Fast and Space-Efficient Parallel Algorithms for Influence Maximization

Letong Wang UC Riverside lwang323@ucr.edu Xiangyun Ding UC Riverside xding047@ucr.edu Yan Gu UC Riverside ygu@cs.ucr.edu Yihan Sun UC Riverside yihans@cs.ucr.edu

Abstract

Influence Maximization (IM) is a crucial problem in data science. The goal is to find a fixed-size set of highly-influential *seed* vertices on a network to maximize the influence spread along the edges. While IM is NP-hard on commonly-used diffusion models, a greedy algorithm can achieve (1-1/e)-approximation, which repeatedly selects the vertex with the highest *marginal gain* in influence as the seed. Due to theoretical guarantee, rich literature focuses on improving the performance of the greedy algorithm. To estimate the marginal gain, existing work either runs Monte Carlo (MC) simulations of influence spread, or pre-stores hundreds of *sketches* (usually per-vertex information). However, these approaches can be inefficient either in time (MC simulation) or space (storing sketches), preventing the ideas from scaling to today's large-scale graphs.

This paper significantly improves the scalability of IM using two key techniques. The first is a *sketch-compression* technique for the independent cascading model on undirected graphs. It allows for combining the simulation and sketching approaches to achieve time-space tradeoff. The second technique includes new data structures for parallel seed selection. Using our new approaches, we implemented *PaC-IM*: Parallel and Compressed IM.

We compare PaC-IM with state-of-the-art parallel IM systems on a 96-core machine with 1.5TB memory. PaC-IM can process large-scale graphs with up to 900M vertices and 74B edges in about 2 hours. On average across all tested graphs, our uncompressed version is 5–50× faster and about 1.5× more space-efficient than existing parallel IM systems. Using compression further saves 3.8× space with only 70% overhead in time on average.

1 Introduction

Influence Maximization (IM) is a crucial problem in data science. The goal is to find a fixed-size set of highly-influential seed vertices on a network to maximize the spread of influence along the edges. For example, in viral marketing, the company may choose to send free samples to a small set of users, in the hope of triggering a large cascade of further adoptions through the "word-of-mouth" effects. Given a network G = (V, E) and a stochastic $diffusion \ model$ to specify how influence spreads along edges, we use n = |V|, m = |E|, and $\sigma(S)$ to denote the expected influence spread on G using seed set $S \subseteq V$. The IM problem aims to find a seed set S with fixed size k to maximize $\sigma(S)$. Given its importance, IM is widely-studied, and we refer the audience to a list of surveys [4, 6, 83] that reviews the numerous applications and a few hundred papers on this topic.

Among various diffusion models, Independent Cascade (IC) [31] (defined in Sec. 2) is one of the earliest and most widely-used models. In this model, only seeds are *activated* initially. In each timestamp, each vertex v that is newly activated in the last timestamp will activate its neighbors u with a probability p_{vu} .

Although IM is NP-hard on the IC model [40], the monotone and submodular properties of IC allow for a greedy algorithm with

(1 - 1/e)-approximation [40]. Given the current seed set S, the greedy algorithm selects the next seed as the vertex with the highest marginal gain, i.e., $\arg\max_{v\in V} \{\sigma(S \cup \{v\}) - \sigma(S)\}$. Due to the theoretical guarantee, this greedy strategy generally gives better solution quality than other heuristics [49]. However, the challenge lies in estimating the influence $\sigma(S)$ of a seed set S. Early work uses Monte-Carlo (MC) experiments by averaging R' rounds of influence diffusion simulation [40, 47]. However, the solution quality relies on using a large value of R', which can be expensive. Later work uses sketch-based approaches [16, 20-22, 57, 71, 72] to avoid MC experiments. Such algorithms pre-store R sketches, where each of them represents a sampled graph—each edge is chosen with the probability in which influence can spread along this edge. When estimating the influence, the diffusion is only simulated on the sampled graphs, which act as the results of the MC experiments. In an existing study [21], using $R \approx 200$ sketches can achieve similar solution quality to $R' = 10^4$ MC experiments, which greatly improves efficiency. The sketches can either be the sampled graphs, and/or memoizing more information from the sampled graphs to accelerate influence computation, such as connectivity [20, 30] or strong connectivity [57].

While numerous sketch-based solutions have been developed, we observed great challenges in scaling them to today's large-scale graphs. In a SIGMOD'17 benchmark paper [2] on nine state-of-the-art (sequential) IM solutions, none of them can process the Friendster graph (65M vertices and 3.6B edges) [45] due to either timeout (more than 40 hours) or out-of-memory. Even the recent parallel algorithms [30, 53, 54, 61] need more than half an hour to process Friendster on a 96-core machine (See Table 4). There are two major challenges in scaling sketch-based approaches to today's billion-scale graphs. The first is the *space issue*. Storing each sketch usually needs some per-vertex information. This indicates O(Rn) space, which is expensive on large graphs (empirically R is a few hundred). The second is insufficient parallelism. In particular, many state-of-the-art IM solutions uses the CELF [47] optimization for seed selection (see details below), which is inherently sequential.

In this paper, we take a significant step to *improve the scalability of sketch-based IM solutions*, and test our algorithms on *real-world billion-scale graphs*. We propose two new techniques to improve both space and time for the IM problem. Our first solution is a *sketch compression* technique for *undirected graphs and the IC model*, which limits the auxiliary space by a user-defined capacity and reduces space usage. Our second technique is *parallel data structures* for greedy seed selection that improves parallelism and reduces running time. Our new data structures work on general graphs and any diffusion model that enables submodularity. Combining the new techniques, we implemented our solutions in *PaC-IM*: <u>Parallel and Compressed IM</u>. We present a heatmap to overview our results in Fig. 1. On the aforementioned Friendster graph ("FT" in Fig. 1), *PaC-IM* only needs 128 seconds to finish

without compression (using 2.5× auxiliary space on top of the input graph), or 609 seconds when limiting auxiliary space in 0.45× input size, using a 96-core machine. This is at least 15× faster than existing parallel solutions, while being much more space efficient, and achieving the same solution quality (see Tab. 4). Below, we briefly overview the key contributions in this paper.

Our first contribution is a **novel compression scheme to store sketches** on undirected graphs and the IC model, which allows for any user-defined compression ratio (details in Sec. 3). Similar with existing algorithms [20, 30], our algorithm also memoizes connectivity information of the sketches, but *carefully avoids the O(Rn) space to store per-vertex information*. Our technique is a combination (and thus a tradeoff) of (partial) memoization with (partial) simulations. The high-level idea is to memoize the connectivity information only for $\rho = \alpha n$ *centers* $C \subseteq V$, where $0 \le \alpha \le 1$ is a user-defined parameter. The influence information of the non-center vertices will be tracked by a local simulation to find a connected center. Theoretically, we show that we can limit the auxiliary space by a factor of α by increasing the time by a factor of $O(1/\alpha)$. Experimentally, such a tradeoff is studied in Fig. 7.

Our second contribution includes two new parallel data structures for seed selection. Recall that many state-of-the-art IM solutions [20, 21, 30, 41, 54] rely on the CELF optimization [47] (details in Sec. 2) for seed selection, utilizing the submodularity of the problem. In a nutshell, CELF is an iterative approach that lazily evaluates the marginal gain of vertices in seed selection, one at a time. While laziness reduces the number of vertices to evaluate, CELF is inherently sequential, which limits the parallelism of existing IM systems. We proposed two novel solutions that achieve high parallelism without introducing much additional work. The key algorithmic challenge is to identify more vertices to evaluate in parallel, while remaining lazy such that most of the "unpromising" vertices remain untouched as in CELF. Our first solution is based on a binary search tree (BST) called P-tree [10, 12, 70] (Sec. 4.1). We highlight its theoretical efficiency (Thm. 4.1 to 4.3 and their analysis). Our second solution is referred to as Win-Tree (Sec. 4.2), which is simpler and has lower space usage, leading to slightly better overall performance. The new data structures are general and work on both directed and undirected graphs and any diffusion model with submodularity. More importantly, we believe they apply to general optimization problems with submodular objective functions. We leave this as future work, and discuss it briefly in Sec. 7.

Our final contribution is an in-depth experimental study of the proposed techniques, and the performance comparison among PaC-IM and state-of-the-art IM systems. Our experiment is on four types of a total of 17 graphs, five of which have more than 1B edges. Besides social networks, we also tested other real-world graphs including web graphs, road graphs and k-NN graphs. One can consider IM on such graphs as studying the influence diffusion between webpages, geologically or geometrically close objects. In all settings, PaC-IM achieves the best running time and space usage on all tested graphs, while guaranteeing better or comparable solution quality to all baselines. Compared to the best baseline, PaC-IM with no compression is $5.6\times$ faster and is $1.5\times$ more space-efficient on average (geometric mean across tested graphs), and is $3.2\times$ faster and $6\times$ more space-efficient using compression with $\alpha=0.1$. Due to space- and time-efficiency, PaC-IM is the only system that can

		Ou			Our			Infuse	_		Ripp	
_		Time	Space		Time	Space		Time	Space		Time	Space
اء	EP	1.00	1.00		1.67	0.39		1.28	1.18	П	14.1	1.53
Web)	SLDT	1.00	1.00		1.76	0.38		1.60	1.17		86.3	1.99
뎔	DBLP	1.00	1.00		1.06	0.21		2.5	1.40		121	0.60
a	YT	1.00	1.00		2.00	0.17		5.09	1.44		58.4	0.94
Si.	OK	1.00	1.00		4.50	0.40		9.13	1.3		37.4	12.8
(Social and	IJ	1.00	1.00		3.42	0.27		10.2	1.76		20.5	3.41
e (TW*	1.00	1.00		4.05	0.41		6.84	1.63		67.0	11.6
Į.	FT*	1.00	1.00		4.75	0.41		15.4	1.64		-	-
Scale-free	SD*	1.00	1.00		4.18	0.36		11.2	1.69		-	-
Š	MEAN	1.00	1.00		2.70	0.32		5.21	1.66		46.1	2.63
Z	GER	1.00	1.00		0.91	0.19		2.86	1.89		250	1.73
and kNN)	USA	1.00	1.00		0.94	0.20		3.63	1.90		933	1.80
밀	HT5	1.00	1.00		0.95	0.18		5.90	1.41		66.2	0.61
l g	HH5	1.00	1.00		0.93	0.17		3.62	1.42		23.7	0.41
(Road	CH5	1.00	1.00		1.51	0.22		62.2	1.80	П	11.1	0.36
	GL5	1.00	1.00		0.92	0.22		5.96	1.87		32.7	0.32
Sparse	COS5*	1.00	1.00		0.82	0.19		6.66	1.88		-	-
Sp	MEAN	1.00	1.00		0.98	0.19		6.60	1.73		71.4	0.67
ľ	ΛΕΑΝ	1.00	1.00		1.76	0.26		5.66	1.56		55.6	1.46
0	.2 .5	1 2	4 8	-	16 >16	MEA	۱,	N = ge	ometr	ic	mear	า

