann function ${ }^{8}$ that is no larger than 4 for all practical inputs [19]. Path compression is almost what we did in the shortcutting step in Borůvka's algorithm: for every find(x) operation, after finding the root $r$, we set the parent of every element $y$ on the path from $x$ to $r$ also to $r$, i.e. $R[y] \leftarrow r$. Intuitively, this makes future find operations on those elements much cheaper without increasing the complexity of the present one. Union-by-size adds an attribute to every root representing the size (number of elements) in its tree. When doing an unite operation, we only graft the tree of smaller size to the larger one, breaking ties arbitrarily, and update the size attribute of the new root. The size field of the smaller tree will never be used again thereafter and can be discarded if necessary. Both techniques are depicted in Figure 3 and Figure 4. They are very easy to implement yet the analysis is highly non-trivial and out of the scope of this thesis. [19] offers a complete analysis and more techniques that can be used to replace path compression and/or union-by-size and are asymptotically equally fast. Furthermore, it has been also proven in [19] that the time bound is tight in a sense: any implementation of the union-find data structure needs $\Omega(n+m \alpha(m+n, n))$ time in the worst case to execute a

[^5]sequence containing $m$ finds and $n$ make-sets for pointer machines ${ }^{9}$ under certain technical assumption. For $m \geqslant n$, which is the case in Kruskal's algorithm, a simpler lower and upper bound of $\Theta(m \alpha(m, n))$ can be proven.

(a) A union-find tree.

(b) After find(4).

Figure 3: Path compression

(a) A union-find data structure.

(b) After unite $(4,0)$. The sizes of of both trees are 2 and 4 respectively, thus 4 is grafted to 0 .

Figure 4: Union-by-size.
Kruskal's algorithm can be efficiently implemented using this union-find data structure. Sorting the edges costs $\mathcal{O}(\mathfrak{m} \log \mathfrak{m})$ time. After that, we have to inspect every edge in the worst case, involving two find operations per edge, and possibly join both components which are $(n-1)$ unite operations in total. Thus we need $\mathcal{O}(m \alpha(m, n))$ time in the worst case. In summary, Kruskal's algorithm has a time complexity of $\mathcal{O}(\mathfrak{m} \log \mathfrak{m})$ or, if the edges are already given in sorted order, $\mathcal{O}(\mathfrak{m} \alpha(m, n))$.

### 2.2 Modern Algorithms

On top of the discussed classical algorithms, many more algorithms that are theoretically or practically more efficient have been proposed. In this section we give a brief introduction to some of them. We start with variants of these classical algorithms and then look at some of the more recent and advanced ones.

[^6]
### 2.2.1 Yao's Algorithm

Borůvka's algorithm makes local decisions during each Borůvka step. If we can speed up these local decisions, we can then accelerate Borůvka step. A natural direction to try is to avoid inspecting every edge during each Borůvka step somehow. It should be possible to implement Borůvka step in less than $\Theta(m)$ time if we can manage this because other steps in a Borůvka step only takes $\mathcal{O}(\mathfrak{n})$ time in contrast to $\mathcal{O}(m)$.

Yao's algorithm [7] is the first MST algorithm that achieves an $\mathcal{O}(m \log \log n)$ time bound in the worst case. It assumes an adjacency array ${ }^{10}$ (or list) representation of graphs. The outbound edges for every vertex are first partitioned into $k=\log n$ equal-sized groups such that every edge in a later group is at least as heavy as all edges in earlier groups for the same vertex. Edges within the same group are not sorted. This can be done in $\mathcal{O}(m \log k)=\mathcal{O}(m \log \log \mathfrak{n})$ time if we recursively apply the classical linear selection algorithm (see e.g. [1]). The goal of the partitioning is to make sure later groups do not have to be checked before all earlier ones are fully exhausted. Therefore only one group, i.e. only $\frac{1}{k}$ of all edges for each vertex have to be checked to determine the minimum edge for this vertex. We remember the current group ID for every vertex across the Borůvka steps. After determining the minimum edge for a vertex, we mark it or (lazily) remove it. If exhausting the whole group did not yield any usable edge, we increment the group ID for the vertex and search again in the next group. The total cost for failed searches of the groups for an unused edge is bounded by the number of edges across all Borůvka steps and is therefore $\mathcal{O}(\mathrm{m})$. We still have $\mathcal{O}(\log \mathfrak{n})$ stages, so the total time for all stages is $\mathcal{O}\left(m+\log n \cdot \sum_{i=1}^{n}\left\lceil\frac{\mid \text { neighbors }(i) \mid}{k}\right\rceil\right)=\mathcal{O}\left(m+\log n \cdot\left(\frac{2 m}{k}+n\right)\right)$ which is $\mathcal{O}(m+n \log n)$ for $k=\log n$. For $m=\Omega(n \log n)$ this is a linear time algorithm. In order to eliminate the $n \log n$ term for the case $m=\mathcal{O}(n \log n)$, we first run normal Borůvka steps (with $k=1$, i.e. no partitioning) for $\log \log n$ stages to reduce the number of vertices by a factor of $\log n$. This preprocessing takes $\mathcal{O}(\mathfrak{m} \log \log n)$ time, which is also the running time of the whole algorithm. Yao's algorithm is often considered impractical because it involves the linear selection algorithm which has a high constant factor [20].

### 2.2.2 Cheriton-Tarjan Algorithm

Another $\mathcal{O}(\mathfrak{m} \log \log \mathfrak{n})$ algorithm for MST is due to Cheriton and Tarjan [21]. The algorithm also maintains a set of connected components implicitly represented by the representatives of every component, just as in Borůvka's algorithm. For every component it manages a meldable priority queue, i.e. a priority queue that supports efficient merging of two queues, of all edges having exactly one endpoint inside the component. Edge weight is the key of the priority queues.

The algorithm works in a round-robin fashion. At the beginning we have $n$ trivial connected components in a (normal, first-in first-out) queue. At each step, the algorithm pops the first element from the queue and then the minimum edge from its priority queue. We find the other endpoint of the edge (the one that is not in this component) and its corresponding component and remove it from the queue. We then merge the two connected components and their associated priority queues. Finally the merged component is appended to the queue. The

[^7]steps are repeated until we have only one connected component in the queue. This connected component represents the MST of the original graph.

Components can be represented by the same union-find data structure as in Kruskal's algorithm. Therefore the merging of two connected components can be efficiently implemented. It remains to find a priority queue implementation that supports efficient merging. Cheriton and Tarjan chose a variant of so-called leftist heaps invented by Crane [22]. A leftist tree is a binary tree such that, for every node in the tree, the right path from the node (the path along the pointers of the right children until there is none) is the shortest among all paths from the node to the bottom. Another way to characterize leftist trees is to define a function $\operatorname{rank}(x)$ for every node $x$ as

$$
\operatorname{rank}(x)= \begin{cases}0, & \text { if } x \text { is an external node } \\ 1+\min \{\operatorname{rank}(\operatorname{left}(x)), \operatorname{rank}(\operatorname{right}(x))\}, & \text { otherwise }\end{cases}
$$

where left $(x)$ and $\operatorname{right}(x)$ are the left and right child of $x$, respectively. A leftist tree is defined as a binary tree satisfying $\operatorname{rank}(\operatorname{left}(x)) \geqslant \operatorname{rank}(\operatorname{right}(x))$ for every internal node $x$, hence its name. A leftist heap is then a leftist tree satisfying the heap order as in binary heaps. In addition to insert and delete-min, leftist heaps also support merge operations that merge two leftist heaps in $\mathcal{O}(\log n)$ time, where n is the total number of elements in both trees, in contrast to ordinary binary heaps, for which it would take $\mathcal{O}(n)$ time. The variant used in Cheriton-Tarjan algorithm uses lazy deletion and lazy merging to achieve the claimed $\mathcal{O}(m \log \log n)$ time bound. The running time of this algorithm can be improved slightly further to $\mathcal{O}\left(m \log \log _{2+m / n} n\right)$ by doing some cleanup operations that remove duplicate or self edges (edges that connect vertices in the same connected component) at appropriate intervals. Further details can be found in [21] and [20]. This algorithm is also deemed impractical because the small speedup over classical algorithms does not overweigh the constant factors due to the use of more complicated pointer based data structures.

### 2.2.3 Variants of Kruskal's Algorithm

The bottleneck of Kruskal's algorithm resides in the sorting step. Intuitively, the number of connected components is often reduced to one (thus the MST has already been found) long before every edge is inspected. A result from the theory of random graphs states that the expected number of edges that need to be checked is about $\frac{1}{2} n \ln n$ for large enough $n$ [23], much fewer than $m$ for dense graphs.

A method to incorporate this observation is to build a priority queue on the edges instead of sorting and then use delete-min to get the next lightest edge until the MST has been built. Building a priority queue takes $\mathcal{O}(m)$ time for many implementations, e.g. binary heaps (see e.g. [1]). Each subsequent delete-min needs $\mathcal{O}(\log m)$ time. The worst-case time complexity is therefore $\mathcal{O}(m+m \log m+m \alpha(m, n))=\mathcal{O}(m \log n)$ just as the original algorithm, but the average case is now only $\mathcal{O}(m+n \ln \mathfrak{n}(\log \mathfrak{m}+\alpha(m, n)))=\mathcal{O}\left(m+n \log ^{2} \mathfrak{n}\right)$. This implementation is called Kruskal's algorithm with demand-sorting by some authors [12].

Another algorithm demonstrating this early-stopping idea works like Quick-sort. If the number of edges is small enough (e.g. $m=\Theta(n)$ ), we run a normal Kruskal's algorithm with union-find. Otherwise, we choose a pivot edge (uniformly at random or the median) and partition the edges into two sets: $\mathrm{E}_{1}=\{e \mid e . w \leqslant$ pivot. $w\}$ and $\mathrm{E}_{2}=\{e \mid e . w>$ pivot.w $\}$. We
then recurse with only the edges in $E_{1}$. After returning from the recursion, we check if the MST has already been built and only recurse with edges in $E_{2}$ if it has not. During the whole algorithm, a global union-find data structure is used. This algorithm, sometimes called the Quick-Kruskal algorithm [24], has the same average time complexity as Kruskal's algorithm with demand-sorting, i.e. $\mathcal{O}\left(m+n \log ^{2} \mathfrak{n}\right)$ [25].

The above two algorithms with early-stopping only work well for random graphs with random weights as early stopping is of no use even if the MST contain a single heavy edge. If the topology is fixed and can be chosen by an adversary, random weights alone cannot guarantee the above average time complexity [24]. An example given in [24] is a "lollipop graph" that consists of a random graph and a tail (a chain) connected to one node. Obviously, all of the edges on the tail must be in the MST, and the probability that an edge is in the heavier half of all edges is $\frac{1}{2}$, implying about half of the edges on the tail are in the heavier half, rendering the early-stopping ineffective.

A simple remedy of the Quick-Kruskal algorithm called the Filter-Kruskal algorithm is given in [24]. It adds a filtering step which removes the edges that have endpoints in the same components after recursing processing the lighter half of the edges, before recursing with the heavier half. This filtering is done with help of the global union-find data structure. Experiments show that his heuristic is indeed much more robust than early-stopping alone. From a theoretical point of view, they proved that the expected running time of Filter-Kruskal algorithm is $\mathcal{O}\left(m+n \log n \log \frac{\mathfrak{m}}{n}\right)=\mathcal{O}(m+n \log n \log \log n)$ for arbitrary graph with random weights. Note the same bound is also achieved by Prim's algorithm with binary heaps under the same condition as mentioned before in Section 2.1.2.

