Efficient Algorithms for Density Decomposition on Large Static and Dynamic Graphs

ABSTRACT

Locally-densest subgraph (LDS) decomposition is a fundamental operation in graph analysis that finds numerous applications in various domains, including community detection, fraud detection, graph querying, and graph visualization. However, the LDS decomposition is computationally challenging for both static and dynamic graphs. Furthermore, the LDS decomposition often produces an excessive number of dense subgraph layers, leading to the unnecessary separation of tightly-connected subgraphs. To address these limitations, an alternative concept called density decomposition was proposed, which can generate a more reasonable number of dense subgraph layers. However, the state-of-the-art algorithm for density decomposition requires $O(m^2)$ time (*m* is the number of edges of the graph), which is very costly for large graphs. In this paper, we conduct an in-depth investigation of density decomposition and propose efficient algorithms for computing it on both static and dynamic graphs. First, we establish a novel relationship between density decomposition and LDS decomposition, allowing us to leverage existing LDS algorithms for density decomposition. Second, we propose a novel algorithm to compute the density decomposition on static graphs, called Flow++, based on carefully designed network flow and divide-and-conquer techniques. The striking feature of our Flow++ algorithm is that its time complexity is $O(m^{3/2} \log p)$ (p is often a very small constant in real-world graphs), thus significantly reducing the complexity of the previous algorithm. Third, for dynamic graphs, we prove a density decomposition updating theorem, based on which we develop a novel dynamic algorithm with O(m) time complexity to maintain the density decomposition. Extensive experiments on several large real-world graphs demonstrate the high efficiency, scalability, and effectiveness of the proposed algorithms.

1 INTRODUCTION

Real-world graphs are typically overall sparse but contain small dense subgraphs. Discovering these dense subgraphs has a wide range of applications, including community detection [1, 19, 31, 39, 51, 67], fraud detection [6], graph querying [21, 38], and graph visualization [3, 68]. Various models have been introduced to characterize dense subgraphs, such as the clique [27], k-core [4, 41, 45, 52], k-truss [22, 37], locally densest subgraph [42, 49, 61], k-edge connected subgraph [15, 16], k-plex [24, 57], k-defective clique [23], k-club [46], and so on.

Among them, k-core perhaps stands as the most popular dense graph model due to its concise definition and linear time complexity. However, the k-core decomposition, defined solely based on degrees, falls short of effectively capturing the density structure of a graph, where density is measured as the ratio of the number of edges to

Permission to make digital or hard copies of all or part of this work for personal or classroom use is granted without fee provided that copies are not made or distributed for profit or commercial advantage and that copies bear this notice and the full citation on the first page. Copyrights for components of this work owned by others than ACM must be honored. Abstracting with credit is permitted. To copy otherwise, or republish, to post on servers or to redistribute to lists, requires prior specific permission and/or a fee. Request permissions from permissions@acm.org.

Conference'17, July 2017, Washington, DC, USA © 2023 Association for Computing Machinery. ACM ISBN 978-x-xxxx-xxxx-x/YY/MM...\$15.00 https://doi.org/10.1145/nnnnnnn.nnnnnn the number of vertices. For example, as shown in Fig. 1a, the top layer of core decomposition C_3 contains 23 edges and 11 vertices, yielding a density of 23/11. Nevertheless, the subgraph with the highest density in this graph is B_2 with a density of 15/7. Clearly, the core decomposition is not density-based, and thus may not accurately capture the dense subgraphs.

To address this limitation, Tatti [61] introduced a concept of locally-densest subgraph (LDS) decomposition, dividing the graph into a set of locally-densest subgraphs with hierarchical features, i.e., a smaller subgraph must be contained in a larger subgraph, and the density of subgraphs strictly decreases as the size of the subgraph increases. LDS decomposition ensures that the smallest subgraph is the densest subgraph in the whole graph. However, despite its ability to fully reflect the density structure, LDS decomposition has three notable drawbacks: (i) The worst-case time complexity for computing it is as high as $O(n^2m)$, where n and m denote the number of vertices and edges, which is costly for large graphs; (ii) The number of decomposed locally-densest subgraphs may be excessive and can only be bounded by O(n), typically leading to unnecessary separation of densely-connected structures. For example, in Fig. 1a, B_0 , B_1 , and B_2 are locally-densest subgraphs. They separate the tightly connected subgraph $\{v_8, \ldots, v_{15}\}$, which is not necessary; (iii) It is very challenging to exactly maintain the locally-densest subgraphs on dynamic graphs due to the strictness of its definition. To the best of our knowledge, there is currently no efficient algorithm for maintaining locally-densest subgraph decompositions, except for recomputing the decomposition from scratch.

On the other hand, Borradaile et al. [14] introduced an alternative density-based decomposition, known as density decomposition, which partitions the vertices based on their indegree and connectivity in a so-called egalitarian orientation [13]. Similar to the LDS decomposition, the density decomposition also exhibits a hierarchical structure, with the density of subgraphs decreasing as the size of layers increases. Consequently, the smallest layer of this decomposition also has extremely high density. In [14], Borradaile et al. proposed an $O(m^2)$ time algorithm to compute the density decomposition, which is clearly costly for large graphs. Moreover, Borradaile et al. did not establish a clear relationship between the density decomposition and other dense subgraph models.