Figure 1: Heatmap of relative running time and space usage, normalized to Ours₁. Ours₁: *PaC-IM* with no compression. Ours_{0,1}: *PaC-IM* with 10× sketch compression. *InfuserMG* [30] and *Ripples* [54]: existing parallel IM systems. Lower/green is better. The graph information is in Tab. 3. The absolute running times are in Tab. 4. *: graphs with more than a billion edges. (Colors are adjusted to be distinguishable in gray-scale.)

process the largest graph ClueWeb [52] with 978M vertices and 75B edges. We believe *PaC-IM* is the first IM solution that scales to tens of billions of edges and close to a billion vertices.

2 Preliminaries

Basic Notations. For graph G = (V, E), we use n = |V| and m = |E|. Since our sketch compression is designed for undirected graphs and IC model, for simple description, throughout the paper, we assume G is undirected and consider the IC model unless otherwise specified. On undirected graphs, a **connected component (CC)** is a maximal subset in V s.t. every two vertices in it are connected by a path. In a max-priority-queue, we use **top** to refer to the element with the largest key, and use function **pop** to find and remove the top element. We use $\tilde{O}(f(n))$ to denote $O(f(n) \cdot \text{polylog}(n))$.

Computational Model. We use the standard binary fork-join model [13, 23] to support parallelism, and the work-span analysis [15, 36]. We assume a set of threads that access a shared memory. A process can fork two child software threads to work in parallel. When both children complete, the parent process continues. A parallel for-loop can be simulated by recursive forks in logarithmic levels. The **work** of an algorithm is the total number of instructions and the **span** (depth) is the length of the longest sequence of dependent instructions. We can execute the computation using a randomized work-stealing scheduler [3, 15] in practice. We assume unit-cost *atomic* operation WRITEMAX(t, v_{new}), which atomically reads the memory location pointed to by t, and write value v_{new} to it if v_{new} is larger than the current value in t. We use the compareand-swap function to implement WRITEMAX in practice.

```
G = (V, E): The input graph.
                                                             k: number of seeds.
                                                                                              Algorithm 1: Sketch-based IM algorithm
\sigma(S) or \sigma_{G,M}(S): The influence spread of seed set S \subseteq V on graph G
   and diffusion model M.
\Delta(v \mid S): The true score (marginal gain) of v on top of S. \Delta(v \mid S) =
   \sigma(S \cup \{v\}) - \sigma(S). We omit S and use \Delta(v) with clear context.
\bar{\Delta}[v]: The stale score (lazily-evaluated marginal gain) of v stored in an
   array. It is an upper bound of \Delta(v \mid S) for the current seed set S.
\Phi_{1..R}~: the sketches. Formally defined in Sec. 3.
                                                                                                  \Phi_r \leftarrow \text{Sketch}(r)
\rho and \alpha : \rho = \alpha n is the number of centers.
Function names:
```

Sketch(G, r) : Compute the r-th sketch from graph G

MARGINAL $(S, v, \Phi_{1..R})$: The marginal gain of vertex v given the current seed set S estimated from R sketches $\Phi_{1...R}$

NextSeed $(S, \Phi_{1...R})$: Greedily determine the next seed based on sketches $\Phi_{1..R}$ given the current seed set S.

Table 1: Notations in the paper.

The Influence Maximization (IM) Problem

Given a graph G = (V, E), an influence diffusion model M specifies how influence spreads from a set of current active vertices to acti*vate* more vertices in V. Given a *seed* set $S \subseteq V$, we use $\sigma_{G,M}(S)$ to denote the expected number of vertices that *S* can activate (including S) under diffusion model M on graph G. The Influence **Maximization (IM)** problem is to find a subset $S^* \subseteq V$ with size k, s.t. S^* maximizes the influence spread function $\sigma_{G,M}$. With clear context, we omit M and G, and use $\sigma(\cdot)$. Several propagation models have been proposed in the literature, including the Independent Cascade (IC) model [31], Linear Threshold (LT) model [33, 65], and more [18, 40, 50, 63]. Since our sketch compression focuses on the IC model, we briefly introduce the IC model here. In the IC model, influence spreads in rounds. Initially, only the seed vertices are active. In round i, each vertex u that was newly activated in round i-1 attempts spread the influence via all incident edges e, and activate the other endpoint v with probability p_e .

Submodularity and the Greedy Algorithm. Kempe et al. [40] proved that IM under the IC model is NP-hard, and that the influence spread function is **submodular**, i.e., for every $X, Y \subseteq V$ where $X \subseteq Y$, and $v \in V \setminus Y$, we have:

$$\sigma(X \cup \{v\}) - \sigma(X) \ge \sigma(Y \cup \{v\}) - \sigma(Y)$$

Due to the submodularity, the following greedy algorithm (later referred to as *GeneralGreedy*) can achieve a (1-1/e)-approximation. The algorithm starts with $S=\emptyset$ and repeatedly adds the vertex with the highest *marginal gain* to S, until |S| = k. The marginal gain $\Delta(v \mid S)$ of a vertex v given the current seed set S is defined as:

$$\Delta(v \mid S) = \sigma(S \cup \{v\}) - \sigma(S) \tag{1}$$

With clear context, we omit *S* and use $\Delta(v)$, and also call it the *score* or the *true score* of v. We call the process to compute the true score of a vertex an *evaluation*. To estimate $\sigma(\cdot)$, early solutions average R' rounds of Monte Carlo (MC) experiments of influence diffusion simulation. However, on real-world graphs, this approach requires a large value of R' to converge, which can be expensive.

Sketch-Based Algorithms. Sketch-based algorithms are proposed to accelerate influence spread simulation. Instead of running independent MC experiments in every evaluation, sketch-based algorithms statically sample R graphs, each reflecting an MC experiment, and always simulate and average the result on the *R* sampled graphs for any evaluation. As mentioned in Sec. 1, using sketches allows

```
Notations: G = (V, E): the input graph. K: the number of seed vertices.
              R: the number of sampled graphs
  Output: S: a set of K seeds that maximizes influence on G
  Notes: \Phi_{1...R}: R sketches computed from R sampled graphs
  // Step 1: Sketch construction
1 ParallelForEach r ← 1 . . . R do
```

```
// Compute the r-th sketch
   // Step 2: Seed selection using CELF
s S \leftarrow \emptyset
4 while |S| < K do
         s^* \leftarrow \mathsf{NextSeed}(S, \Phi_{1..R}) \quad \textit{//Find} \ \mathsf{arg} \ \mathsf{max}_{v \in V} \mathsf{Marginal}(S, v, \Phi_{1..R})
                                                           // Mark s* as a seed in the sketches
         MarkSeed(s^*, \Phi_{1..R}, S)
         S \leftarrow S \cup \{s^*\}
8 return S
```

Algorithm 2: Sequential Seed Selection with CELF

```
Notes: Q: max-priority-queue on all vertices v \in V with key \bar{\Delta}[v]
        Initially \bar{\Delta}[v] = \text{Marginal}(\emptyset, v, \Phi_{1..R})
1 Function NextSeed (S, \Phi_{1..R})
                                                 // S: current seed set; \Phi_{1..R}: R sketches
2
        Repeat
3
             s^* \leftarrow Q.Pop()
                                                         // Pop: find and remove the top
              \bar{\Delta}[s^*] \leftarrow \text{Marginal}(S, s^*, \Phi_{1..R})
4
              if \bar{\Delta}[s^*] < Q.Top() then return s^*
              else Q.Insert(s*)
                                                        // insert s* back with new score
```

for much faster convergence with the number of simulations (i.e., sketches) R than that in MC experiments. Therefore, sketch-based algorithms are widely studied [16, 20-22, 57, 71, 72]. We summarize sketch-based algorithms in two steps (see Alg. 1): sketch construction and seed selection. Next, we briefly introduce both steps, and some useful techniques in previous work. We summarize some related approaches in Tab. 2, and review more existing work in Sec. 6. Some useful notations are given in Tab. 1.

Sketch Construction. In the earliest sketch-based algorithm StaticGreedy, R sampled graphs [21] are explicitly stored as sketches. In the IC model, the *r*-th sketch corresponds to a sampled graph $G'_r = (V, E'_r)$, where $E'_r \subseteq E$, such that each edge $e \in E$ is sampled with probability p_e , meaning a successful activation. An evaluation will average the number of reachable vertices on all sampled graphs from the seed set S. A later paper Infuser proposed the fusion optimization [30], which uses hash functions to avoid explicitly storing the sampled graphs G'_r . They sample an edge e in a sketch G'_r with a random number generated from seed $\langle e, r \rangle$, such that whether an edge is selected in a certain sketch is always deterministic, and a sampled graph G'_r can be fully reconstructed from the sketch id r. We also use this idea in our sketch compression algorithm.