The filtering idea can be used without recursive partitioning. We simply do a one-time partitioning with Quick-select (see e.g. [1]) to find the (e.g. $\Theta(n)$ ) lightest edges, run Kruskal's algorithm (or any other MST algorithm that return a union-find data structure) with those edges, do a filtering on the heavier edges, and invoke Kruskal's algorithm on the remaining heavier edges. This is what the parallel implementation of Kruskal's algorithm in the Problem Based Benchmark Suite (PBBS) [26] does. More details on that algorithm is given in Section 2.3.4.

### 2.2.4 Other Algorithms

In the last few decades, a couple of algorithms that are asymptotically faster than Yao's and Cheriton and Tarjan's $\mathcal{O}(m \log \log n)$ have been proposed. Most of them are highly complicated and thus deemed impractical, or are only applicable to graphs with certain special properties. We only give an incomplete listing here.

Fredman and Tarjan [10] gave an $\mathcal{O}\left(m \log ^{*} n\right)$ time algorithm for the MST problem that invokes Prim's algorithm implemented with Fibonacci heaps iteratively, where the iterated logarithm $\log ^{*} n:=\min \{i \mid \underbrace{\log \log \cdots \log n}_{i \text { times }} n \leqslant 1\}$ and $\log ^{*} n \leqslant 5$ for all practical input. The rationale of the algorithm is to limit the number of elements that coexist in the priority queue to reduce the $n \log n$ term in the time complexity of Prim's algorithm (which is $\mathcal{O}(m+n \log n)$ ). The algorithm runs in passes, each of which executes Prim's algorithm until the number of elements in the priority queue exceeds a certain threshold $k$ or after it just added a vertex into the priority queue that has been marked by a previous Prim instance, at which point it starts a new instance of Prim's algorithm in another vertex. The pass ends when every vertex belongs to the tree of an instance of Prim's algorithm. Every tree found by a Prim instance
is contracted into a super-vertex. The $\mathfrak{i}$-th pass can be implemented in time $\mathcal{O}\left(\mathfrak{m}+\mathfrak{n}_{\mathfrak{i}} \log k\right)$ where $n_{i}$ is the number of vertices before the pass. If we set $k=k_{i}=2^{\frac{2 m}{n_{i}}}$, that would be $\mathcal{O}\left(\mathfrak{m}+n_{i} \log \left(2^{\frac{2 m}{m_{i}}}\right)\right)=\mathcal{O}\left(\mathfrak{m}+n_{i} \cdot \frac{2 m}{n_{i}}\right)=\mathcal{O}(m)$. Furthermore, it can be shown that no more than $\frac{2 m}{k_{i}}$ super-vertices remain after pass $i$, because the total degree of every Prim tree after the pass is at least $k_{i}$ and the total degrees of the whole graph is 2 m . Therefore we have $k_{i+1}=2^{\frac{2 m}{n_{i+1}}} \geqslant 2^{\frac{2 m}{m / k_{i}}}=2^{k_{i}}$. Thus the sequence of $k_{i}$ increases tetrationally, indicating the number of passes is $\mathcal{O}\left(\log ^{*} \mathfrak{n}\right)$, proving a total time complexity $\mathcal{O}\left(m \log ^{*} \mathfrak{n}\right)$ for the algorithm. This algorithm is then (very) slightly improved by Gabow et al. [27] to $\mathcal{O}\left(m \log \log ^{*} n\right)$ shortly thereafter.

The bound for deterministic algorithms for the MST problem was again lowered by Chazelle [28] to $\mathcal{O}(\mathfrak{m} \alpha(\mathfrak{m}, \mathfrak{n}))$ by utilizing an interesting data structure called soft heaps also invented by him [29]. A soft heap is a meldable priority queue implementation that may corrupt elements stored in it by increasing their keys. Soft heaps support delete-min and merge in constant amortized time and insert in $\mathcal{O}\left(\log \frac{1}{\epsilon}\right)$, satisfying the additional property that no more than $\epsilon \mathfrak{n}$ elements in the data structure are corrupted at any time, where $0<\epsilon \leqslant \frac{1}{2}$ is the error rate. The errors are introduced to break the information-theoretic lower bound on priority queues because otherwise insertions and deletions could be used to do comparison-based sorting in $\mathrm{o}(\mathrm{n} \log \mathrm{n})$ time. A similar algorithm with the same bound has also been independently proposed by Pettie [30].

Pettie and Ramachandran [31] ultimately have broken the bound again by providing an asymptotically optimal deterministic comparison-based algorithm for the MST problem on pointer machines. An intriguing fact about the algorithm is that although the running time of their algorithm is proven to be matching the decision tree lower bound for pointer machines, this bound itself is not known. The best upper and lower bound of the running time of the algorithm to date are $\mathcal{O}(\mathfrak{m} \alpha(\mathfrak{m}, \mathfrak{n}))$ (achieved by Chazelle [28] as mentioned above) and $\Omega(\mathfrak{m})$, respectively. They have also proved that their algorithm works in linear time with high probability for random graphs even if the lower bound should later be shown to be superlinear. Roughly, their algorithm first generates optimal decision trees for graphs of no more than $r=\log \log \log n$ vertices in $\mathcal{O}(n)$ time. Then it partitions the graph into subgraphs of about $r$ vertices in $\mathcal{O}(m)$ time. After that, the MST for each subgraph is calculated by using the optimal decision trees, for which we do not know the exact time bound. The found MSTs are contracted into individual super-vertices in linear time. The remaining graph is dense because the number of vertices is reduced to $\frac{n}{r}$. For such a graph, the MST can be found in linear time by invoking previous algorithms (e.g. Fredman and Tarjan's $\mathcal{O}\left(m \log ^{*} n\right)$ time algorithm). Therefore the total running time is dominated by the application of the decision trees and other steps take linear time in total.

Note that the lower bound for the MST problem, should it prove to be superlinear, does not necessarily hold for models other than pointer machines. For example, on the so-called trans-dichotomous model, a unit-cost RAM model where a word cannot hold unreasonably much data, the MST can be found in deterministic linear time for integral edge weights [32]. Deterministic linear time can also be achieved for special graphs. For example, the MST problem for planar graphs can be solved in $\mathcal{O}(n)$ time and for dense graphs (where $\mathfrak{m}=\Omega\left(n^{1+\epsilon}\right)$ for some $\epsilon>0)$ in $\mathcal{O}(m)$ time [21]. The limit on the density can be relaxed to $\mathfrak{m}=\Omega(n \log \log \log n)$ while retaining a linear time bound for all asymptotically faster algorithms described in this section [31]. This fact is exploited in the optimal algorithm described above.

If we have access to a stream of perfectly random bits with uniform distribution, the MST
problem can be solved in expected linear time by an algorithm given by Karger et al. [33]. Their algorithm relies on a linear verification algorithm, an algorithm that verifies the minimality of a spanning tree in linear time and returns witnesses for edges that do not belong in the true MST. One such verification algorithm on word RAM model is given by King [34] and one for pointer machines by Buchsbaum et al. [35]. With the latter, this MST algorithm can be completely implemented on pointer machines. The MST algorithm first executes Borůvka step twice to reduce the number of vertices by a factor of 4 . It then chooses a random sample of all edges by including each edge with $\frac{1}{2}$ probability and finds the MSF F of the chosen subgraph recursively. The heavy edges with respect to the found MSF F are then filtered out with help of the verification algorithm. This is an application of the Cycle property. If we denote the remaining light edges as L , the algorithm finally returns the MST of $\mathrm{F} \cup \mathrm{L}$ with a recursive call. The key observation that leads to the claimed running time is that $L$ has an expected size of only $\frac{n}{2}$. Therefore the expected running time is $T(m, n)=T\left(\frac{m}{2}, \frac{n}{4}\right)+O(n+m)+T\left(\frac{n}{2}+\frac{n}{4}, \frac{n}{4}\right)=O(n+m)$ where $T(m, n)$ denotes the running time of the algorithm on a graph with $m$ edges and $n$ vertices.

Katriel et al. [36] focused on a practical modification of the randomized linear time algorithm for dense graphs by ignoring the Borůvka steps, selecting a sample of edges of size $\sqrt{\mathrm{mn}}$ instead of $\frac{m}{2}$, utilizing Prim's algorithm instead of recursion and reducing the verification step to range minimum queries (RMQ) that make use of a byproduct of the first Prim instance, namely the order in which vertices are added into the MST. The resulting algorithm has expected time complexity $\mathcal{O}(m+n \log n+\sqrt{m n})$ if Prim's algorithm is implemented with Fibonacci heaps. They reported favorable performance with pairing heaps.

There are more algorithms that are based on the Cycle property. For example, the so-called reverse-delete algorithm that Kruskal described in the same paper as Kruskal's algorithm [18] works by looking at the edges in descending order by weight and delete an edge if it does not disconnect the graph. As a practical implementation would need an efficient algorithm for dynamic connectivity supporting edge deletion which is highly non-trivial, this algorithm is not usually used. An implementation by Thorup [37] achieves a running time of $\mathcal{O}\left(m \log n(\log \log n)^{3}\right)$.

### 2.3 Parallel algorithms

This section discusses some parallel algorithms for the MST problem. For that, we first give some fundamental definitions and constructs in parallel computing.

### 2.3.1 Preliminaries

Models of Computation The most-used computation models in shared-memory parallel computation are the Parallel Random-Access Machine (PRAM) models, an analogy to the RAM model for sequential algorithms. On a PRAM, we have a set of $p$ processors identified by unique indices called processor IDs. The number of processors may depend on the problem size. All processors have access to a global shared memory and have their own private registers and memory that others cannot access. All processors work synchronously and each can perform a standard arithmetic or logic operation on a memory cell within one clock cycle. There are multiple PRAM variants that differ in the way they handle concurrent memory accesses to the same memory cell within the same clock cycle: Exclusive-Read Exclusive-Write
(EREW): No concurrent reads or writes to the same memory location are allowed; ConcurrentRead Exclusive-Write (CREW): Processors may read a memory cell concurrently and will receive the same value, but no concurrent writes by multiple processors are allowed; and Concurrent-Read Concurrent-Write (CRCW): Processors may read or write to the same memory location at the same time. Technically it is also possible to define an Exclusive-Read Concurrent-Write (ERCW) PRAM, but this is rarely used because write access is generally assumed to imply read access. In the case of CRCW PRAM, it is important to define the result of concurrent writes. Again here are several variants given in the increasing order of ability: CRCW-Common where all processors writing to the same cell in the same clock cycle must write the same value and this value is stored into the cell; CRCW-Arbitrary where processors may write different values and an arbitrary (random) processor would succeed in writing its value into the cell while the values other processors want to write are ignored; CRCW-Priority where the processor with the smallest ID (thus highest priority) succeeds; and CRCW-Reduction where an associative operator (a reduction) somehow combines the written values the processors attempt to write into a single one and writes it into the cell, e.g. taking the sum, minimum or maximum. Although the latter two seem much stronger than CRCW-Common, which in turn looks stronger than EREW PRAM, it is possible to simulate a concurrent memory access of CRCW-Reduction or CRCW-Priority PRAM on an EREW PRAM within only $\mathcal{O}(\log \mathfrak{p})$ parallel time by exploiting the fact that sorting $n$ elements can be done on EREW PRAM in $\mathcal{O}(\log n)$ time (see [38]).

Within PRAM models, we can define quantities that characterize the performance of algorithms. Basically we have the depth which is the number of clock cycles that have elapsed before all processors (hence the program or algorithm) terminate, and work which is the sum of clock cycles of every processor in which the processor performed operations (in contrast to being idle). The main advantages of using depth and work to specify the complexity of a PRAM algorithm is the simplicity to translate them into the running time of the algorithm on a real-world computer via Brent's theorem. Brent's theorem, also called the Work-Time Scheduling Principle, states that we can simulate a PRAM algorithm of time $T(n)$ and work $\mathcal{W}(n)$ with unlimited number of processors on $p$ processors within $T_{p}(n)=\mathcal{O}\left(\frac{W(n)}{p}+T(n)\right)$ time on the same PRAM model [38].