Many existing algorithms also use memoization on sketches to avoid influence spread simulation on sketches. On undirected graphs and IC model, the MixGreedy paper [20] first observed that a vertex v's influence on a sketch is all vertices in the same connected component (CC) as v, but they only used this idea to select the first seed. The Infuser system adopts this idea to select all seeds, and proposes InfuserMG, which memoizes the CC information of each sampled graph as the sketch. A vertex v's score is then the average of the (inactivated) CC sizes on all R sketches, which can be

	Randomization	Compute Influence	Select Seed	#vertics per seed	Space	Parallel
GeneralGreedy [40]	R' Monte Carlo experiments	Simulation	Evaluate all	O(nR'T)	O(n)	no
MixGreedy [20] 1st Seed	Fixed R sampled graphs	Memoization	Evaluate all	O(nR)	O(n)	no
MixGreedy [20] Others	R' Monte Carlo experiments	Simulation	CELF	$O(n_c R'T)$	O(n)	no
StaticGreedy [21]	Fixed R sampled graphs	Simulation	CELF	$O(n_cRT)$	O(n)	no
InfuserMG [30]	Fixed R sampled graphs	Memoization	CELF	$O(n_c R)$	O(nR)	yes
PaC-IM (this work)	Fixed R sampled graphs	Simulation + Memoization	n CELF	$O(n_c R \cdot \min(T, 1/\alpha))$	$O((1 + \alpha R)n)$	yes

Table 2: Existing and our new approaches. MixGreedy uses different approaches to select the first seed and the other seeds, so we list them separately. "#vertices per seed": number of vertices to visit in all re-evaluations involved to find a seed. n: number of vertices. n_c : number of re-evaluations needed in CELF. T: the average number of reachable vertices in a simulation. Empirically, T is large, and $n_c \ll n$. To achieve similar quality, $T \ll n$.

obtained in O(R) cost. This approach avoids simulation, but leads to O(Rn) space that is expensive for large graphs. Sec. 3 shows how our sketch compression approach reduces this high space usage.

Seed Selection with CELF Optimization. Another useful optimization for the greedy algorithm is CELF [47], which avoids evaluating all vertices to select the next seed. CELF uses lazy evaluation for submodular functions, where a vertex u is evaluated only if it becomes "promising" as a candidate of the next seed. We show the CELF-based seed selection algorithm in Alg. 2. The algorithm uses a priority queue *Q* to maintain all vertices with their scores as the key. Due to submodularity, $\Delta(v)$ is non-increasing with the expansion of the seed set S. With lazy evaluation, the scores in Q may be stale, but are always upper bounds of the true score. To distinguish it from the true score, we call this lazily-evaluated score the **stale score** of v, and denote it as $\bar{\Delta}[v]$ stored in an array. To select the next seed, CELF keeps popping the top element v from Q, re-evaluating its true score $\Delta(v)$, and inserting it back unless $\Delta(v)$ is greater than the current top $\bar{\Delta}[\cdot]$ value in Q (Line 3-Line 6). In this case, we can set v as the next seed without further examining the remaining vertices in *Q*, since their true score can only be lower than the values in Q. In many cases, CELF can reduce the number of re-evaluations, but it is essentially sequential and evaluates all vertices one by one. In Sec. 4, we present our new data structures that allow for parallel re-evaluations in CELF.

3 Space-Efficient Sketches

This section presents our new technique to construct compressed sketches in parallel for the IC model on undirected graphs. In this setting, as discussed in Sec. 2, a vertex v can activate all vertices in the same CC on a certain sketch. Thus, memoizing per-vertex CC information in sketches [30] can accelerate the influence evaluation, but requires O(Rn) space, which does not scale to large input graphs. Alternatively, one can avoid memoization and run a simulation by traversing the sampled graph to find the CC when needed. This requires no auxiliary space, but can take significant time.

Our approach combines the benefit of both, using bounded-size auxiliary space while allowing for efficiency. Our key idea is to only partially memoize CC information in each sketch, and retrieve this information by partial simulations. In PaC-IM, we only store the CC information for $\rho = \alpha n$ center vertices $C \subseteq V$, where $0 \le \alpha \le 1$ is a user-defined parameter. Each sketch Φ_r is a triple $\langle r, label[\cdot], size[\cdot] \rangle$ corresponding to an implicit sampled graph G'_r from G, where each edge e = (u, v) is retained with probability p_e .

- *r*: the sketch id. Similar to the fusion idea mentioned in Sec. 2, the sketch id fully represents the sampled graph.
- $label[1..\rho]$: the CC label of center c_i on this sketch. It is the smallest center id j where c_j is in the same CC as c_i .

```
Algorithm 3: Our sketch algorithm with compression
   Global Variables: G = (V, E): the input graph
        C = \{c_1, c_2, \dots, c_{\rho}\} \subseteq V: randomly selected centers. \rho = |C|
        R: the number of sketches
   Notes: A sketch \Phi_r is a triple \langle r, size[1...\rho], label[1...\rho] \rangle:
        r: the id of the sketch.
         size[1...\rho]: size[i] is the connect component size of center i.
         label[1..\rho]: label[i] is the connect component id of center i.
1 Function Sketch(r)
                                                                          // r: sketch id
        Compute the connected components of graph G' = (V, E'), where
          E = \{(u, v) \mid (u, v) \in E, SAMPLE(u, v, r) = true\}
        ParallelForEach c_i \in C do
           label[i] \leftarrow \min_{i} \{c_i \text{ is in the same CC as } c_i\}
4
        ParallelForEach c_i \in C do
5
             if label[i] = i then size[i] \leftarrow the CC size of center c_i
        return \langle r, label[1...\rho], size[1...\rho] \rangle
   // sample an edge e with probability p_e for sketch r
8 Function Sample(e, r)
                                              // e: edge identifier, r: sketch identifier
        p \leftarrow random(e, r)
                                       // Generate p \in [0, 1] from random seed e, r
10
        return p \leq p_e
   // \delta: marginal influence of v on sketch \Phi_r. l: label of v's CC in sketch \Phi_r if v
      is connected to any center; otherwise l = -1. S: the current seed set.
11 Function \langle \delta, l \rangle = GetCenter(Φ<sub>r</sub>, v, S)
        Start BFS from v on a sampled subgraph of G, where an edge e \in E
          exists if Sample(e, r) = true. Count the #reached-vertices as n'
        if a center c_i \in C is encountered during BFS then
13
14
             l \leftarrow \Phi_r.label[i]
                                                         // Find the label of the center
15
             return \langle \Phi_r.size[l], l \rangle
                                                // the CC size and label of the center
        if any v' \in S has been visited by BFS then return (0, -1)
16
        else return \langle n', -1 \rangle
                                               // influence is #visited vertices in BFS
17
   // Marginal gain of v given seed set S on sketches \Phi_{1..R}
18 Function Marginal(S, v, \Phi_{1..R})
        ParallelForEach r \leftarrow 1..R do
19
         \langle \delta_r, \cdot \rangle \leftarrow \text{GetCenter}(\Phi_r, v, S)
20
        return (\sum_{r=1}^{R} \delta_r)/R
                                             // the sum can be computed in parallel
21
22 Function MarkSeed (v, \Phi_{1..r}, S)
        ParallelForEach r \leftarrow 1..R do
                                                                  // For each sketch \Phi_r
23
             \langle \delta, l \rangle \leftarrow \text{GetCenter}(\Phi_r, v, S)
24
             if l > 0 then \Phi_r.size[l] \leftarrow 0 // Clear the corresponding CC size
```

• $size[1..\rho]$: if label[i] = i (i.e., i represents the label of its CC), size[i] is the influence of center c_i on this sketch. It is initially the CC size of c_i , and becomes 0 when any vertex in this CC is selected as a seed.

Algorithm Overview. We first present the high-level idea of our sketch compression algorithm. An illustration is presented in Fig. 2. As mentioned, we only store the CC information (label and size)

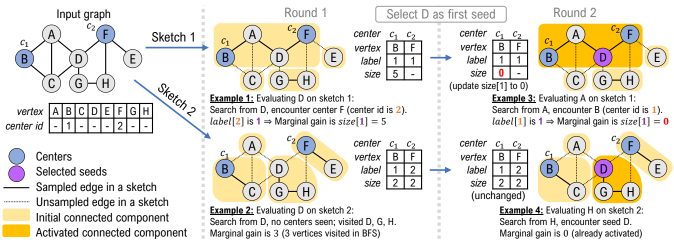


Figure 2: An example of our sketch compression on a graph with 8 vertices and R = 2 sampled graphs.

for $\rho = \alpha n$ centers for each sketch. We further use a global flag array to indicate if a vertex is a center, and therefore the total space usage is $O((1+\alpha R)n)$. To query the CC size of an arbitrary vertex v on sketch Φ_r , we will start a breadth-first search (BFS) from v. When any center c_i is encountered, we know that v should activate the same set of vertices as c_i on this sketch. As such, we can stop searching and use the influence (CC size) of c_i as the influence for v. If v is not connected with any center on the sampled graph, the CC containing v is likely small, and the BFS can visit all of them quickly. In either case, the number of vertices visited by the BFS can be bounded. We show in Thm. 3.1 that limiting the auxiliary space by a factor of α roughly increases the evaluation time by a factor of $O(1/\alpha)$. By controlling the number of centers $\rho = \alpha n$, we can achieve a tradeoff between the time to query the score of each vertex and the space usage.

We present our algorithm in Alg. 3. Next, we will elaborate on the three functions needed in Alg. 1: Sketch(G, r), which constructs the r-th sketch from input graph G, Marginal($S, v, \Phi_{1..R}$), which computes the score (marginal gain) of a vertex v on top of S using sketches $\Phi_{1..R}$, and MarkSeed($s^*, \Phi_{1..R}$), which adjusts the sketches $\Phi_{1..R}$ when s^* is selected as a seed.

Sketch Construction (Sketch(G,r)). We start by selecting $\rho = \alpha n$ center vertices $C = \{c_1, c_2, \dots, c_\rho\}$ uniformly at random. We will only maintain the CC information of the ρ centers on all sketches.

To construct a sketch Φ_r , we first compute the CC information of the sampled graph G'_r , which can be performed by any parallel connectivity algorithm [25]. We store the CC information for all centers in two arrays. $\Phi_r.label[i]$ records the label of CC of the i-th center. For multiple centers that are in the same CC, we simply use the smallest CC id as the label for all of them to represent this CC, so all such centers refer to the label to find the CC information. For a center c_i , If i is the label of its CC, we use $\Phi_r.size[i]$ to record the size of this CC. An example of these arrays is given in Fig. 2.

Computing the Marginal Gain (Marginal $(S, v, \Phi_{1..R})$). Given the sketches $\Phi_{1..R}$ and the current seed set S, the function Marginal $(S, v, \Phi_{1..R})$ computes the marginal gain of a vertex v. Our algorithm loops over all sketches and sums up the marginal gains

of v on all of them. We use a helper function $\langle \delta, l \rangle = \text{GetCen}$ TER(Φ_r , v, S), which returns δ as the marginal gain of v on sketch Φ_r , and l as the label of centers connected to v (l=-1 if no center is connected to v). This function will run a breadth-first search (BFS) from v on G'_r (i.e., only using edges $e \in E$ s.t. SAMPLE(e, r)is true). If any center is encountered during this BFS (Examples 1 and 3 in Fig. 2), then the influence of v is the same as c_i on this sketch. The information of c_i is retrieved by its label $l = \Phi_r.label[i]$, and thus $\delta = \Phi_r.size[l]$. The influence δ is either the size of the CC containing v when no vertices in this CC are seeds (Example 1 in Fig. 2), or 0 otherwise, as is updated in MARKSEED (Example 3 in Fig. 2). Otherwise, if the BFS terminates without visiting any centers, the returned label l will be -1. If any seed is visited during BFS, then the marginal gain is $\delta = 0$ (Line 16, Example 4 in Fig. 2). Otherwise δ is the number of vertices n' visited during BFS, which is also the size of CC containing v (Example 2 in Fig. 2). Using the GetCenter function, the marginal influence of v on sketch Φ_r can be obtained as the first return value δ of GetCenter(Φ_r, v, S).

Marking a Seed (MarkSeed($s^*, \Phi_{1..R}$)). This function updates the sketches when $s^* \in V$ is selected as a seed. For each sketch Φ_r , the CC label of s^* is the second return value l of GetCenter(Φ_r, s^*, S). If $l \neq -1$, we set $\Phi[r]$. size[l] as 0—since s^* is selected, all other vertices in this CC will get no marginal gain on this sketch.

Our sketch compression scheme allows for a tradeoff between the space usage and the query time in Marginal(): a smaller value of ρ (fewer centers) leads to lower space usage, but also increases the cost to evaluate vertices, as it may take longer to find a center. It is worth noting that PaC-IM unifies and is a hybrid of StaticGreedy and InfuserMG. Theoretically, using $\alpha=1$, our sketch Φ is equivalent to InfuserMG where the CC information for all vertices on all sketches are memoized; using $\alpha=0$, our algorithm is equivalent to StaticGreedy where no CC information is maintained, and evaluating the influence is simply done by traversing the sampled graph. In practice, PaC-IM is much faster than StaticGreedy and InfuserMG even when with no compression due to better parallelism and other new techniques (e.g., those in Sec. 4). We summarize the theoretical guarantees in Tab. 2, and state them in Thm. 3.1.

Theorem 3.1. PaC-IM with parameter α requires $O((1 + \alpha R)n)$ space to maintain R sketches, and visit $O(R \cdot \min(1/\alpha, T))$ vertices to re-evaluate the marginal gain of one vertex v, where T is the average CC size of v on all sketches.

Proof. Assume that no connectivity is stored in the sketches, then running a simulation for a vertex will visit RT vertices, based on the definition of T. This is also what the baseline algorithm StaticGreedy does. We now consider the case that $\rho = \alpha n$ centers are selected. Note that all centers are picked independently and uniformly at random. This means that each visited vertex in the BFS has the probability of α that is a center. Let us first focus on a search on a specific sketch r, and assume the CC size of v is T_r . The search terminates either 1) all T_r vertices have been visited, or 2) a center is encountered. Therefore, when visiting the T' vertices in the BFS order, each of them stops the search with a probability of α . The expected number of visited vertices in the BFS is therefore also bounded by $\sum_{i=1}^{\infty} (1-\alpha)^{i-1}\alpha \cdot i$, which solves to $1/\alpha$. The total number of visited vertices in all BFS is $\sum_r \min(T_r, 1/\alpha) = \min(RT, R/\alpha) = R \cdot \min(T, 1/\alpha)$.

4 Parallel Priority Queues for Seed Selection

We now present the parallel seed selection process in PaC-IM. We call each iteration in seed selection (selecting one seed) a round. Recall that prior solutions use the CELF optimization (see Tab. 2 and Sec. 2), which maintains (possibly stale) scores of all vertices in a priority queue Q, and updates them lazily. In each round, CELF pops the vertex with the highest (stale) score from Q, and re-evaluates it. The process terminates when a newly evaluated score is higher than all scores in O: otherwise, the new score will be inserted back to O. For simplicity, we assume no tie between scores. In the cases when ties exist, both our approach and CELF break the tie by vertex id. Let $F_i = \{v \mid \bar{\Delta}_{i-1}[v] \geq \Delta^*\}$ be the set of vertices re-evaluated by CELF in round *i*, where $\bar{\Delta}_{i-1}[v]$ is the stale score of *v* after round i-1, and Δ^* is the maximum true score in round i (i.e., the score of the chosen seed in round i). Empirically, $|F_i|$ is usually large in the first several rounds, but can be rather small in later rounds, in which case CELF greatly reduces the numbers of re-evaluations.

However, the CELF process is inherently sequential. Hence, some existing parallel implementations (e.g., [30, 54]) only parallelize the re-evaluation function Marginal (Line 4 in Alg. 2), but leave the CELF process sequential, i.e., all $|F_i|$ evaluations are performed one by one. When F_i is large, this results in low parallelism.

Prior Work on Parallel Priority Queue. As a fundamental data type, parallel priority queues are widely studied [9, 12, 26, 64, 69, 70, 77]. However, as far as we know, all these algorithms/interfaces require knowing the batch of operations (e.g., the threshold to extract keys, or the number of keys to extract) *ahead of time*. This is not true in CELF—the set F_i is only known during the execution. We need different approaches to tackle this challenge.

Overview of Our New Approaches. We first formalize the interface of the parallel priority queue needed in the IM framework in Alg. 1. The data structure maintains an array $\bar{\Delta}[\cdot]$ of (stale) scores for all vertices. It is allowed to call Marginal function to re-evaluate and obtain the true score of any vertex. The interface needs to support NextSeed function, which returns the vertex id with the highest true score.

Algorithm 4: Seed Selection based on P-tree

```
Maintains: A parallel binary search tree T for all \bar{\Delta}[v].
1 Function NextSeed (S, \Phi_{1..R})
                                                                           // The best seed so far
2
         s^* \leftarrow \bot
          j \leftarrow 0
3
4
         repeat
               // Extract (remove & output) the top 2^{j} elements in T into array B_{i}
               B_i[1..2^j] \leftarrow T.SplitAndRemove(2^j)
               // Evaluate the true score for each v \in B_i
 6
               ParallelForEach v \in B_i do
                 \bar{\Delta}[v] \leftarrow \text{Marginal}(S, v, \Phi_{1..R})
               t \leftarrow \arg\max_{v \in B_i} \bar{\Delta}[v]
                                                               // can be computed in parallel
8
               if (s^* = \bot) OR (\bar{\Delta}[t] > \bar{\Delta}[s^*]) then s^* \leftarrow t
               j \leftarrow j + 1
10
          until \bar{\Delta}[s^*] > T.\text{Max}()
                                                         //\bar{\Delta}[s^*] is better than the top in T
11
          T.BATCHINSERT(\bigcup_{j'=0}^{j-1} B_{j'} \setminus \{s^*\})
12
         return s*
```

We present two parallel data structures to maintain the lazily evaluated scores in parallel. Our first approach is based on a parallel binary search tree (BST), the P-tree [10, 12, 70]. We prove that using P-trees, our approach has work (number of re-evaluations) asymptotically the same as that in CELF, while is highly parallel: the *i*-th seed selection finishes in roughly $\log |F_i|$ iterations of reevaluations (each iteration evaluates multiple vertices in parallel), instead of $|F_i|$ iterations (each iteration evaluate one vertex) in CELF. We also propose a new data structure based on parallel winning tree (aka. tournament tree), and refer to it as Win-Tree. Win-Tree does not maintain the total order of the scores, and is simpler but potentially more practical than P-trees. We introduce the P-treebased approach in Sec. 4.1 and Win-Tree-based approach in Sec. 4.2, and compare their performance in Sec. 5.3. These two approaches are independent of the sketching algorithm and apply to seed selection on any submodular diffusion models (not necessarily the IC model and/or on undirected graphs). Note that almost all diffusion models for IM are submodule (e.g., Independent Cascade (IC), Linear Threshold (LT), Triggering (TR [40]), and more [40, 73, 80]).

4.1 Parallel Priority Queue Based on P-tree

To parallelize CELF, our first approach is to maintain the total (decreasing) order of the scores of all vertices, using a parallel binary search tree (BST) called P-tree [10, 12]. We will use two functions on a P-tree T: 1) T.SplitAndremove(k), which extracts (removes and outputs to an array) the first k tree nodes (k largest scores) from T, and 2) T.BatchInsert(B), which inserts a set of keys B to T. Both algorithms are parallel with polylogarithmic span.