In order to describe parallel algorithms, the following well-studied fundamental operations are defined and used as primitives later for describing parallel algorithms.

Parallel Map. Perhaps the simplest parallel construct is the map function. map(f,L) applies a function $f$ to every element in the sequence $L$ and returns a sequence containing the results. Formally, $\operatorname{map}(f, L):=\{f(x) \mid x \in L\}$. map is normally required to return the sequence of new values in the same order as their corresponding values in $L$. If the function $f$ is pure, i.e. it does not have any side-effect and its value only depends on the argument, $f(x)$ can be evaluated for every $x \in \mathrm{~L}$ in parallel. Due to the simplicity to parallelize such a map operation, it is sometimes called "embarrassingly parallel". If function $f$ does not have a return value, but does something to the environment based on its argument like writing to an array, map is also called parallel foreach or parallel for-loop. map is present in most functional programming languages, though it is not always executed in parallel.

Parallel Reduction (Fold). Another frequently used primitive is reduction or fold. For a pure associative binary function $f$ and a sequence $L$, reduce applies $f$ to the first two elements in L, and then successively applies $f$ to the last function value and the next element of L. Formally,
its functionality is defined as reduce $(f, L[0 . . n-1]):=f($ reduce $(f, L[0 . . n-2]), L[n-1])$ and reduce $(f, L[0 . .1]):=f(L[0], L[1])$. Often, we also define the result of reduce on an one-element list to be the list itself. Though any associative f can be used, the most frequently used ones by far are min, max and sum. The associativity of $f$ comes into play when we try to implement reduce in sub-linear time. Since reduce $(f, L[0 . . n-1])=f\left(\right.$ reduce $\left(f, L\left[0 . . \frac{n}{2}\right]\right)$, reduce $\left.\left(f, L\left[\frac{n}{2}+1 . . n-1\right]\right)\right)$ by associativity and both arguments of the outer application of $f$ are independently computable, reduce can be implemented in depth $\mathcal{O}(\log \mathfrak{n})$ on EREW PRAM. By Brent's theorem, we can simulate the algorithm in $\mathcal{O}\left(\frac{n}{p}+\log n\right)$ time on a machine with $p$ processors. If $p<\log n$, we can achieve a depth of $\mathcal{O}\left(\frac{n}{p}+p\right)$ easily by breaking the list into $p$ part and letting each processor compute the sum of one sublist. A single processor computes adds the sums of the sublists together to produce the final result.

Parallel Prefix-Sum (Scan). Prefix-sum, also called scan, is a stronger primitive than reduction and is present in many parallel algorithms. It also forms the foundation of other parallel primitives. This operation computes all prefix sums of a given sequence. Two versions of prefix-sum exist, namely inclusive and exclusive prefix-sums. They differ in whether $L[i]$ is included in the prefix-sum of position $i$. Prefix-sum is stronger than reduction because the last value of an inclusive prefix-sum is exactly the reduction. At first glance, it is not obvious that prefix-sum can be computed efficiently in parallel because its definition seems inherently sequential. However, it can also be computed in $\mathcal{O}(\log \mathfrak{n})$ time too by a two-phase algorithm sometimes called Blelloch scan [39]. Prefix-sums can also be generalized to any associative operators besides addition.

Parallel Filtering. With help of prefix-sum, we can implement another useful primitive: filtering. filter (pred, L) gives a sequence containing the elements of $L$ for which the predicate pred gives true in their original relative order, i.e. filter $(\operatorname{pred}, L):=\{x \in L \mid \operatorname{pred}(x)=\operatorname{true}\}$. For an efficient implementation of filter, we first use a parallel map to compute an array flags of 0 or 1 where flags $[i]=1$ if and only if $\operatorname{pred}(\mathrm{L}[i])$ is true. An exclusive prefix-sum of flags is calculated. This gives the count of elements before $\mathrm{L}[i]$ where the predicate gives true, and is thus the final position of $\mathrm{L}[\mathrm{i}]$ in the resulting sequence if $\operatorname{pred}(\mathrm{L}[i])$ is true. A final parallel for-loop copies these values to the corresponding position of the target array. A special case of filtering, pack(flags, L), simply stores elements of L whose flag values are 1 into another array. If we also want to get those elements with flags of 0 , we may use a partition or split which is slightly more efficient than two filtering operations because it can reuse some intermediate computations.

Parallel Sorting. As in sequential computation, sorting is also a fundamental building block of parallel algorithms. Many efficient algorithms exist for PRAM (see e.g. [38]). On real-world multi-core CPUs, people seem to prefer sample sort which proves to be efficient [40]. The algorithm chooses a sample of the original array, sorts the sample, chooses $m-1$ equidistant values as splitters and partitions the original array into $m$ disjoint buckets with those splitters. Buckets are recursively sorted and then concatenated to form the sorted array. We omit the details here.

The most natural choice of classical algorithm for parallelization is Borůvka's algorithm because of the way it makes decisions: every Borůvka step finds the minimum outgoing edge for a component locally, thus can be executed for all components in parallel. That is probably why

Borůvka's algorithm received far more attention in the era of parallel computation than Prim's and Kruskal's in the literature. We first look at some implementations of Borůvka's algorithm and then a few of Prim's and Kruskal's algorithm.

### 2.3.2 Borůvka's algorithm

Bader and Cong [41] gave several parallel implementations of Borůvka's algorithm and are among the first that reported reasonable speedup on shared-memory multiprocessors. All of these implementations do a true graph compaction (also known as contraction) after grafting in a Borůvka step, i.e. contract every component to a super-vertex, instead of the implicit one given in Algorithm 1. They use different representations of graphs and thus have slightly different implementations of Borůvka steps.

Variant 1. One implementation (called Bor-EL) uses an edge-list representation as in Algorithm 2 but every edge is stored twice in the list (one for each direction). The contraction step is simply sorting the edge list with the super-vertex (or representative as in Algorithm 2) of the first endpoint as the primary, that of the second endpoint as the secondary and the edge weight as the tertiary key. Self-loops within the same component and multi-edges between components are removed with a subsequent parallel prefix-sum.

Variant 2. A second implementation (Bor-AL) uses adjacency array as the graph representation where every edge is again in both lists of outgoing edges of its endpoints. The contraction step is accomplished by sorting the array of vertices by their representatives, which is equivalent to sorting all the edges by the representatives of their first endpoints, and then concurrently sort the outgoing edges for every vertex by the representatives of the second endpoints with a sequential sorting algorithm (insertion sort for short lists and merge sort for longer ones). Redundant edges are then removed with prefix sums.

Variant 3. A last implementation of Borůvka's algorithm which is perhaps more interesting uses a flexible adjacency lists (array) representation of graphs (denoted Bor-FAL). This implementation reduces the cost for compaction by allowing every vertex to hold a list of adjacency lists (arrays). After grafting, every vertex plugs its list of adjacency arrays to that of its representative and edges themselves do not have to be moved around. The find-min step has to check for redundant edges because self and multi-edges are not removed with this representation.

A fundamental problem with all these implementations is that their find-min steps are done concurrently for all vertices but sequentially for each vertex. That means there can be massive load imbalance if the degrees of the vertices differ too much, as is the case for graphs containing star-shaped subgraphs. Furthermore, the first two implementations uses the expensive parallel sort to build the contracted graph. The Bor-FAL implementation does not have this problem, but may degenerate to the cache-unfriendly adjacency-list representation if vertex degrees are evenly distributed.

Another implementation of Borůvka's algorithm using adjacency-array representation is given by da Silva Sousa et al. [42]. In the contraction step, the implementation uses atomic incrementing instructions to count the new number of outgoing edges to get the index of the first edge for each vertex. Subsequently it copies edges between to the new edge array, again using atomic increments. Self-edges within a component are removed but multi-edges between components are not. This implementation also suffers from the load-imbalance problem in the
find-min step and in the contraction step due to atomic increments.
A very recent work of Cong and Tanase [43] aims to reduce the cost for memory accesses and improve locality in Borůvka's algorithm, generalizing ideas proposed in [44]. Their implementation (independently) exploits some ideas similar to the present thesis. We feel this thesis is still justified because the new algorithm proposed here has other new ideas and the design was finalized prior to the publication of their work. When describing the new algorithm we will give reference to their work that demonstrates the same or similar idea.

### 2.3.3 Prim's algorithm

Prim's algorithm is apparently inherently sequential and hard to scalably parallelize. Known algorithms run several instances of Prim's algorithm starting from different vertices simultaneously and stop to do some kind of merging when two trees touch. Such algorithms can be fast on many graphs, but are vulnerable to adversarial or non-random input in which case the algorithms can be forced to run sequentially. One implementation by Bader and Cong [41] that combines Prim's algorithm with Borůvka's is given here.

The algorithm works by running multiple instances of Prim's algorithm with binary heaps simultaneously with different starting vertices. Every Prim instance performs sequentially as normal and colors the found vertices to its unique color when they are first added into its heap. It runs until the heap becomes empty or it extracts a vertex from its heap that is colored by another processor or has any neighbor that is.

After all Prim instances are stopped, we add the found trees to the MST. For every vertex that is not removed from the heap by any Prim instance, we also add its shortest outgoing edge to the MST. Then we shrink the components induced by the MST edges to super-vertices in parallel. Super-vertices with no outbound edges are removed because they represent connected components of the original graph.

After the shrinking we start over again with multiple Prim instances. The whole process is repeated until the number of remaining super-vertices goes below a threshold, at which point we find its MST by executing Prim's algorithm sequentially.

Since every processor running a Prim instance may visit different number of vertices during an iteration, load-imbalance may occur. This is remedied with a work-stealing technique. They also noted that the algorithm may make no progress during an iteration for very special graphs. This can be solved if vertices are shuffled in advance. A definite worst-case for this algorithm is star-shaped graph where almost no progress can be made in the Prim instances and they loop through all neighbors of a vertex by a single processor, essentially sequentializing the algorithm. This problem is not easily solvable by work-stealing in their algorithm as stated.

### 2.3.4 Kruskal's algorithm

Of the two main phases of Kruskal's algorithm, sorting is very well parallelizable (see e.g. [38]). In the contrary, the second part is hard to scalably parallelize because whether to accept or to reject an edge depends on the decisions made earlier. Known implementations can be forced to run sequentially by an adversary and will do more work than the sequential implementation. However, a good speedup on graphs that arise in practice is still achievable. One such practical implementation is given by Blelloch et al. [26] and included in the PBBS.

Blelloch et al.'s algorithm utilizes a technique called deterministic reservation. First, the following structures and operations are introduced as building blocks of the algorithm:

Priority write. For a memory cell $x$, a priority write, denoted $x$. pwrite $(v)$, sets the value in $x$ to $v$ if $v$ is smaller than the original value stored in $x$ (thus having a higher priority). If multiple processors perform this operation on the same memory cell simultaneously, the memory cell will contain the smaller of the original value stored in the cell and the smallest value of all those write operations. On modern shared-memory architectures, this operation can be implemented with the atomic compare-and-swap (CAS) instruction within a loop. More on that is given later in Section 3.