Different from many priority queue implementations (e.g., binary heap) that extract the top element one at a time, P-tree maintains the total order of all vertices, and thus we can extract a batch of vertices with top (stale) scores and evaluate them in parallel. The key challenge here is to design a similar stop condition as CELF to avoid evaluating too many vertices, since we are unaware of the number of "useful" re-evaluations, i.e., $|F_i|$, ahead of time. Our idea is to use $prefix\ doubling\ [14, 24, 34, 35, 66]$ to achieve both work efficiency and high parallelism. The pseudocode is given in Alg. 4 with an illustration in Fig. 3. To find the next seed, the P-tree T starts with the stale scores $\bar{\Delta}[\cdot]$ from the previous round. It will

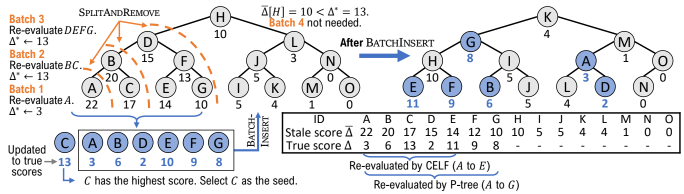


Figure 3: Example of *P-tree-***Based Seed Selection.** The letters in the tree nodes represent vertices, and the numbers below them are their stale scores. *P-tree* maintains decreasing order of the (stale) scores. By prefix-doubling, we extract batches of 1, 2, 4 vertices and evaluate each batch in parallel. After batch 3, the highest true score (13) is higher than the current best in the tree (10), and the algorithm stops. We will select the node with the highest true score and insert the rest back to the tree with their new score. *P-tree* may evaluate more vertices than CELF, but the extra work can be bounded (Thm. 4.2).

then extract the top (largest stale score) nodes in batches of size 1, 2, 4, 8, ... from T (Line 5). Within each batch B_j , we re-evaluate all vertices in parallel (Line 7). These new scores are used to update the current best seed s^* . The loop terminates when the score of s^* is better than the best score in T (Line 11). Finally, the algorithm selects s^* as the seed and inserts the rest of the new true scores $\bigcup B_j \setminus \{s^*\}$ back to T.

Due to prefix doubling, each seed selection finishes in at most $O(\log n)$ rounds. Note that our approach evaluates more vertices than CELF, but due to the stop condition (Line 11), the extra work is bounded by a constant factor (proved in Thm. 4.2). Next, we prove the correctness and efficiency of the algorithm.

Theorem 4.1 (*P-tree* Correctness). Alg. 4 always selects the next seed with the largest marginal gain, i.e., $\Delta(s^*) = \max_{v \in V} \{\Delta(v)\}$. Proof. Let s^* be the vertex selected by the algorithm, we will show $\Delta(s^*) \geq \Delta(v)$ for all $v \in V$ when the stop condition on Line 11 is triggered. We first show that $\Delta(s^*) \geq \Delta(v)$ for all $v \in V$ that has been split from the tree. This is because all such vertices have been re-evaluated, and s^* by definition has the highest true score among them. We then show s^* has a higher true score than any other vertices still in T. For any $u \in T$, the stop condition indicates $\Delta(s^*) = \bar{\Delta}[s^*] > \bar{\Delta}[u] \geq \Delta(u)$ (due to submodularity). Therefore, $\Delta(s^*)$ is the highest true score among all vertices.

THEOREM 4.2 (P-TREE EFFICIENCY). Alg. 4 has the total number of re-evaluations at most twice of the number by CELF.

Proof. Recall that F_i is the set of re-evaluated vertices by CELF when selecting the i-th seed, and let F_i' be the set of re-evaluated vertices by P-trees using Alg. 4. We first show a simple case—if both CELF and Alg. 4 start with the same stale scores $\bar{\Delta}_{i-1}[\cdot]$, then $|F_i'| \leq 2|F_i|$. Let us reorder vertices in V as v_1, v_2, \ldots, v_n by the decreasing order of their stale score $\bar{\Delta}_{i-1}[\cdot]$. Assume v_l is the last vertex evaluated by CELF, so $|F_i| = l$. v_l must be in the last batch in P-tree. Assume the last batch is batch j with 2^j vertices. This indicates that all $2^j - 1$ vertices in the previous j - 1 batches are before v_l . Therefore $2^j - 1 < l = |F_i|$, and $|F_i'| = 2^{j+1} - 1$, which proves $|F_i'| \leq 2|F_i|$.

We now consider the general case. We first focus on a specific seed selection round *i*. Due to different sets of vertices re-evaluated in each round, at the beginning of round *i*, CELF and *P-tree* may

not see exactly the same stale scores. We denote the stale score at the beginning of round i in CELF as $\bar{\Delta}_{CELF}[\cdot]$ and that for P-tree as $\bar{\Delta}_{BST}[\cdot]$. We reorder vertices by decreasing order of $\bar{\Delta}_{CELF}[\cdot]$ as v_1, v_2, \ldots, v_n , and similarly for $\bar{\Delta}_{BST}[\cdot]$ as u_1, u_2, \ldots, u_n . Denote Δ^* as the highest true score in round i. Let v_{x_i} be the last vertex in $v_{1...n}$ such that $\bar{\Delta}_{CELF}[v_{x_i}] \geq \Delta^*$, and u_{y_i} the last vertex in $u_{1...n}$ such that $\bar{\Delta}_{BST}[u_{y_i}] \geq \Delta^*$. Namely, x_i is the rank of Δ^* in $v_{1...n}$, and y_i is the rank of Δ^* in $u_{1...n}$. By definition, F_i is exactly the first x_i vertices in sequence v, and thus $x_i = |F_i|$; F_i' contains all vertices smaller than u_{y_i} and possibly some more in the same batch with u_{y_i} , so $|F_i'| \leq 2y_i$. In the simple case discussed above where we assume $\bar{\Delta}_{CELF}[\cdot] = \bar{\Delta}_{BST}[\cdot]$, we always have $x_i = y_i$.

We will use amortized analysis to show that $\sum y_i \leq \sum x_i$, which further indicates $\sum |F_i'| \le 2 \cdot \sum |F_i|$. We can partition $u_{1...y_i}$ into two categories: U_1 as the intersection of $u_{1...y_i}$ and $v_{1...x_i}$, and U_2 as the rest. Note that for $t \in U_2$, $\bar{\Delta}_{BST}[t] \geq \Delta^* > \bar{\Delta}_{CELF}[t]$. This happens iff. t is evaluated by CELF in a previous round, but not by *P-tree*. Namely, there exists a round j, such that $t \in F_j$ but $t \notin (F'_j \cup F'_{j+1} \cup ... \cup F'_{i-1})$. Let $v^*_{1..n}$ and $u^*_{1..n}$ be the u and v sequences in round j, respectively. Note that $F_j = v_{1...x_j}^*$, and $u_{1...y_j}^*\subseteq F_j'$. Since $t\in F_j$, then vertex t was counted in x_j . Since $t\notin F_j'$, t is not counted in any y_j . Therefore, we can save a token for such t when it is evaluated by CELF in round j but not by P-tree, such that when later t is counted in $u_{1...y_i}$ in round i, we will use the token to count t for free. Note that $t \notin (F'_i \cup F'_{j+1} \cup ... \cup F'_{i-1})$, for the same reason, t is not counted in any $y_{j'}$ for $j \le j' < i$, so the token must still available in round i. In summary, all vertices in U_1 are counted in x_i , and all vertices in U_2 can be counted by the saved tokens (charged to some previous x_i). Therefore, using amortized analysis, we have $\sum y_i \leq \sum x_i$.

Recall that $x_i = |F_i|$, and $|F_i'| \le 2y_i$. Since $\sum y_i \le \sum x_i$, we proved that $\sum |F_i'| \le \sum 2y_i \le 2 \sum x_i = 2 \sum |F_i|$.

Theorem 4.3 (P-tree Cost Bound). Given the same sketches $\Phi_{1..R}$, our seed selection based on P-tree will select the same seed set as CELF with $\tilde{O}(W_{CELF})$ work and $\tilde{O}(kD_{\Delta})$ span, where k is the number of seeds, W_{CELF} is the work (time complexity) by CELF, and D_{Δ} is the span to re-evaluate one vertex.

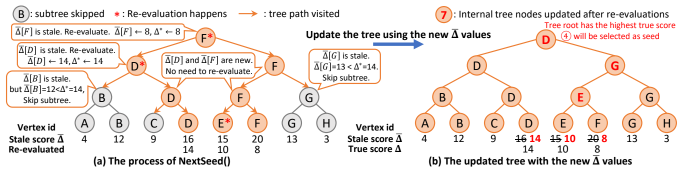


Figure 4: Seed selection based on *Win-Tree*. Each leaf stores a vertex id, and each internal node stores the vertex in its subtree with the highest (stale) score. (a) An example of finding the maximum score. For illustration purposes, we assume the parallel threads work at the same speed and all tree nodes on the same level are processed in parallel (in reality the threads run asynchronously in fork-join parallelism). Therefore, the subtree at ⑦ will see $\Delta^* = 14$ updated by ④, and this subtree will be skipped. (b) Updating the internal nodes with the new $\bar{\Delta}[\cdot]$ values. Finally, the root of the tree has the highest true score.

Thm. 4.3 can be proved by Thm. 4.2 and the cost bound of *P-trees*. For page limit, we present the proof in the full version of this paper.

Proof. According to analysis of *P-tree* [12], SPLITANDREMOVE have $O(\log n)$ work and $O(\log n)$ span, where n is the tree size. BATCHIN-SERT has $O(n' \log n)$ work and $O(\log n' \log n)$ span, where n' is the size of unordered insertion batch. For each seed selection round that evaluates a set of vertices F'_i , Alg. 4 mainly comes from 1) constructing the tree with $O(n \log n)$ work and polylogarithmic span, 2) $\log |F_i'| = O(\log n)$ splits and output the elements to an array, in $O(F'_i + \log^2 n)$ total work and polylogarithmic span, 3) $O(|F'_i|)$ reevaluations, and 4) inserting a batch back to the tree with size $|F'_i|-1$, costing $O(|F_i'| \log n)$ work and polylogarithmic span. Considering in CELF, the work involves 1) at least O(n) work to construct the priority queue, 2) $O(|F_i|)$ re-evaluations and 3) at least O(1) cost per evaluation in operating (popping and re-inserting) the priority queue. From Thm. 4.2, we know $\sum |F_i'| = O(\sum |F_i|)$, therefore, the overhead of P-tree compared to CELF is at most a polylogarithmic factor, which proves the $\tilde{O}(W_{CELF})$ bound.

For the span, note that *P-tree* needs $O(\log n)$ batches, each batch requires a split, and evaluates multiple vertices in parallel. Therefore, the span is $O(D_{\Delta})$ + polylog(n)) in each round. Combining with the polylogarithmic span for batch-insertion, and considering that there are $O(\log n)$ batches, we can prove the $\tilde{O}(D_{\Delta})$ span bound for each round.