Priority reserve. A data type (called a reservation station) is introduced which supports three operations: priority reserve ( $\chi$.reserve $(p)$ ), check $(x . \operatorname{check}(p))$ and check-and-release ( $x . \operatorname{checkR}(p)$ ), where $p$ is conceptually the priority of the operation. $x . r e s e r v e(p)$ reserves the memory location $x$ with priority $p$. This reservation fails (or will be canceled) if other reservations with higher priority have reserved (or should reserve later) the same location. The function returns whether the reservation was successful. $x . \operatorname{check}(p)$ checks if $x$ is reserved with priority $p$. $x . \operatorname{checkR}(p)$ checks if $x$ is reserved with priority $p$ and cancels the reservation if so. Cancellation is done by storing a sentinel priority $\perp$ to the cell to denote the cell is now not reserved. This value is also used to initialize the data type.

We now describe their algorithm. Edges are first sorted into nondecreasing order using a parallel sample sort. An array of reservation stations is initialized with $\perp$. A union-find data structure for the $n$ vertices is initialized to contain singletons. The algorithm takes a prefix of the array of edges, and for an edge between $u$ and $v$ of index $i$ in the sorted order, it tries to reserve stations $u$ and $v$ with priority $i$ if $u$ and $v$ do not belong to the same connected component according to the current union-find structure. This can be done with a parallel for-loop. After all reservations are submitted, a commit step performs a parallel loop over that prefix to check if the respective edge successfully reserved at least one of $u$ and $v$. If so, it releases the reservation on the other station, marks the edge as an MST edge, and merges the components of $u$ and $v$ as in the normal Kruskal's algorithm by linking the smaller of $u$ and $v$ to the larger. This ensures no cycle is formed. Note that we lose the benefit of union-by-size by doing this. After that, a prefix-sum is used to move the unsettled (neither discarded nor successfully committed) edges together to the front of the remaining unprocessed edges and a new iteration is started with another prefix of the same length of the unsettled edges and some new edges.

Another optimization mentioned before can also be used. Namely we only choose a small number (say $\frac{4}{3} \mathfrak{n}$ ) of the lightest edges and run the algorithm on this reduced list of edges. After that we filter out the heavy edges with respect to the current spanning forest and run the algorithm on the remaining edges. They reported a relative speedup of 18 and absolute speedup of 10 (against an optimized serial implementation of Kruskal's algorithm with the same filtering optimization) on a machine with 32 cores ( 64 threads with hyper-threading). This algorithm is used as the main rival of the new algorithm introduced in Section 3.

Another parallel algorithm based on Kruskal's algorithm is the Filter-Kruskal algorithm described in Section 2.2.3. The parallelism resides in the base case where $\mathfrak{m}=\mathcal{O}(\mathfrak{n})$, where a parallel sorting algorithm can be used, the partitioning and the filtering. In fact, the base case can be any other parallel MST algorithm as long as it maintains the global union-find data structure in the Filter-Kruskal algorithm. This algorithm with the above parallel Kruskal's
algorithm of PBBS as the base case is also included in the experiments in Section 4.
Katsigiannis et al. [45] tried to speed up Kruskal's algorithm using helper threads that discard heavy edges on cycles while the main threads executes the normal Kruskal's algorithm. The edges are still considered by the main thread in non-decreasing order. Helper threads examine the edges that have not yet been checked by the main thread and mark discarded the edges forming cycles in the present forest with help of the union-find data structure. The main thread first consults an boolean array that the helper threads write to to see whether the edge is already discarded before checking for cycle itself. Because multiple helper threads are running, one may hope that many edges are already discarded when the main thread comes to those edges. Though they reported a speedup of 5 on 8 cores on some synthetic graphs, it can be seen this algorithm does not scale well as the number of threads goes up because of low utilization of the helper threads. In fact, it even demonstrated slow-down when the number of helper threads increases to some (not very high) point.

Apart from parallel implementations of originally sequential algorithms, there are also algorithms designed to be parallel, especially on PRAM models. We give an incomplete summary of the results in Table 2.1. Many of these PRAM algorithms share strong similarities with the classical sequential algorithms, especially Borůvka's algorithm.

A notable example is due to Awerbuch and Shiloach [46]. The algorithm exploits a variant of the CRCW-Priority PRAM model where the priority of processors are not defined by their IDs, but by another fixed attribute. The algorithm needs one processor for each edge and one for each vertex. That priority attribute of the processors representing edges is edge weight. Each iteration of the algorithm is also roughly divided into find-min, grafting and breaking symmetry, and shortcutting where the find-min is implicitly performed by exploiting the concurrent write capability. The difference from Algorithm 1 is that the shortcutting step only does one linking step for each vertex instead of a full compression of the trees. The number of iterations can still be proven to be $\mathcal{O}(\log n)$. Zaroliagis [47] modifies this algorithm by incorporating the partitioning idea in Yao's algorithm and achieves the same depth within the CRCW-Common PRAM model. The algorithm is more complicated and calls the above algorithm to reduce the number of vertices for sparse ( $m<n \log ^{2} n$ ) graphs. It also makes use of a number of other algorithms for different tasks, notably a algorithm that simulates CRCW-Priority PRAM on CRCW-Common PRAM.

Both algorithms and most of the other algorithms in Table 2.1 do not permit easy efficient implementation on real-world machines because simulating CRCW PRAM on computers is generally not practical, and the partitioning would incur large constant factor anyway. The EREW ones are often complicated. In retrospect, the ideas in the present thesis (priority writes) might be applicable to implement these algorithms on real computers. This could be a direction for future work.

There are also algorithms designed for other architectures. For example, [48, 49, 50, 51, 42] aim at GPUs and $[52,53,54]$ at distributed-memory architectures. These are out of the scope of this thesis.

| Algorithm | Depth - Work | Note* |
| :---: | :---: | :---: |
| Chin et al. [55], 1982 | $\mathcal{O}\left(\log ^{2} n\right)-\mathcal{O}\left(n^{2}\right)$ | EREW |
| Cole and Vishkin [56], 1986 | $\mathcal{O}(\log \mathfrak{n})-\mathcal{O}((m+n) \log \log \log \mathfrak{n})$ | CRCW-P |
| Awerbuch and Shiloach [46], 1987 | $\mathcal{O}(\log n)-\mathcal{O}((m+n) \log n)$ | CRCW-P |
| Karger [57], 1992 | $\mathcal{O}(\log \mathfrak{n})-\mathcal{O}\left(m+n^{1+\varepsilon} \log n\right)$ | EREW |
| Johnson and Metaxas [58], 1992 | $\mathcal{O}\left(\log ^{3 / 2} n\right)-\mathcal{O}\left((m+n) \log ^{3 / 2} n\right)$ | EREW |
| Cole et al. [59], 1994 | $\mathcal{O}\left(2^{\log ^{*} n} \log n\right)-\mathcal{O}(m+n)$ | CRCW-A, <br> randomized |
| Cole et al. [60], 1996 | $\mathcal{O}(\log \mathfrak{n})-\mathcal{O}(m+n)$ | CRCW-A, randomized |
| Poon and Ramachandran [61], 1997 | $\mathcal{O}\left(2^{\log ^{*} n} \log n \log \log n\right)-\mathcal{O}(m+n)$ | EREW, <br> randomized |
| Zaroliagis [47], 1997 | $\mathcal{O}\left(\log ^{2} \mathfrak{n}\right)-\mathcal{O}((m+n) \log \mathfrak{n})$ | EREW |
| (same as above) | $\mathcal{O}(\log n)-\mathcal{O}((m+n) \log n)$ | CRCW-C |
| Pettie and Ramachandran [62], 1999 ${ }^{\text {a }}$ | $\mathcal{O}(\log \mathfrak{n})-\mathcal{O}(m+n)$ | EREW, <br> randomized |
| Chong et al. [63], 2001 | $\mathcal{O}(\log n)-\mathcal{O}((m+n) \log n)$ | EREW |
| Chong et al. [64], 2003 | $\mathcal{O}(\log \mathfrak{n})-\mathcal{O}((m+n) \sqrt{\log n})$ | EREW |
| (same as above) | $\mathcal{O}(\log \mathfrak{n})-\mathcal{O}((m+n) \log \log \mathfrak{n})$ | CRCW-A |

Table 2.1: MST algorithms on PRAM.

* CRCW-P stands for CRCW-Priority, CRCW-A for CRCW-Arbitrary, CRCW-C for CRCW-
Common.
A preliminary version of the work appeared in 1999. The referenced version was published
in 2002 .


## 3 The New Algorithm

In this section we describe a new parallel algorithm based on Borůvka's algorithm. The main steps of the algorithm do not differ from the original Borůvka's algorithm: we still have an outer loop that keeps invoking Borůvka step until we have a single connected component. The Borůvka step still has three main steps, i.e. find-min, grafting and shortcutting as in Algorithm 1. We first describe the necessary building blocks of the algorithm and then describe the new algorithm incorporating these building blocks.

### 3.1 Priority Write

A main ingredient of the new algorithm is priority write or priority update. This primitive is used in Blelloch et al.'s parallel implementation of Kruskal's algorithm and briefly described in Section 2.3.4. Priority write of the value $v$ to a memory cell identified by $x$ (thus $x$ is a pointer or address) is denoted $x$. pwrite $(v)$ or pwrite $(x, v)$. The operation replaces the value at the destination of $x$ with $v$ if $v$ is smaller than the original value stored there. More formally, the operation is functionally equivalent to Algorithm 3:

```
Algorithm 3: Priority Write
    Function pwrite(x: pointer to value, v: value)
        if \(* x>v\) then \(/ *{ }^{*} * x\) " is the value stored in the destination of \(x * /\)
            \(* x \leftarrow v\)
```

Problems arise when this operation is used in parallel settings. When multiple processors do this operation simultaneously, data race will occur. Take the following processor interleaving as an example where $* x$ is initialized to 5 and Processor 1 and 2 call pwrite $(x, 3)$ and pwrite $(x, 1)$, respectively:

```
// *x is initialized to 5.
Processor 1: if 5 > 3 then // true
Processor 2: if 5 > 1 then // true
Processor 2: *x <- 1 // *x is now 1
Processor 1: *x <- 3 // *x is now 3
// *x = 3 at the end
```

Here *x is incorrectly set to 3 after both pwrites. We therefore require pwrite to be atomic. On modern shared-memory architectures, pwrite is not readily available but can be easily implemented with atomic compare-and-swap (CAS) instructions. A CAS operation is functionally equivalent to Algorithm 4, but guaranteed to be atomic by the hardware (see
e.g. [65]):

```
Algorithm 4: Compare-and-Swap
    Function cas(x: pointer to value, old: value, new: value): boolean
        if \(* x=\) old then
            \(* x \leftarrow\) new
            return true /* success */
        else
            return false /* failure */
```

With help of CAS instruction, the priority write operation can be implemented free of data race as follows in Algorithm 5:

```
Algorithm 5: Priority Write with Compare-and-Swap
    Function pwrite(x: pointer to value, v: value)
        do
            old \(\leftarrow * x\)
        while \(v<\) old and cas \((x\), old,\(v)=\) false /* short-circuiting in effect */
```

This piece of code repeatedly reads the current value of $* x$ to a variable old, checks if the intended new value $v$ is still smaller than old, and does a CAS to store $v$ into $* x$ if it is. The loop ends when at some point the newly read value old is no larger than the intended new value $v$ or a CAS has succeeded. Note that the second condition is checked only if the first is true because of short-circuiting. From now on, pwrite denotes this version of priority write.

Priority writes may be defined for any value type with a total ordering. Due to hardware limitation on atomic compare-and-swap instruction, however, the size of supportable value types is normally restricted to that the largest supported primitive integral type, e.g. 16 bytes on current mainstream 64 -bit processors (see e.g. [65, 66]).