4.2 Parallel Priority Queue: Win-Tree

While *P-tree* provides theoretical efficiency for seed selection, it maintains the total order of all vertices, which is not needed in many implementations for priority queues (e.g., a binary heap) and may cause performance overhead. Also, *P-tree* is an explicit tree that maintains the parent-child pointers, which causes additional space usage. We now propose a more practical data structure based on a winning tree that overcomes these two challenges, though it does not have the same bounds as in Thm. 4.3.

A classic *winning tree* (aka. tournament tree) is a complete binary tree with n leaf nodes and n-1 interior nodes. The data are stored in the leaves. Each interior node records the larger key of its two children. Since a winning tree is a complete binary tree, it can be stored *implicitly* in an array T[1..2n-1], which consumes smaller space. In our case, each tree node stores a vertex id (noted as t.id). The key of a node t is the (stale) score of this vertex, i.e., $\bar{\Delta}[t.id]$.

Algorithm 5: Seed Selection based on *Win-Tree*

Maintains: Global variable Δ^* : the highest true score evaluated so far A winning tree T with n leaf nodes each storing a record. For a tree node $t \in T$, we use the following notations: t.id: the id of the vertex stored in this node t.parent / t.left / t.right: the parent/left child/right child of node t. The winning tree is a max-priority-queue based on (stale) score $\bar{\Delta}[t.id]$ for each vertex.

```
1 Function FINDMax(tree node t, current seed set S, sketches \Phi_{1-R})
        if t.id = t.parent.id then stale \leftarrow false
                                                                 // evaluated by parent
        else stale \leftarrow true
        // Skip a subtree if the max is stale and is smaller than the current best
           score; no re-evaluation needed for the entire subtree
        if stale = true and \bar{\Delta}[t.id] < \Delta^* then return
4
        if stale = true then
                                      // Current value is stale, re-evaluation needed
5
              \bar{\Delta}[t.id] \leftarrow \text{MarginalGain}(S, t.id, \Phi_{1..R})
                                                                           // Re-evaluate
 6
             WRITEMAX(\Delta^*, \bar{\Delta}[t.id])
                                                          // update the best score so far
        if t is a leaf then return
        In Parallel:
             FINDMAX(t.left, S, \Phi_{1..R})
10
             FINDMax(t.right, S, \Phi_{1..R})
11
        // compare two branches and reset max
        if \bar{\Delta}[t.left.id] > \bar{\Delta}[t.right.id] then t.id \leftarrow t.left.id
12
         else t.id \leftarrow t.right.id
13
14 Function NextSeed(S, \Phi_{1..R})
         \Delta^* \leftarrow 0
15
        FINDMAX(T.root, S, \Phi_{1..R})
16
         s^* \leftarrow T.root.id
17
         return s*
```

Each interior node stores the id from its children with a larger score. As a result, the vertex at the root has the highest (stale) score.

To support CELF seed selection efficiently, we use the information in the internal nodes of *Win-Tree* to prune the search process. Suppose the best true score we evaluated so far is Δ^* , then if we see a subtree root t with a stale score smaller than Δ^* , we can skip the *entire subtree*. This is because all nodes in this subtree must have smaller stale scores than t.id, which indicates even smaller true scores. Although this idea is simple, we must also carefully maintain the *Win-Tree* structure, with the newly evaluated true scores. We presented our algorithm in Alg. 5 with an illustration in Fig. 4, and elaborate on more details below.

To implement NextSeed, we keep a global variable Δ^* to track the largest true score obtained so far, initialized to 0. The algorithm calls the FINDMax(t,...) subroutine starting from the root, which explores the subtree rooted at t. We first determine if the score of the vertex at t is stale: when t's id is the same as its parent, this vertex has been re-evaluated and has the true score ready (Line 2). Based on the node's status, there are three cases. First, if the score is stale, and is already lower than Δ^* , accordingly to the discussion above, we can skip the entire subtree and terminate the function (Line 4). Second, if the score is stale, but is higher than Δ^* , we have to re-evaluate the vertex *t.id*, since it may be a candidate for the seed (Line 6). We then use the atomic operation WRITEMAX to update Δ^* by this true score if it is better. The third case is when the score is not stale. Although no re-evaluation is needed on t.id, we still have to further explore the subtree, since with the newly evaluated score, the subtree structure (i.e., the ids of the internal nodes) may change. Therefore, in both cases 2 and 3, we recursively explore the two subtrees in parallel (Lines 9 to 11). After the recursive call returns, we re-compute the vertex id by comparing its left and right children, and use the one with a higher score.

Unlike *P-trees*, we cannot prove strong bounds for the number of re-evaluations in *Win-Tree*—since the parallel threads are highly asynchronous, the progress of updating Δ^* and pruning the search cannot be guaranteed. However, we expect *Win-Tree* to be more practical than *P-tree* for a few reasons. First, *Win-Tree* is a complete binary and can be maintained in an array, which requires smaller space (no need to store metadata such as pointers in *P-trees*). Second, the *P-tree* algorithm requires $O(\log n)$ batches and synchronizing all threads between batches. Such synchronization may result in scheduling overhead, while the *Win-Tree* algorithm is highly asynchronous. Most importantly, it does not maintain the total order, and the construction time is O(n) instead of $O(n \log n)$. In Sec. 5.3, we experimentally verify that although *Win-Tree* incurs more re-evaluations than *P-trees*, it is still faster in most tests.

5 Experiments

Setup. We implemented *PaC-IM* in C++. We run our experiments on a 96-core (192 hyperthreads) machine with four Intel Xeon Gold 6252 CPUs, and 1.5 TB of main memory. We use numact1 -i all in experiments with more than one thread to spread the memory pages across CPUs in a round-robin fashion. We run each test for 4 times and report the average of the last 3 runs.

We tested 17 graphs with information shown in Tab. 3. To understand the generality of our approaches, we include real-world graphs with a wide range of sizes and distributions, including **five billion-scale or larger graphs**. In addition to the commonly-used benchmarks of social networks, we also include web graphs, road networks, and *k*-NN graphs (each vertex is a multi-dimensional data point connecting to its *k*-nearest neighbors [76]). Solving IM on such graphs simulates the influence spread between websites (web graphs), geologically connected objects (road networks), and geometrically close objects (*k*-NN graphs). Based on graph patterns, we call social and web graphs *scale-free* graphs, and the rest *sparse* graphs. We symmetrize the directed graphs to make them undirected. *PaC-IM* (with compression) is the only tested system that can process the largest graph ClueWeb [52] with 978M vertices and 74B edges. Even with 1.5TB memory, *PaC-IM* can process Clueweb

		n	m	Influence	Notes
	EP	0.08M	0.81M	5332	Epinions1 [62]
	SLDT	0.08M	0.94M	6342	Slashdot [48]
	DBLP	0.32M	2.10M	1057	DBLP [79]
귵	YT	1.14M	5.98M	29614	com-Youtube [79]
Social	OK	3.07M	234M	1460000	com-orkut [79]
Š	LJ	4.85M	85.7M	376701	soc-LiveJournal1 [5]
	TW^*	41.7M	2.40B	11776629	Twitter [45]
	FT*	65.6M	3.61B	19198744	Friendster [79]
- q	SD*	89.2M	3.88B	15559737	sd_arc [52]
Web	CW*	978M	74.7B	-	ClueWeb [52]
-pe	GER	12.3M	32.3M	384	Germany [1]
Road	USA	23.9M	57.7M	370	RoadUSA [1]
	HT5	2.05M	13.0M	1018	HT [27, 76], k=5
-	HH5	2.05M	13.0M	2827	Household [27, 76], k=5
k-NN	CH5	4.21M	29.7M	355065	CHEM [29, 76], k=5
×	GL5	24.9M	157M	11632	GeoLife [76, 81], k=5
	COS5*	321M	1.96B	4753	Cosmo50 [46, 76], k=5

Table 3: Graph Information. n: number of vertices. m: number of edges. Influence: the maximum influence spread of 100 seeds selected by PaC-IM (R=256), Influence (R=256), and Ripples (R=256), rounded to integers. Most influence values are evaluated by 20000 simulations. The underline numbers on very large graphs are evaluated by 2000 simulations. "-": unable to evaluate within 50 hours using 2000 simulations.

only when $\alpha \le 0.25$ (4× or more compression ratio in sketches), which shows the necessity of compression.

We select k = 100 seeds in all tests. We use the IC model with a constant propagation probability p in the same graph. For scale-free graphs, we set p = 0.02. For sparse graphs, we set p = 0.2 as the vertices have smaller degrees. When comparing the *average* numbers across multiple graphs, we use the *geometric mean*.

Software Libraries. Our implementation uses ParlayLib [11] for fork-join parallelism and some parallel primitives (e.g., sorting). We use the *P-tree* implementation from the PAM library [68, 70]. When computing connectivity in sketch construction, we use the union-find implementation UniteRemCAS from ConnectIt [25].

Tested Algorithms. We tested both *P-tree* and *Win-Tree* for seed selection. In most cases, *Win-Tree* is more efficient in both time and space, so use *Win-Tree* as the default option in *PaC-IM*. We present more results comparing the two options in Sec. 5.3.

We compare to three existing parallel IM systems: InfuserMG [30], No-Singles [61] and Ripples [53, 54], and call them the baselines. As introduced in Sec. 2, InfuserMG uses a similar sketch-based approach as PaC-IM but does not support compression or parallel priority queues. No-Singles and Ripples both use the idea of Reverse Influence Sampling (RIS) [16]. In our experiments, Ripples is always better than No-Singles on running time and memory usage, so we only report the results of Ripples. We note that InfuserMG and our PaC-IM require undirected input graphs while Ripples make no such assumptions. We observe that InfuserMG and Ripples have severe scalability issue when the number of threads increase (see examples in Fig. 6). Hence, we report their shortest time among all the tested numbers of threads.