Priority writes have been first introduced in [26] and extensively analyzed by Shun et al. [67]. They may seem very inefficient at first glance due to the presence of the loop and CAS instruction. More careful reasoning reveals that the CAS (and a potential write) only occurs when the present value in $* x$ is larger than the intended new value and the loop is broken immediately when this is not the case. Therefore the performance of the primitive depends on how often a new value is written by the CAS instruction by all processors. Intuitively, this does not happen all that often because a written value will prevent many future writes. In fact, if every of $p$ processors attempts to call pwrite $\left(x, v_{i}\right)$ where $v_{i}$ is drawn uniformly at random from a range or $v_{i}$ is the $\mathfrak{i}$-th value in a random permutation of values, the expected number of writes is only $\ln p+\mathcal{O}(1)$. That is because a random permutation of length $p$ only has $H_{p}:=\sum_{i=1}^{p} \frac{1}{i}=\ln p+\mathcal{O}(1)$ prefix-minima in average with high probability, where $H_{p}$ is the $p$-th harmonic number [68]. This is also the expected running time if $p$ processors write to a single location with random values at the same time. The worst case happens when the values are written in decreasing order. This is extremely unlikely because that would mean the CAS supporting hardware somehow orders all writes in the hardware queue in descending order. Because it is the operating system that schedules concurrent processes invoking pwrite, and perfect synchronization, which does not exist in real world, would be needed for an adversary to forge such a situation, we may safely assume the expected running time is $\mathcal{O}(\ln p)$ regardless of the input. If we use $p$ processors to make $n$ pwrites and we assume the $n$ operations are divided into batches of size $p$ as is roughly the
case with parallel for-loop, each of the batch takes $\mathcal{O}(\log \mathfrak{p})$ time. Thus the expected total time for all operations is $\mathcal{O}\left(\frac{n}{p} \log p\right)$. Shun et al. [67] gave a result of $\mathcal{O}\left(\frac{n}{p}+c \log n+c p\right)$ where $c$ is a constant characterizing the cost of the ensuring cache-coherence.

The situation becomes more obscure when $p$ processors execute $n$ pwrites to $m$ different locations. If the pwrites are made to random locations, we can expect the contention on a single cell to reduce. The exact running time seems to be open at this time. Shun et al. [67] gave a bound of $\mathcal{O}\left(\frac{n}{p}+\mathfrak{c m} \log \frac{\mathfrak{n}}{\mathfrak{m}}+(c p)^{2}\right)$. This result seems unfavorable, especially because of the second and third terms which do not decrease or even rise as $p$ goes up. That is probably because their result makes a pessimistic assumption that a successful CAS by a processor results in a full invalidation of the cache of all other processors incurring time cost $c$. This is pessimistic because other processors do not have to be aware of the write unless they want to write to the same location. Despite the imperfect theoretical bound, they demonstrated good performance in practice, as we will also see later in the context of the new MST algorithm.

Note that there is a similarity between priority writes and the PRAM model with concurrent writes by priority (CRCW-Priority). The difference is that the argument defines the priority of a pwrite operation and not the processor ID which is the case with CRCW-Priority. Therefore pwrite is intuitively more flexible. On the other hand, CRCW-Priority can simulate pwrite, too, by first sorting the sequence of operations by their priorities and let the processor with the ID corresponding to the rank of the arguments perform the write. This, however, incurs extra cost due to the sorting.

### 3.2 Compaction

The new algorithm is based on Borůvka's algorithm, therefore it also has a shortcutting step that compacts the found connected components into super-vertices. We discuss this compaction step in this section.
Definition 3.1 (compaction). Formally, the compaction operation receives the array of parents $\mathrm{R}[0 . . \mathrm{n}-1]$ as input and is supposed to shortcut the trees implicitly defined by the values so that at the end of the compaction process, every vertex $\mathfrak{i}$ is either a root (characterized by $R[i]=i)$ or is the child of a root $(R[R[i]]=R[i])$.

This can be solved in parallel on CREW PRAM with a pointer jumping (or path doubling) technique within $\mathcal{O}(\log \mathfrak{n})$ depth and $\mathcal{O}(n \log \mathfrak{n})$ work [38] as shown in Algorithm 6.

```
Algorithm 6: Pointer Jumping
    Input: Parent array R[0..n-1]
    Output: Compacted parent array: a node is either a root or a child of a root
    foreach \(0 \leqslant \mathfrak{i}<\mathrm{n}\) do in parallel
        while \(R[i] \neq R[R[i]]\) do \(\quad / *\) if \(i\) 's parent is not i's grandparent */
            \(R[i] \leftarrow R[R[i]] \quad / *\) graft \(i\) to its grandparent */
```

Note that for parallel loops on PRAM, program code within the loop body is run by all processors simultaneously with perfect synchronization. That means all processors run line 2 at the same time and, after all processors are done checking the condition, all processors for which the condition holds run line 3 simultaneously. The synchronization is crucial for the correctness
of the theoretical time bound $\mathcal{O}(\log \mathfrak{n})$. Indeed, consider a chain: if we let all processors wait until the processor representing the leaf node finally reaches the root, the time is already $\Omega(n)$.

There are more efficient algorithm for this compaction problem that only needs $\mathcal{O}(\mathrm{n})$ operations in total (see e.g. [38]), but it turns out that this step only takes a marginally small proportion of the time of our MST algorithm, so we do not go further optimizing it. In fact, we may even run line 2 and line 3 without synchronization in practice because the operating system normally schedules processes more or less evenly so that every process benefits from the shortcutting work of other processes thanks to cache-coherency protocols. This way, we can also relieve ourselves of costly locks.

### 3.3 The Algorithm

On a high level, the new algorithm described here does not differ much from the sequential Borůvka's algorithm given in Algorithm 2. The difference resides in how the Borůvka step is implemented. Assuming the same edge-list representation, we run through every edge in a parallel loop in the Borůvka step to determine the best edge for every current super-vertex, namely the representative of its current connected component. This is done by performing a priority write to each of the endpoints of every edge with the weight and ID as the argument. By the definition of priority writes, we will have the best edges for every connected component. After that we graft trees along the best edges, breaking symmetry just like in the sequential version, but this time in parallel. After that we perform a full compaction. A new set of active super-vertices is found with a filter operation. To improve locality of memory accesses, we change the endpoints of every edge to their corresponding representatives. Self-edges are filtered out at the end of the Borůvka step.

Now we describe our MST algorithm using the edge-list representation in detail. We say a tuple $\left(a_{1}, a_{2}, \ldots, a_{k}\right)$ is smaller than another tuple $\left(b_{1}, b_{2}, \ldots, b_{k}\right)$ if there is an $i, 1 \leqslant i \leqslant k$, so that $\forall_{1 \leqslant j<i}: a_{j}=b_{j}$ and $a_{i}<b_{i}$. This also known as the lexicographical order. Two edges can be compared according to the lexicographical order of their corresponding (weight, index) pair, where index is the position of the edge in the original edge list. Operator "par " stands for parallel assignment, i.e. all available processors divide the intended assignments evenly and execute their own portion in parallel. The full algorithm in given in Algorithm 7. Note that before every Borůvka step, V always only holds vertices that are representatives of their components.

As we can see, the algorithm does not differ much from the sequential implementation in Algorithm 1 and Algorithm 2. The first main difference is the use of pwrite in the find-min step in line 7 and line 8 . Conceptually, those pwrites try to store the edge into the locations for both its endpoints in best array. This solves the load-imbalance issue with known implementations which loop through all vertices in a parallel but process all edges for a vertex sequentially. Since CAS instructions and thus pwrites have limit on the size of the operands, we actually write the pair (weight, index) of the edge with pwrite. This is well suitable for double-precision or 64-bit integral weights (both are 8 bytes) and 4 to 8 -byte indices since most of current 64 -bit architectures support CAS instructions on 16 -byte operands. This is also better than only storing the index because we save two random memory accesses when comparing an edge with the two present in the best array (for both endpoints).

The grafting step almost stays the same except that it is now parallel. If best $[i]=$ sentinel

```
Algorithm 7: New Parallel MST/MSF Algorithm
    Input: Graph \(\mathrm{G}=(\mathrm{V}, \mathrm{E})\)
    Output: The MST of G
    begin
        \(\mathrm{R}[\mathrm{i}] \stackrel{\mathrm{par}}{\leftrightarrows} \mathfrak{i}, \mathfrak{i} \in \mathrm{V}\) /* initialization */
        MST \(\leftarrow \varnothing \quad / *\) all the MST/MSF edges */
        while \(|\mathrm{V}|>1\) and \(|\mathrm{E}|>0\) do
            /* loop invariant: V only contains root vertices. */
            /* loop invariant: E contains no self-loops. */
            best \([\mathrm{i}] \stackrel{\mathrm{par}}{\rightleftarrows}\) sentinel, \(\mathfrak{i} \in \mathrm{V} \quad / *\) sentinel is an edge of weight \(\infty\) */
            /* find-min step */
            foreach \(e \in E\) do in parallel
            best[e.u]. pwrite(e)
            best[e.v].pwrite(e)
            /* best[i] is now the lightest edge leaving i's component;
            best \([i]=\) sentinel if there is none. */
            /* grafting step */
            foreach \(\mathfrak{i} \in \mathrm{V}\) do in parallel
            index \(\leftarrow\) index of \(\mathfrak{i}\) in the current V
            if best \([i]=\) sentinel then \(\quad / *\) no edge found for component \(i * /\)
                \(\mathrm{R}[\mathrm{i}] \leftarrow-1 \quad / *\) inactivate that component */
                mst_edge \([\) index \(] \leftarrow\) sentinel
            else
                \(\mathrm{j} \leftarrow\) the other endpoint of edge best \([\mathrm{i}]\)
                if best \([i]=\) best \([j]\) and \(i<j\) then
                    Do nothing with \(R[i] \quad / *\) break symmetry; \(R[i]\) stays \(i\) */
                    mst_edge \([\) index \(] \leftarrow\) sentinel
                else
                    \(R[i] \leftarrow j \quad / * \operatorname{graft} i\) to \(j\) */
                    mst_edge \([\) index \(] \leftarrow\) best \([i]\)
            MST \(\leftarrow\) MST \(+\{\) mst_edge \([j] \neq\) sentinel \(|0 \leqslant \mathfrak{j}<|\mathrm{V}|\} \quad / *\) filtering */
            /* shortcutting step

Perform pointer jumping (Algorithm 6) on \(R\) for vertices in \(V\) to compact trees.
            /* relabeling step */
            \(\mathrm{E} \leftarrow\{(\mathrm{R}[e . u], \mathrm{R}[e . v], e . w) \mid e \in \mathrm{E}\}\)
            /* filtering step */
            \(E \leftarrow\{e \in E \mid e . u \neq e . v\}\)
                            /* filtering */
            \(V \leftarrow\{i \in V \mid R[i]=i\} \quad / *\) new set of vertices; filtering */
        return MST
for some vertex \(\mathfrak{i}\) (recall that \(\mathfrak{i}\) is always a representative), that means the component of \(\mathfrak{i}\) did not get any edge in the find-min step. Thus we inactivate the component by setting the parent of \(\mathfrak{i}\) to a negative value in line 12. If we graft a component to another, we also mark the edge we just used to be in the MST (line 21). Subsequently, all those marked edges are added to the set of MST edges in line 22 with a parallel filtering.

As in the sequential case, we perform a shortcutting step to make every vertex either a root or a child of a root in line 23 . Note we should only do this for roots vertices that are not inactivated in line 12 (i.e. those with \(R[i] \geqslant 0\) ).

In order to further improve locality in the find-min step, we relabel the endpoints of the edges by changing the endpoints of every edge to their parents in line 24 . This way, the find-min step in the next iteration does not have to consult the \(R\) array to determine the representative of the endpoints for an edge, saving two random memory accesses. Cong and Tanase [43] have independently come up with the same idea and applied to their variant of Borůvka's algorithm very recently.

Finally, a filtering step takes place to remove self-edges, reducing the length of the edge list and saving time in the find-min step. The vertex set is also updated to the new set of roots (thus removing exhausted components and components that are grafted to other components). The loop invariant is restored this way and wo start the loop over unless the edge set becomes empty or only a single vertex remains.

Note that this algorithm also computes the minimum spanning forest if the original graph is not connected. That is because a component is inactivated in line 12 if it is exhausted, and it will be removed from the vertex set, so it will not prevent further progress of the algorithm.

An important remark is that the parallelism in this algorithm is very coarse-grained: we are only using data parallel primitives and no locks or other manual synchronization are needed within each such construct. This is favorable because locks are too expensive to guarantee good performance.

Since the algorithm is essentially Borůvka's algorithm, we still have \(\mathcal{O}(\log \mathfrak{n})\) iterations where the number of active vertices at least halves after every iteration. In every iteration, the grafting takes \(\mathcal{O}\left(\frac{n_{i}}{p}+\min \left(p, \log n_{i}\right)\right)\) time where \(n_{i}\) is the number of vertices at the beginning of the \(\mathfrak{i}\)-th iteration and \(p\) is the number of available processors. Shortcutting takes \(\mathcal{O}\left(\frac{\mathfrak{n}_{i} \log n_{i}}{\mathfrak{p}}\right)\) time if we implement pointer jumping with synchronization. In the practice, however, we do not have to (and want to) use expensive locks to ensure synchronization, as discussed in Section 3.2. The parallel filtering and parallel for-loops take time \(\mathcal{O}\left(\frac{m_{i}}{p}+\min \left(p, \log m_{i}\right)\right)\) in total, where \(\mathfrak{m}_{\mathfrak{i}}\) is the number of edges at the beginning of the iteration. The running time of the find-min step is still open as discussed in 3.1, but we conjecture it to be expected \(\mathcal{O}\left(\frac{\mathfrak{m}_{\mathrm{i}} \log \mathfrak{m}_{\mathrm{i}}}{\boldsymbol{p}}\right)\). In practice it runs very fast and is invulnerable to adversarial inputs. Summing up all terms, noticing \(n_{i+1} \leqslant \frac{n_{i}}{2}\), the MSF algorithm is conjectured to finish in expected \(\mathcal{O}\left(\frac{n \log n+m \log m \log n}{p}\right)\) time regardless of input, assuming \(n \gg p\). The \(\frac{m \log m \log n}{p}\) term looks too large and does not seem to match the performance in practice. It is of theoretical interest to prove the bound or even a better one, but in practice, it appears that we can safely use pwrite without much thoughts on performance. Note that if the algorithm is run with a single thread, it degenerates to normal Borůvka's algorithm. The running time in that case becomes deterministically \(\mathcal{O}(\mathrm{m} \log \mathfrak{n})\) because each Borůvka step does no more than \(\mathcal{O}(\mathfrak{m})\) work. This makes the algorithm favorable for situations where we do not always have the full computing resources of the system.

Other optimizations can be applied to the algorithm to improve the performance even further for some graphs. One such optimization has been introduced in Section 2.3.4, namely we can run the algorithm on the lightest portion of all the edges, run a filtering on the remaining edges with help of the acquired information during the first run and run the algorithm for a second time on the edges surviving the filtering. Finding the lightest edges can be done with ideas similar to a parallel sample sort: we take a sample of all edges, sort them with a sample sort, and take the respective value in the sample according to the portion we want, and do a partitioning/splitting with that value as the pivot. The lightest edges are thereby moved to the beginning of the array. Note that we have to reactivate the inactivated components after the first run by setting \(R[i] \leftarrow i\) for all \(i\) with negative \(R[i]\) because they are inactivated only because we exhausted edges in the lightest portion. Cong and Tanase [43] have gone even further by partitioning the edges into many buckets (instead of two) where edges in earlier buckets are lighter than all edges in later buckets and always doing a filtering on the edges in a bucket before running their MST algorithm on it.

\section*{4 Experimental Results}

We have introduced the new algorithm for computing minimum spanning trees in Section 3. In this section, we compare the new algorithm with some of the existing sequential and parallel algorithms introduced in Section 2 on synthetic and real-world graphs and present experimental results that demonstrate the practical performance of our new algorithm.

\subsection*{4.1 Benchmark Configuration}

Experiments are conducted on a workstation with the following technical characteristics:
\begin{tabular}{l|l}
\hline Name & Value \\
\hline OS & Ubuntu 14.04.5 64-bit \\
CPU & Intel(R) Xeon(R) CPU E7-8867 v4 @ 2.40 GHz \\
Sockets & 4 \\
Cores per socket & 18 \\
Threads per core & 2 \\
Cache line size & 64 bytes \\
Cache alignment & 64 -byte boundary \\
L2 cache size & \(256 \mathrm{~KB} /\) core \\
L3 cache size & \(45 \mathrm{MB} /\) socket \\
RAM & 1 TB \\
\hline
\end{tabular}

Table 4.1: System specifications.

The test programs and benchmark are based the Problem Based Benchmark Suite (PBBS) [69]. It is a framework that offers various data parallel primitives and can be used to compare different solutions to the same problem by their performance. Correctness can be checked against a reference implementation in a black-box manner.

The following implementations are included in the benchmark:
seq_pbbs_kruskal. A sequential implementation of Kruskal's algorithm provided in PBBS. It uses union-find with path-compression and union-by-size to maintain the current components.
seq_pbbs_filtering_kruskal. Same as above, but it uses the filtering technique described in Section 2.3.4 and Section 3, i.e. we first uses a partitioning algorithm to get the lightest edges, run Kruskal's algorithm on it, filter out self-edges with the current union-find data structure, and execute once more Kruskal's algorithm on the remaining edges.
par_pbbs_kruskal. The parallel Kruskal's algorithm provided in PBBS, introduced in Section 2.3.4, without filtering.
par_pbbs_filtering_kruskal. Same as above but with filtering.
par_filter_kruskal. An implementation of the recursive Filter-Kruskal algorithm described in [24] and briefly introduced in Section 2.2.3. The parallel Kruskal's algorithm with filtering from PBBS serves as the base case of the algorithm and the needed partition and filter primitives are provided by PBBS.
seq_prim_binary. An implementation of Prim's algorithm with binary heaps.
seq_prim_pairing. An implementation of Prim's algorithm with pairing heaps. The pairing heaps are from the Policy-Based Data Structures included in the GNU C++ compiler [70].
seq_boruvka. An implementation of Borůvka's algorithm given in Algorithm 2. par_boruvka_d. A variant of Borůvka's described in Section 2.3.2 and [42]. Small changes were made to make it suitable for the PBBS framework including support for double-precision weights and graph format conversion. Time for conversion is not counted towards its running time.
par_new_boruvka. An implementation of the new Borůvka's algorithm presented in this thesis.
par_new_filtering_boruvka. Same as above but with filtering.
All source codes are written in C++ and are compiled using GNU C++ compiler Version 5.4.1 with relevant compilation flags -03 -march=native. Parallelization is generally achieved with the Cilk++ [71] except for par_boruvka_d, which uses OpenMP and partly Intel Threading Building Blocks (TBB) for parallelization.

Various synthetic and real-world graphs are used in the benchmarks. These are listed below:
randLocal_20M. A random local graph with 20 million vertices and degree 5 for each vertex. The 5 edges for each vertex is chosen uniformly at random. This is generated with the graph generation utility provided in PBBS.
rMat_20M. A graph generated with the Recursive Matrix ( \(R\)-MAT) algorithm proposed in [72]. The algorithm models real-world graphs like social networks nicely and produces ones with small diameter and power-law degree distributions. This is generated with the graph generation utility provided in PBBS.
2Dgrid_20M. A regular square 2-D grid with 20 million vertices. The side length, i.e. the number of vertices on a side, is therefore \(\sqrt{20}\) million. The vertices on the borders are also adjacent to their counterparts on the other side of the grid. This is generated with the graph generation utility provided in PBBS.
3Dgrid_20M. A regular 3-D grid with 20 million vertices with side length \(\sqrt[3]{20 \text { million. }}\) The same edge-wrapping as above is also present. This is generated with the graph generation utility provided in PBBS.
stars_20M. A graph consisting of stars with 20 million edges. This is generated with the graph generation utility provided in PBBS.
chain_20M. A graph containing a single chain of increasing 20 million vertex IDs and edge weights. This is generated with the graph generation utility provided in PBBS.
delaunay_20M. A graph generated by randomly scattering 20 million points onto the unit square and building the Delaunay triangulation of the point set. The number of edges is roughly 60 million because Delaunay triangulations are in a sense planar graphs with the maximum number of edges and connected simple planar graphs never have more than \(3 n-6\) edges (see e.g. [73]). The edge weights are the Euclidean distances of the endpoints. This graph is generated with MathWorks Matlab [74].
delaunay_20M-n. Same as above, but \(n\) (which is 20 million) random edges of the triangulation are removed. This is to model a planar graph of "half fullness".
delaunay_20M-2n. Same as above but with 2 n edges removed. The resulting graph
only has about \(n\) edges.
delaunay3d_10M. Similar to delaunay_20M-n, but now 10 million points instead of 20 are drawn from a unit cube and the edges are those formed by a threedimensional Delaunay triangulation.
delaunay3d_10M-2n. Same as above, with \(2 n\) edges removed.
delaunay3d_10M-4n. Same as above, with \(4 n\) edges removed. Note the resulting graph still has about 40 million edges since the graph is not planar anymore and hence does not satisfy the same \(3 n-6\) upper bound as in the two-dimensional case.
uniform_20M_20M. A graph uniformly drawn from the universe of all graphs of 20 million vertices and 20 million edges. The generation is done with the readily available functions RandomGraph and UniformGraphDistribution of Wolfram Mathematica [75].
uniform_2M_20M. Same as above but with 2 million vertices and 20 million edges, so that the density of the graph slightly increases.
uniform_200K_20M. Same as above but with only 200 thousand vertices.
uniform_20K_20M. Same as above but with only 20 thousand vertices.
nlpkkt240. A graph from the SuiteSparse Matrix Collection, formerly The University of Florida sparse matrix collection [76].
USA. A real-world road network of the USA, provided by the 9th DIMACS Implementation Challenge [77]. Weights are the physical distances.
livejournal. A graph taken from the Stanford Network Analysis Platform (SNAP) [78] representing the friendship network of the LiveJournal social network.
Unless noted otherwise, the edges weights in the graphs are uniformly random. Table 4.2 summarizes the used graphs, their sizes and the final MST (MSF) edges.

All time measurements are calculated by executing the respective implementations ten times and taking the average of the running times after removing the minimum and the maximum.
\begin{tabular}{l|r|r|r}
\hline Name & Vertices & Edges & MST edges \\
\hline randLocal_20M & 20000000 & 100000000 & 19999999 \\
rMat_20M & 33554432 & 100000000 & 29355409 \\
2Dgrid_20M & 19998784 & 39997568 & 19998783 \\
3Dgrid_20M & 19902511 & 59707533 & 19902510 \\
stars_20M & 20000004 & 20000000 & 20000000 \\
chain_20M & 20000000 & 19999999 & 19999999 \\
delaunay_20M & 20000000 & 59999950 & 19999999 \\
delaunay_20M-n & 20000000 & 39999943 & 19932186 \\
delaunay_20M-2n & 20000000 & 19999961 & 17214746 \\
delaunay3d_10M & 10000000 & 77586968 & 9999999 \\
delaunay3d_10M-2n & 10000000 & 57587804 & 9999991 \\
delaunay3d_10M-4n & 10000000 & 37593541 & 9997509 \\
uniform_20M_20M & 20000000 & 20000000 & 16762252 \\
uniform_2M_20M & 2000000 & 20000000 & 1999999 \\
uniform_200K_20M & 200000 & 20000000 & 199999 \\
uniform_20K_20M & 20000 & 20000000 & 19999 \\
\hline nlpkkt240 & 27993601 & 746478752 & 27993599 \\
USA & 23947347 & 28854312 & 23947346 \\
livejournal & 4036538 & 34681189 & 3997961 \\
\hline
\end{tabular}

Table 4.2: Sizes of the graphs included in the benchmark and the number of MST edges in these graphs. If the number of MST edges is smaller than \(|\mathrm{V}|-1\), the value is then the number of MSF edges.

\subsection*{4.2 Benchmark Results}

The results of the benchmark are presented in this subsection. We first list the graphics of the benchmarks and then provide a detailed analysis. The \(x\)-axis of the graphics is always the number of used threads, and \(y\) the speedup over the fastest sequential implementation for that particular graph. Note that the \(y\)-axis has an unusual scaling to enhance the contrast below the \(y=1\) line.

For a more precise reference, the running times for 1 and 144 threads are also tabulated after the graphics in Table 4.3 and Table 4.4. The relative and absolute speedups are given in Table 4.5.

\subsection*{4.2.1 Graphics and Tables for the Experiments}


Figure 5: Benchmark results for randLocal_20M graph. The lines for seq_boruvka and seq_prim_pairing overlap.


Figure 6: Benchmark results for rMat_20M graph.


Figure 7: Benchmark results for 2Dgrid_20M graph.

3Dgrid_20M


Figure 8: Benchmark results for 3Dgrid_20M graph. The lines for seq_boruvka, seq_prim_binary and seq_prim_pairing are closely next to each other.


Figure 9: Benchmark results for stars_20M graph. Prim's algorithm performs very poorly on this graph because it needs \(\mathcal{O}(n \log n)\) time to execute \(n\) inserts or decrease-keys. Par_boruvka_d is excluded because the code the authors provide online [42] seems to contain a bug and did not terminate for stars.


Figure 10: Benchmark results for chain_20M graph. Kruskal's algorithm works badly here. The lines for seq_boruvka and seq_prim_binary overlap and these for seq_pbbs_kruskal and seq_pbbs_filtering_kruskal overlap.


Figure 11: Benchmark results for delaunay_20M graph.
delaunay_20M-n


Figure 12: Benchmark results for delaunay_20M-n graph.


Figure 13: Benchmark results for delaunay_20M-2n graph. Kruskal's algorithm, with or without filtering, is the fastest sequential algorithm.
delaunay3d_10M


Figure 14: Benchmark results for delaunay3d_10M graph.


Figure 15: Benchmark results for delaunay3d_10M-2n graph.


Figure 16: Benchmark results for delaunay3d_10M-4n graph.


Figure 17: Benchmark results for uniform_20M_20M graph. Kruskal's algorithm with and without filtering share the same straight line as the fastest sequential algorithms.
uniform_2M_20M


Figure 18: Benchmark results for uniform_2M_20M graph.


Figure 19: Benchmark results for uniform_200K_20M graph. Seq_pbbs_kruskal is slow for this graph because of the full sorting, but it works better with filtering.
uniform_20K_20M


Figure 20: Benchmark results for uniform_20K_20M graph. Seq_pbbs_kruskal is slow for this graph because of the full sorting, but filtering remedies a bit.


Figure 21: Benchmark results for nlpkkt240 graph. Kruskal's algorithm without filtering is slow because of the full sorting. With filtering, it becomes the fastest sequential algorithm.

USA


Figure 22: Benchmark results for USA graph. Lines for seq_pbbs_kruskal, seq_boruvka and seq_prim_binary overlap.

\section*{livejournal}


Figure 23: Benchmark results for livejournal graph.
\begin{tabular}{|c|c|c|c|c|c|}
\hline & \begin{tabular}{l}
seq \\
pbbs kruskal
\end{tabular} & ```
    seq
    pbbs
filtering
    kruskal
``` & \begin{tabular}{l}
seq \\
prim \\
binary
\end{tabular} & seq prim pairing & seq boruvka \\
\hline randLocal_20M & 18.300 & 11.450 & 33.041 & 46.291 & 43.638 \\
\hline rMat_20M & 18.813 & \(\underline{13.500}\) & 51.646 & 79.168 & 44.280 \\
\hline 2Dgrid_20M & 8.404 & \(\underline{6.246}\) & 16.844 & 15.138 & 22.585 \\
\hline 3Dgrid_20M & 11.963 & \(\underline{7.798}\) & 26.658 & 28.791 & 31.948 \\
\hline stars_20M & 3.285 & 3.110 & 25.281 & 47.348 & 1.891 \\
\hline chain_20M & 1.315 & 1.311 & 0.246 & 0.645 & 0.247 \\
\hline delaunay_20M & 13.113 & \(\underline{11.538}\) & 15.670 & 14.292 & 22.692 \\
\hline delaunay_20M-n & 8.535 & \(\underline{7.668}\) & 14.935 & 13.591 & 25.060 \\
\hline delaunay_20M-2n & \(\underline{3.930}\) & 4.073 & 7.905 & 8.519 & 14.501 \\
\hline delaunay3d_10M & 14.375 & 13.213 & 13.943 & \(\underline{11.987}\) & 20.353 \\
\hline delaunay3d_10M-2n & 10.325 & 8.410 & 12.267 & 11.798 & 22.965 \\
\hline delaunay3d_10M-4n & 6.828 & \(\underline{4.708}\) & 10.417 & 10.844 & 15.587 \\
\hline uniform_20M_20M & \(\underline{3.700}\) & 3.700 & 12.520 & 15.063 & 8.365 \\
\hline uniform_2M_20M & 2.986 & \(\underline{1.360}\) & 1.776 & 3.729 & 2.627 \\
\hline uniform_200K_20M & 3.050 & 0.942 & \(\underline{0.519}\) & 0.690 & 1.632 \\
\hline uniform_20K_20M & 2.730 & 0.888 & \(\underline{0.219}\) & 0.298 & 0.778 \\
\hline nlpkkt240 & 152.125 & \(\underline{45.700}\) & 53.995 & 66.692 & 59.181 \\
\hline USA & 6.086 & 5.370 & 6.159 & \(\underline{5.033}\) & 6.089 \\
\hline livejournal & 5.459 & \(\underline{2.430}\) & 4.112 & 7.656 & 4.815 \\
\hline
\end{tabular}