Each algorithm has a parameter that controls the solution quality: R for InfuserMG and our algorithms and ϵ (0 < ϵ \leq 0.5) for Ripples. The solution qualities increase with larger R or smaller ϵ . To ensure

		Relative Influence			Total Running Time (second)				Memory Usage (GB)				
		Ours	InfuserMG	Ripples	Ours ₁	$\mathbf{Ours}_{0.1}$	InfuserMG	Ripples	CSR	\mathbf{Ours}_1	$\mathbf{Ours}_{0.1}$	InfuserMG	Ripples
	EP	100%	99.9%	98.9%	0.29	0.49	0.38	4.14	0.01	0.14	0.06	0.17	0.22
	SLDT	100%	99.8%	99.1%	0.30	0.53	0.48	26	0.01	0.15	0.05	0.17	0.29
	DBLP	100%	99.0%	98.3%	0.35	0.37	0.86	41.9	0.02	0.48	0.10	0.67	0.29
a.	YT	100%	99.9%	98.2%	1.22	2.44	6.20	71.1	0.05	1.63	0.28	2.36	<u>1.54</u>
Social	OK	100%	100%	99.8%	8.79	39.6	80.3	329	1.77	6.13	2.44	8.07	78.5
Š	LJ	100%	99.9%	99.4%	6.00	20.5	61.3	123	0.68	5.98	1.63	10.5	20.4
	TW	100%	100%	-	93.4	378	639	6258	18.2	63.0	25.9	103	729
	FT	100%	100%	-	128	609	1973	-	27.4	<u>97.9</u>	39.9	161	-
Web	SD	100%	97.1%	-	150	627	1684	-	29.6	125	44.8	211	-
×	CW	100%	-	-	-	9776	-	-	564	-	738	-	-
ad	USA	100%	74.7%	92.8%	14.6	13.7	53.1	13628	0.61	25.8	5.05	49.0	46.4
Ro	GER	100%	70.4%	94.4%	9.35	8.53	26.7	2334	0.33	13.3	2.58	25.1	23.0
	HT5	100%	85.7%	94.8%	0.72	0.68	2.73	47.5	0.11	1.37	0.24	1.93	0.83
7	HH5	100%	79.5%	97.6%	2.28	2.11	8.26	53.9	0.11	3.00	0.51	4.26	1.22
k-NN	CH5	100%	92.3%	97.8%	3.52	5.32	124	39.1	0.25	4.87	1.05	8.77	1.77
¥	GL5	100%	76.9%	98.9%	19.5	17.9	116	637	1.36	27.6	6.02	51.6	8.71
	COS5	100%	37.7%	-	348	284	2319	-	17.0	<u>355</u>	66.1	666	-

Table 4: Running time, memory usage, and influence spread (normalized to the maximum) of all tested systems on a machine with 96 cores (192 hyperthreads). "-": out of memory (1.5 TB) or time limit (3 hours). Ours₁ is our implementation with *Win-Tree* without compression. Ours_{0.1} is our implementation with $\alpha = 0.1$ (10× compression for sketches). *InfuserMG* [30] and *Ripples* [53, 54] are baselines. We report the *best time* of *InfuserMG* and *Ripples* by varied core counts (the scalability issue of *InfuserMG* and *Ripples* are shown in Fig. 6). CSR is the memory used to store the graph in CSR format (see more in Sec. 5.1). The bold numbers are the fastest time/smallest memory among all implementations on each graph. The underlined numbers in memory usage are the smallest memory among systems that do not use compression (Ours₁, *Ripples* and *InfuserMG*). A heatmap of the full result is in Fig. 1.

fairness, when comparing running time, we guarantee that PaC-IM always gives a better solution than the baselines. For InfuserMG and PaC-IM, we set the number of sketches R=256. Our solution using R=256 is on average more than 99% of the quality when using $R=2^{15}$, which is consistent with the observation in the StaticGreedy paper [21] (see our full version for quality analysis). For Ripples, smaller ϵ means better accuracy but more computation. We tested Ripples under ϵ in range 0.13–0.5 as used in their paper. PaC-IM under R=256 yields about the same solution quality as $\epsilon=0.13$ (the best tested setting in their paper). When reporting time, we use $\epsilon=0.5$ since this gives the fastest running time, and the quality is still reasonably high (at least 93% of our best influence).

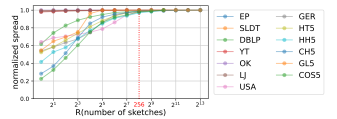


Figure 5: Spread curves for our algorithm. In each plot, *x*-axis is the number of sketches (R) and *y*-axis is the spread normalized to the maximum spread of each graph. Higher is better.

We observe that on sparse graphs, the influence spread of InfuserMG is only 38–92% of the best achieved by PaC-IM and Ripples. Although theoretically PaC-IM and InfuserMG should give the same output (assuming a fixed seed for the random number generator), we note that InfuserMG uses many optimizations that sacrifices solution quality. We tried to increase R and various other attempts in InfuserMG, but they did not improve the solution quality. Therefore, we keep the same value R = 256 for both PaC-IM and InfuserMG.

5.1 Overall Time and Space

Tab. 4 shows the running time, memory usage, and normalized influence spread of all systems. Ours₁ and Ours_{0.1} are our PaC-IM with $\alpha=1$ (no compression) and $\alpha=0.1$ ($10\times$ sketch compression), respectively. To better illustrate the tradeoff between time and space for all systems, we present a heatmap in Fig. 1, where all the numbers (time and space) are normalized to Ours₁. Ours_{0.1} is the only algorithm that can process the largest graph CW [52]. With similar spread quality, $Ours_1$ is faster than all baselines on all graphs, and $Ours_{0.1}$ is just slower than InfuserMG on the two smallest graphs. Meanwhile, $Ours_{0.1}$ has the smallest space usage on all graphs. The advantage of PaC-IM is more significant on larger graphs, both in time and space.

Running Time. PaC-IM is significantly faster than the baseline algorithms on almost all graphs. As mentioned, we report the best running time among all tested core counts for InfuserMG and Ripples, since they may not scale to 192 threads (see Fig. 6 and Sec. 5.2). Even so, PaC-IM is still faster on all graphs. On average, $Ours_1$ is $5.7 \times faster$ than InfuserMG and $56 \times faster$ than Ripples. With compression, $Ours_{0.1}$ is slightly slower than $Ours_1$, but is still $3.2 \times faster$ than InfuserMG and $32 \times faster$ than Ripples.

In general, the compression in PaC-IM saves space by a tradeoff of increasing running time. When $\alpha=1$, the CC sizes for all vertices are stored in the sketches, and a re-evaluation only needs a constant time to query on each sketch. When $\alpha=0.1$, each query involves a search to either find a center, or visit all connected vertices, which roughly costs $O(1/\alpha)$ on each sketch. Indeed, on all scale-free graphs, $Ours_{0.1}$ takes a longer time than $Ours_1$. Interestingly, on most of the sparse graphs, we observe that $Ours_{0.1}$ is also faster than $Ours_1$. This is because seed selection only takes a small fraction of the total running time (except for CH5), so the

slow-down in the seed selection step is negligible for the overall performance. Meanwhile, avoiding storing O(Rn) connectivity sizes reduces memory footprint and makes the sketching step slightly faster, which overall speeds up the running time.

Memory Usage. We show space usage of PaC-IM and baselines in Fig. 1 and 7 and Tab. 4. We also show the size of representing the graph in standard Compressed Sparse Row (CSR) format as a reference in Tab. 4 and Fig. 7, which roughly indicates the space to store the input graph. CSR uses 8 bytes for each vertex and each edge. Using $\alpha=0.1$, PaC-IM uses the least memory on all graphs. Even without compression, our algorithm uses less memory than the baselines on 10 out of 16 graphs. Note that although the compression rate for sketches is $10\times$ in $Ours_{0.1}$, the total space also includes the input graph and the data structure for seed selection. Therefore, we cannot directly achieve a $10\times$ improvement in space. In most cases, the total memory usage is about $5\times$ smaller.

Summary. Overall, PaC-IM has better performance than the baselines in both time and space. We note that the space usage of Ripples can be (up to $3\times$) better than $Ours_1$ on certain graphs (but still worse than $Ours_{0.1}$), but in these cases the running time is also much longer (by $10\times$ or more). On scale-free graphs, $Ours_{0.1}$ is $2.5\times$ slower than $Ours_1$ on average, but uses $3\times$ less space. On sparse graphs, $Ours_{0.1}$ is almost always better in both time and space.

5.2 Scalability

We also study the scalability of all systems based on core counts on six representative graphs. Fig. 6 presents the running time with varying core counts *P*. We separate the time for sketch construction (the "sketch time") and seed selection (the "selection time") to analyze the two components independently. For *PaC-IM*, both sketch and selection time decrease with more cores and achieve almost linear speedup. Unfortunately, such scalability trend is not always true for both *InfuserMG* and *Ripples*. *InfuserMG* and *Ripples* scale to 8–16 cores on many graphs, but the performance may get worse with more cores. On LJ, SD, and GE, the running time of *InfuserMG* is better than *PaC-IM* when running sequentially, but due to poor scalability, it becomes worse than *PaC-IM* when running on more than 1 core. This indicates that a major performance gain of *PaC-IM* is better scalability.

The scalability curves also indicate the necessity of our new parallel data structures for seed selection proposed in Sec. 4. For Ours_{0.1}, since the re-evaluation time becomes longer than Ours₁ due to compression, the selection time may be longer than sketch construction, where parallelizing seed selection is important in improving the overall performance. Even for Ours₁ where the selection time is a small fraction of the total running time when P = 1, the sequential selecting time is still much longer than the total running time when P > 16. This means that if this step is not well-parallelized, it will become the performance bottleneck when *P* is large. For both InfuserMG and Ripples, the sketch time parallelizes better than the selection time. This is because both algorithms use well-parallelized algorithms to construct sketches. For example, InfuserMG uses a standard coloring [58, 67] idea for parallel connectivity. However, for seed selection, both systems use sequential CELF, and only parallelize the re-evaluation on R sketches. When the core count

becomes comparable to R, such approach may cause scheduling overhead and have negative impacts on the performance.

5.3 Analysis of the Proposed Techniques

Next, we evaluate the two proposed techniques in this paper: sketch compression and data structures for seed selection.

Compression. We evaluate the running time and memory usage of different compression ratios by controlling the parameter α , and present the results in Fig. 7. The gray dashed horizontal line represents the CSR size of each graph, which is the space to store the input. Compression always reduces memory usage, but may affect running time differently. The memory usage always decreases with the value α decreases. As mentioned previously, the actual compression rate can be lower than $1/\alpha$, since our compression only controls the memory for storing sketches, and there are other space usage in the algorithm. Roughly speaking, using $\alpha=0.05$ can save space on graphs by up to $8\times$ and shrink the space very close to the input graph size.

Compression affects the running time differently for sketch construction and seed selection. Smaller α indicates longer running time in seed selection, but may improve the sketch time slightly due to the reduced memory footprint. The effect of compression on total running time depends on the ratio of sketching and selection time, where scale-free and sparse graphs exhibit different patterns. In the sampled graphs, scale-free graphs usually have one or several large CCs since they are dense, while the sparse graphs usually have many small CCs. This can also be seen by the overall influence in Tab. 3: even we use smaller p on scale-free networks, the total influence is still much larger than the sparse graphs. On scale-free graphs, due to large CCs on the sampled graph, selecting any seed in this CC may significantly lower the score of other vertices, leading to much large total re-evaluations than sparse graphs. As a result, smaller α causes a clear time increase for seed selection on scale-free networks since each re-evaluation becomes slower, while the compression only has a small impact on most of the sparse

P-tree vs. *Win-Tree*. We now experimentally study both data structures for seed selection. Recall that our goal is to achieve high parallelism without introducing much overhead on work. In particular, *P-tree* has the theoretical guarantee that the total number of re-evaluations is no more than twice of the sequential CELF. To evaluate the work overhead is small, we plotted the number of re-evaluations for both data structures compared to CELF Fig. 8(a). For each graph, we count the number of re-evaluations by CELF as x, the number by *P-tree* as y_1 , and the number by *Win-Tree* as y_2 , and draw all $(x, y_1/x)$ as blue round points and all $(x, y_2/x)$ as orange triangle points. The number of re-evaluations by *P-trees* is very close to that by CELF $(1.03 \times \text{ on average})$, while *Win-Trees* require slightly more $(1.7 \times \text{ on average})$. The result is consistent with our theoretical analysis.

As mentioned in Sec. 4.2, we expect *Win-Trees* to perform better in practice due to various reasons. To study this, we plotted Fig. 8(b) as a comparison for selection time between *P-tree* and *Win-Tree* under different compression ratios. Each data point represents a graph, and the value is the ratio of selection time between *P-tree* and *Win-Tree*. The average ratios are always larger than 1, indicating

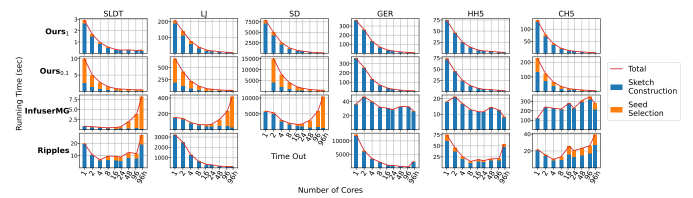


Figure 6: Running time using different core counts for different IM algorithms. In each plot, the *x*-axis shows core counts (96h means 96 cores with hyperthreading) and the *y*-axis is running time in seconds.

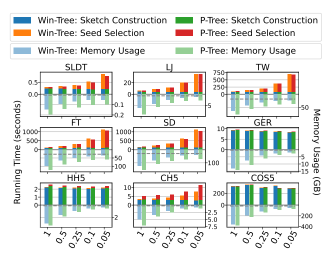


Figure 7: Running time and memory with different values of α . The x-axis represents the compression rate. The growing up y-axis represents the running time (in seconds) of the sketching and selecting process. The growing down y-axis represents the total memory usage (in GB). The gray horizontal line represents the CSR size of each graph, which is the basic memory we need to load the graph.

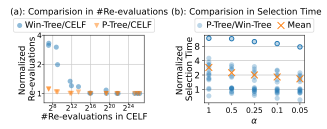


Figure 8: Compare *P-tree* and *Win-Tree* in the number of re-evaluations and selection time.

better performance for *Win-Tree*, but the advantage decreases as α becomes smaller (higher compression). This is because when α is large, the re-evaluation is fast, and the major time is on the tree operations, where *Win-Trees* is more advantageous due to the reasons mentioned in Sec. 4.2. With higher compression (small α), the re-evaluation becomes more expensive. Since the *P-tree* evaluates fewer vertices, the overall selection time is more likely to be better.

		n	CELF	P-Tree	Win-Tree
	HP	12008	7936	7964	8322
	EP	75879	57980	58010	58398
	SLDT	77360	66618	66649	67035
7	DBLP	317080	3255	3269	4075
social	YT	1134890	656014	656068	656727
SC	OK	3072627	3066983	3066995	3067514
	LJ	4847571	3609995	3610105	3611583
	TW	41652231	41200415	41200437	41200774
	FT	65608366	61946405	61946434	61947064
q	SD	89247739	84554009	84576755	84576236
web	CW				
pe	USA	23947348	143	159	465
road	GER	12277375	139	155	416
	HT5	928991	516	536	1007
7	HH5	2049280	3931	3943	4717
k-NN	CH5	4208261	628918	629501	653157
Ä	GL5	24876978	11042	11430	12996
	COS5	321065547	431	449	1264

Table 5: Number of Re-evaluations

To further understand this, we focus on the five topmost points (circled) in Fig. 8(b): those data points are from COS5, where P-tree is 8–12× slower than Win-Tree. For COS5, the number of vertices n is large, but only hundreds of vertices are re-evaluated in total. Thus, the seed selection time is dominated by constructing the data structures, i.e., $O(n \log n)$ work for P-tree to maintain total ordering, and O(n) for Win-Tree that is a lot faster. Since the number of re-evaluations is small, most of the vertices in the tree are never touched, and thus maintaining their order wastes the work.

In Fig. 7, we also compare the time and space between P-tree and Win-Tree. Similar to the discussions above, when α is small, Win-Tree is almost always faster than P-tree, but P-tree may perform better when α is large. However, the advantage is quite small, since the number of re-evaluations of Win-Tree is still close to P-tree (see Fig. 8(a)). Win-Tree also uses smaller memory than P-tree. As discussed in Sec. 4.2, this is because P-tree needs to explicitly maintain tree pointers and balancing criteria, while each Win-Tree node only needs to store the vertex id (2n integers in total).

In summary, P-tree almost always incurs fewer re-evaluations than Win-Tree, but is worse in running time in most cases, especially with low compression (α close to 1). When higher compression (smaller α), P-tree can perform slightly better. Win-Tree is also more space-efficient than P-tree. Based on these observations, we always use Win-Tree as the default data structure in PaC-IM, but also provides the interface for users to choose P-trees.

6 Related Work

Influence Maximization (IM). IM has been widely studied for decades with a list of excellent surveys [4, 6, 49, 60, 83] that review the applications and papers on this topic. IM is also of high relevance of the data management community, and many excellent papers published recently regarding benchmarking [2, 56], new algorithms [37], new propagation models and applications [8, 38, 73, 78], and interdisciplinary extensions [80, 84].

According to [49], IM algorithms can roughly be categorized into three methodologies: simulation-based, proxy-based, and sketch-based. Among them, (Monte Carlo) simulation-based approaches (e.g., [32, 40, 47, 75, 82]) are the most general and apply to the most settings (i.e., graph types and diffusion models); however, they do not take advantage of the specific settings, so generally their performance is limited. Proxy-based approaches (e.g., [19, 39, 41, 42, 51, 59]) use simpler algorithms/problems (e.g., PageRank or shortest-paths) to help to solve IM. While the solutions can be fast in practice, their solution quality has no theoretical guarantees. Sketch-based solutions, as we introduced in Sec. 1 and 2, generally have good performance and theoretical guarantees, and they are specific to diffusion models. As a sketch-based solution, *PaC-IM* mainly focuses on the IC model (although the parallel data structures in Sec. 4 are general).

Sketch-based algorithms can further be categorized into forward (influence) sketches and reverse (reachable) sketches [49]. As the names suggest, forward sketches record the influence that each vertex can propagate to in sampled graphs. Most algorithms [20–22, 30, 40, 57] mentioned in this paper, including PaC-IM, use forward sketches. Reverse sketches find a subset of vertices T and keep the sets of vertices that can reach each vertex in T. Many IM algorithms (e.g., [16, 71, 72, 74]), including No-Singles [61] and Ripples [53, 54] that we compared to, use reverse sketches. These algorithms can trade off (lower) solution quality for (better) performance/space, by adjusting the size of T.

Space-Efficient Connectivity. We are aware of a few algorithms that can compute graph connectivity using o(n) space [7, 17, 28, 43], which share a similar motivation with PaC-IM. However, PaC-IM does not require computing connectivity in o(n) space, but only requires storing it in O(n) space. Hence, the goal here is essentially different, although these approaches are inspiring.

7 Conclusion and Future Work

In this paper, we address the scalability issues in existing IM systems by novel techniques including sketch compression and parallel data structures for seed selection. Our sketch compression technique (in Sec. 3) applies to the IC model and undirected graphs, and avoids the O(Rn) space usage in existing systems, which allows PaC-IM to run on much larger graphs, without sacrificing the performance much. Our new data structures, P-tree and Win-Tree (in Sec. 4), are

the first solutions to parallelize the CELF seed selection step, to the best of our knowledge, which is general to any submodular diffusion models.

In addition to new algorithmic ideas, our new techniques are carefully analyzed (see Thm. 3.1 and 4.3) and have good theoretical guarantees regarding work, parallelism, and space usage. These analyses not only lead to good practical performance (see Fig. 1), but also helps to understand how the techniques interplay. The techniques are also experimentally verified in Fig. 6 to 8. The theory and our careful implementation also lead to stable speedup with increasing core counts (see Fig. 6). This means that *PaC-IM* will likely have even better performance in the future given stronger processors with more core counts.

We note that our approaches based on *P-tree* and *Win-Tree* can be much more general than parallelizing CELF seed selection in IM. In fact, CELF is a general greedy approach to accelerate optimization problems with submodular objective functions [44, 47, 55]. We leave this as future work.

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