Table 4.3: Running times of sequential algorithms in seconds. The best time for each particular graph is underlined.
\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|c|c|c|}
\hline &  & \begin{tabular}{l}
par \\
pbbs \\
filtering \\
kruskal
\end{tabular} &  &  &  & \begin{tabular}{l}
par \\
new filtering boruvka
\end{tabular} &  & \begin{tabular}{l}
par \\
pbbs filtering kruskal
\end{tabular} &  &  &  & \begin{tabular}{l}
par \\
new filtering boruvka
\end{tabular} \\
\hline \#threads & 1 & 1 & 1 & 1 & 1 & 1 & 144 & 144 & 144 & 144 & 144 & 144 \\
\hline randLocal_20M & 34.500 & 19.800 & 17.238 & 88.823 & 30.600 & 12.400 & 0.765 & 0.544 & 0.636 & 20.075 & 1.026 & 0.624 \\
\hline rMat_20M & 37.400 & 25.800 & 22.788 & 83.999 & 26.400 & 16.900 & 0.809 & \(\underline{0.663}\) & 0.735 & 18.229 & 0.887 & 0.776 \\
\hline 2Dgrid_20M & 16.563 & 13.625 & 14.288 & 30.317 & 11.025 & 8.019 & 0.396 & 0.351 & 0.458 & 2.564 & 0.389 & 0.366 \\
\hline 3Dgrid_20M & 22.513 & 15.500 & 14.988 & 44.499 & 18.000 & 9.296 & 0.503 & \(\underline{0.419}\) & 0.507 & 5.382 & 0.626 & 0.460 \\
\hline stars_20M & 8.143 & 7.890 & 7.779 & - & \(\underline{2.290}\) & 2.305 & 0.176 & 0.173 & 0.191 & - & 0.080 & 0.080 \\
\hline chain_20M & 1.623 & 1.590 & 1.836 & 1.475 & \(\underline{0.660}\) & 0.665 & 0.163 & 0.159 & 0.211 & 0.592 & 0.039 & 0.038 \\
\hline delaunay_20M & 23.688 & 20.938 & 19.688 & 39.243 & 12.813 & 11.600 & 0.541 & 0.532 & 0.582 & 3.207 & 0.417 & 0.441 \\
\hline delaunay_20M-n & 16.988 & 15.300 & 16.188 & 30.872 & 10.400 & 9.730 & 0.411 & 0.420 & 0.461 & 2.570 & 0.370 & 0.384 \\
\hline delaunay_20M-2n & 9.369 & 9.223 & 9.155 & 16.709 & \(\underline{5.333}\) & 5.355 & 0.236 & 0.233 & 0.250 & 1.822 & 0.200 & 0.200 \\
\hline delaunay3d_10M & 24.063 & 20.600 & \(\underline{\underline{9.584}}\) & 42.447 & 15.500 & 12.800 & 0.515 & 0.573 & 0.434 & 5.708 & 0.501 & 0.509 \\
\hline delaunay3d_10M-2n & 18.363 & 14.900 & 8.680 & 33.764 & 14.100 & 11.375 & 0.413 & 0.407 & 0.398 & 4.506 & 0.485 & 0.444 \\
\hline delaunay3d_10M-4n & 13.263 & 9.320 & 8.085 & 23.375 & 9.601 & 6.858 & 0.299 & 0.265 & 0.312 & 3.103 & 0.347 & 0.286 \\
\hline uniform_20M_20M & 9.369 & 9.411 & 8.620 & 18.509 & \(\underline{5.569}\) & 5.650 & 0.242 & 0.239 & 0.265 & 2.292 & \(\underline{0.207}\) & 0.210 \\
\hline uniform_2M_20M & 5.340 & 2.061 & 1.440 & 11.752 & 3.070 & \(\underline{\underline{1.016}}\) & 0.155 & 0.097 & 0.144 & 5.691 & 0.133 & 0.086 \\
\hline uniform_200K_20M & 5.170 & 0.544 & 0.566 & 8.822 & 2.210 & \(\underline{0.403}\) & 0.141 & 0.032 & 0.080 & 6.309 & 0.095 & 0.035 \\
\hline uniform_20K_20M & 4.500 & 0.268 & 0.496 & 5.013 & 1.200 & 0.259 & 0.128 & 0.024 & 0.060 & 6.395 & 0.081 & 0.026 \\
\hline nlpkkt240 & 264.375 & 89.713 & 39.988 & 278.142 & 59.100 & \(\underline{\underline{22.588}}\) & 5.423 & 2.325 & 2.046 & 72.458 & 3.129 & 1.498 \\
\hline USA & 12.213 & 12.300 & 12.488 & 11.025 & \(\underline{\underline{3.645}}\) & 4.448 & 0.329 & 0.321 & 0.358 & 2.022 & \(\underline{0.171}\) & 0.253 \\
\hline livejournal & 10.575 & 3.919 & 2.848 & 14.941 & 3.600 & \(\underline{\underline{1.830}}\) & 0.234 & 0.148 & 0.215 & 6.504 & 0.153 & 0.129 \\
\hline
\end{tabular}

Table 4.4: Running times of parallel algorithms with 1 and 144 threads in seconds. Underlined numbers mark the best times for the particular graph and particular number of threads. If the time for a parallel algorithm executed with one thread is even faster than the best sequential algorithm, that time is doubly underlined.
\begin{tabular}{l|r|r|r|r|r|r|r|r|r|r|r|r|r|r}
\hline
\end{tabular}

Table 4.5: Relative and absolute speedups of parallel algorithms. Underlined numbers mark the best relative and absolute speed-ups for the particular graph with 144 threads.

\subsection*{4.2.2 Analysis for Sequential Algorithms}

For sequential algorithms, seq_pbbs_filtering_kruskal is almost always the fastest. Notable exceptions include:
- stars_20M (Figure 9). Borůvka's algorithm only needs a single Borůvka step here because the leaves of the star will choose their only outgoing edge as the minimum edge, connecting the graph in one step. Thus Borůvka's algorithm runs in linear time here. Kruskal's algorithm needs a full sorting of random weights, and Prim's algorithm has to perform \(n\) decrease-keys or inserts, therefore both algorithms have a running time of \(\Theta(n \log n)\). Note that \(m=\mathcal{O}(n)\) for stars. The highly optimized sorting implementation from the standard library of GCC has a much lower constant factor, making Kruskal's algorithm considerably faster than Prim's algorithm and even comparable to the linear-time Borůvka's algorithm here. The case would be slightly different if Prim's algorithm first builds a priority queue in linear time and then calls \(n\) constant-time decrease-keys. However, a large constant factor would be inevitable.
- chain_20M (Figure 10), where seq_boruvka and seq_prim_binary are equally fast. On chains with increasing weights, Borůvka's algorithm finds the MST in a single Borůvka step and requires only linear time. Prim's algorithm is also fast because the priority queue only has a single element in it during the whole execution. Seq_prim_pairing is slower because of the hidden constants in its time complexity. Kruskal's algorithm needs a sorting and is thus slower. However, because the edge weights are already in increasing order, the sorting is much cheaper than sorting random numbers, which is why its running time is less than half of that for stars_20M.
- Dense uniform graphs like uniform_200K_20M (Figure 19) and uniform_20K_20M (Figure 20). That is because the number of vertices is too small compared to the number of edges, which makes sorting of the edges too costly. Prim's algorithm excels here because it does not even have to process all the edges before finding the MST. As mentioned before in Section 2.1.2, Prim's algorithm with binary heaps has an expected running time of \(\mathcal{O}(m+n \log n \log \log n)\) for graphs with random weights, which is \(\mathcal{O}(\mathfrak{m})\) for denser graphs like these two.

On real-world graphs like nlpkkt240 (Figure 21) and livejournal (Figure 23), the effect of filtering is well demonstrated because vertices have a larger average degree and hence the graphs are relatively dense. Sorting the whole set of edges is too costly, but sorting only a portion and then doing a filtering reduces the total time to only one third to one half.

It is also interesting to note that Prim's algorithm with pairing heaps is generally slower than with binary heaps except on several graphs. That is because the effect of a faster decrease-key is only visible when it is needed many times due to larger constant hidden in the Big-Oh notation.

Due to the lack of removal of used edges, Borůvka's algorithm generally does not work well with the sole exceptions of stars_20M (Figure 9) and chain_20M (Figure 10) where the Borůvka step is executed a single time anyway. However, with filtering and removal of self-edges, the new parallel algorithm works quite well with one thread. That is because the algorithm only does linear work in each Borůvka step when it is executed with a single thread. As shown in Table 4.4, the new parallel algorithm with or without filtering even beats the fastest sequential algorithm on all three real-world graphs and two uniform graphs. On other graphs, par_new_boruvka and par_new_filtering_boruvka also have comparable performance to the
fastest sequential implementation. This suggests the new algorithm can be applied even when the number of available processors is small. Other algorithms do not have such a property. The only graphs where other parallel algorithms perform better with one thread are delaunay3d_10M (Figure 14) and delaunay3d_10M-2n (Figure 15), where par_filter_kruskal excels because its multi-stage filtering removes most of the edges.

\subsection*{4.2.3 Analysis for Parallel Algorithms}

We now turn to the analysis of the running times of parallel algorithms.
par_boruvka_d [42] does not perform well on our test graphs. On randLocal_20M (Figure 5), rMat_20M (Figure 6), nlpkkt240 (Figure 21) and livejournal (Figure 23), the parallel algorithm with all available cores even runs longer than most of the sequential algorithms. Reading through the detailed logs reveals that more than \(\frac{2}{3}\) of the time is spent on the compaction step where atomic instructions are used in an unavoidably inefficient manner as described in Section 2.3.2, in contrast to our new parallel algorithm where most of the atomic instructions are only reads rather than writes. Even on graphs where par_boruvka_d exhibits speedup against sequential algorithms, the speedup comes quite late, namely when at least 9 cores are used where 3 are enough for other algorithms most of the times. Even when a speedup is visible, the algorithm is not as fast as other algorithms. For example, on graph USA (Figure 22), the algorithm needs 1.344 seconds with 72 cores, but all other algorithms are at least twice as fast. Its bottlenecks seem to be the compaction and, surprisingly, a step that marks MST edges. The latter involves a parallel for-loop of length \(n\) with one random read and one random write memory access in each Borůvka iteration.

The effect of filtering manifested itself also in the parallel settings: Borůvka's algorithm and Kruskal's algorithm with filtering are generally faster than without. The only notable exceptions are delaunay3d_10M (Figure 14) for Kruskal's algorithm and USA (Figure 22) for Borůvka's. The former is because the filtering only removes about half of the edges and thus does not justify the extra cost for partitioning and filtering. The latter is because USA is very sparse so that the first partition already contains most of the edges. Therefore the cost for this partitioning does not pay off.

As shown in the graphics, par_new_filtering_boruvka or par_new_boruvka are the fastest parallel implementation on more than half of the graphs, especially on stars_20M (Figure 9) and chain_20M (Figure 10), where the difference is large due to the reasons described above in the sequential part. On real-world graphs, i.e. nlpkkt240 (Figure 21), USA (Figure 22) and livejournal (Figure 23), the new algorithm leads by a remarkable margin and reaches an absolute speedup of more than 30. The sole exceptions where the new algorithm loses by a noticeable margin are delaunay3d_10M (Figure 14) and delaunay3d_10M-2n (Figure 15). Par_filter_kruskal is faster on these graphs due to the random weights and the graph topology. The progressive multi-stage filtering of par_filter_kruskal works in its full ability and removes most of the edges. Par_pbbs_filtering_kruskal only does one filtering and cannot achieve the same effect.

A remark on par_pbbs_filtering_kruskal and par_pbbs_kruskal is that they even need slightly more time on the union-find loop than the sequential implementation on chain_20M (Figure 10) because the algorithm is forced to run sequentially after the sorting step by the structure and weights of the graph. Though this behavior would unlikely cause problems on
graphs of practical size, it is of theoretical interest to note that this implementation can be forced to run sequentially by an adversary. The remarkable relative speedups shown in Table 4.5 are due to the highly efficient sample sort implementation in PBBS and the slowness of the algorithm with one thread.

The experiments have demonstrated that our new algorithm, together with filtering, is indeed efficient on a wide range of graphs. Unlike other parallel algorithms, the algorithm is also very efficient when only a small number of processors are available. The same even holds when only a single processor is present.

\section*{5 Conclusions and Outlooks}

In the present thesis we have briefly described known sequential and parallel algorithms for computing minimum spanning trees (MST) and forests (MSF) on shared-memory architectures. We also have presented a new conceptually quite simple yet remarkably efficient parallel algorithm for MST/MSF computation based on Borůvka's algorithm. The algorithm utilizes priority writes (pwrite) as a primitive to achieve its simplicity and reduced contention. pwrites can be easily and efficiently implemented with atomic compare-and-swap (CAS) which is widely supported by modern processors. Coarse-grained and balanced parallelism is realized nicely this way. Several optimizations that aim at improving locality of memory accesses are applied to achieve further speedup.

Experimental results on a rich set of synthetic and real-world graphs have demonstrated the extraordinary efficiency of the new algorithm. The new algorithm is faster than or as fast as its rivals on almost every graph in the benchmark. A reasonable speedup with respect to classical sequential algorithms is achieved even with only a few processors. The parallel algorithm outperforms many sequential implementations even with a single processor.

Though the algorithm proved to be efficient on many graphs, there is still room for improvement and future work. The following list gives some possible directions for future research:

Outlook 1. Theoretical time bound of the find-min step. The algorithm exhibits simplicity and excellent performance in practice. However, not much is known about its theoretical efficiency. Though we conjecture stars are the worst case for the find-min step, it remains to be carefully analyzed.

Outlook 2. Applying priority writes to other algorithms or problems. Priority writes (pwrites) are the key to the efficiency of the new MST algorithm. They seems to have the potential to be utilized to implement some known algorithms for the CRCW-Priority PRAM model on real computers. Whether this is possible or beneficial stays open for future research. It is also conceivable that they can be used in other algorithms that involve finding some minimum across multiple processors. It would be interesting to see more such examples.

Outlook 3. More thorough experiments. We have conducted benchmarks with a number of classical algorithms and state-of-the-art parallel ones. The new algorithm works very well on a wide range of test graphs. Yet the experiments did not, and could not, cover all known algorithms. An interesting future work might be a more thorough benchmark that includes more algorithms and more graphs.

Outlook 4. Generalization to other architectures. The algorithm is targeted at sharedmemory architectures and achieves good performance there. It would be natural to ask whether it can be generalized to other architectures like distributed-memory ones. At first glance it is not obvious how this can be done because synchronization is harder and more expensive on these architectures and atomic instructions are virtually nonexistent.

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[^0]:    ${ }^{1}$ The problem is defined as follows: given n points with a distance function $\mathrm{d}(\cdot, \cdot)$ satisfying the triangle inequality, i.e. $d(a, c) \leqslant d(a, b)+d(b, c)$ for all $a, b$ and $c$, find a tour with the minimum total traveled distance that visits every point exactly once and then leads back to the starting point.

[^1]:    ${ }^{2}$ The connectivity problem is the problem of identifying all connected components of a given graph.
    ${ }^{3}$ This holds as long as the input is also taken into account.

[^2]:    ${ }^{4}$ Though the graph is undirected, we sometimes assign conceptual directions to the edges. No actual modification of the edges is performed.
    ${ }^{5} \log n$ is always base- 2 logarithm and $\ln n$ is e-based throughout this thesis. Logarithms of other bases are given explicitly.

[^3]:    ${ }^{6}$ In this representation, the edge tuples are stored in a simple array of length $m$, denoted as $E[0 . . m-1]$.

[^4]:    ${ }^{7}$ The reduction to the latter is proved by discussing the two cases $m \geqslant n \log n \log \log n$ and $m<n \log n \log \log n$.

[^5]:    ${ }^{8}$ Formally, $\alpha(\mathfrak{m}, \mathfrak{n}):=\min \left\{i \geqslant 1, \mathfrak{i} \in \mathbb{Z} \left\lvert\, A\left(i,\left\lfloor\frac{m}{n}\right\rfloor\right) \geqslant \log _{2} n\right.\right\}$ where the Ackermann function is defined to be $A(1, j)=2^{j}$ for $\mathfrak{j} \geqslant 1, A(i, 1)=A(i-1,2)$ for $i \geqslant 2$, and $A(i, j)=A(i-1, A(i, j-1))$ for $i, j \geqslant 2$. The latter has an explosively fast growth, which is why the growth of its inverse is extremely slow.

[^6]:    ${ }^{9}$ Informally, pointer machines are a model of computation where no direct memory addressing is allowed. Memory cells must be found by following pointers and not by arithmetic.

[^7]:    ${ }^{10} \mathrm{An}$ adjacency array representation has an edge list sorted by their starting vertex (thus every undirected edge is stored twice in the list), and an array of length $n$ representing the index of the first outgoing edge for every source index. This way, we can efficiently find all neighbors of any given vertex.