Motivated by this, we for the first time conduct a thorough investigation of density decomposition, providing insights and a comprehensive understanding of its properties and applications. Specifically, we first establish a novel and interesting connection between density decomposition and LDS decomposition, which enables us to use the existing highly-optimized LDS decomposition algorithm to compute the density decomposition [42]. Then, we propose a new algorithm based on carefully designed network flow and divide-and-conquer techniques to efficiently compute the density decomposition with reduced time complexity. Subsequently, we also develop novel algorithms with linear time complexity to efficiently maintain the density decomposition on dynamic graphs. Finally, we conduct extensive experiments to evaluate our algorithms using several large real-life graphs, and the results demonstrate the efficiency, scalability, and effectiveness of the proposed algorithms. In summary, the main contributions of this paper are as follows.

New theoretical results on density decomposition. We show that each layer of the density decomposition is exactly a locallydensest subgraph, making it part of the LDS decomposition. This crucial relationship reveals the fundamental similarity between these two decompositions. On the other hand, their key distinction lies in that density decomposition, through a round-up relationship, can naturally group locally-densest subgraphs with similar densities. This grouping significantly reduces the number of subgraphs, addressing the issue of unnecessary layers in the LDS decomposition. In particular, we show that the number of non-trivial layers generated by the density decomposition is equal to the pseudoarboricity of a graph, denoted as p, typically a small number. In addition, we also reveal an interesting approximation relationship between density decomposition and k-core decomposition, indicating that core decomposition can be considered as a 2-approximation of the density decomposition.

Novel algorithms for computing density decomposition. We first propose a novel algorithm, called Flow, to compute the density decomposition, based on a carefully-designed network flow technique. Compared to the state-of-the-art $O(m^2)$ -time algorithm, Flow achieves a superior time complexity of $O(p \cdot m^{3/2})$. To further improve the efficiency, we use a powerful divide-and-conquer technique, developing a new network flow-based algorithm, called Flow++. We show that such a new algorithm further reduces the time complexity to $O(m^{3/2} \log p)$.

Novel algorithms for maintaining density decomposition. We for the first time study the problem of maintaining the density decomposition on dynamic graphs. We discover a density decomposition update theorem, revealing that the insertion or deletion of an edge requires analyzing changes in just one layer of the density decomposition. Based on this, we develop an insertion algorithm, called Insert to handle edge insertions. This algorithm only needs to invoke the Breadth-First Search (BFS) algorithm at most twice, yielding a worst-case time complexity of O(m). For edge deletion, we develop the Delete algorithm along with its enhanced version, Delete++, which incorporates several carefully designed pruning strategies. Both of these algorithms have O(m) time complexity, thus they are very efficient for handling large dynamic graphs.

Extensive experiments. We conduct comprehensive experiments to evaluate the proposed algorithms using 8 real-world datasets. The results are summarized as follows: (1) Both the Flow and Flow++ algorithms are substantially faster than the state-of-the-art algorithm by at least two orders of magnitude for density composition. Moreover, even when compared to the highly-optimized LDS decomposition algorithm [42], our Flow++ can still achieve a remarkable speed improvement of one order of magnitude. (2) For density decomposition maintenance, both our insertion and deletion algorithms outperform the baselines by at least 5 orders of magnitude. We also conduct three case studies to show the effectiveness of density decomposition. The results indicate that core decomposition is likely to result in a higher density of lower-level cores than higher-level cores. Conversely, density-based decomposition ensures a monotonically decreasing trend in density for each layer, facilitating the successful discovery of dense subgraphs. Moreover, LDS decomposition can be well approximated by density decomposition while significantly reducing the number of redundant layers. Our experimental results demonstrate that density decomposition serves as a computationally efficient model for dense subgraph decomposition, accurately capturing the density structure of the graph. For reproducibility purposes, we release the source code of this paper at an anonymous link: https://anonymous.4open.science/r/DD-83C6.

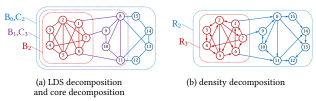


Figure 1: Example graph G = (V, E) and its decompositions.

2 PRELIMINARIES

Consider an undirected graph G = (V, E), where V and E are the vertex set and edge set respectively. Let n = |V| and m = |E| be the number of vertices and edges in G. An orientation of G can be obtained by assigning a direction to every edge in E, resulting in a directed graph $\vec{G} = (V, \vec{E})$. For example, the directed graph in Fig. 1b is an orientation of the undirected graph in Fig. 1a. To eliminate confusion, we use angle brackets $\langle u, v \rangle$ to represent a directed edge in \vec{G} , while round brackets (u, v) denote an undirected edge in G. For a directed (resp., undirected) graph G and a vertex u, we use $d_u(G)$ (d_u for brevity) to denote the indegree (resp., degree) of u. A path in a directed graph $\vec{G} = (V, \vec{E})$ is a sequence of vertices $s = v_0, v_1, ..., v_{l-1}, t = v_l$, where $\langle v_{i-1}, v_i \rangle \in \vec{E}$, for i = 1, ..., l. We denote such a path as $s \rightsquigarrow t$, and the length of $s \rightsquigarrow t$ is l. If there exists a path $s \rightsquigarrow t$, we say that s can reach t. Besides, if an edge $\langle u, v \rangle$ is reversed, then it becomes $\langle v, u \rangle$. When a path is reversed, all edges in the path undergo a reversal. Before introducing the density decomposition, we first give the definitions of reversible path and egalitarian orientation as follows.

DEFINITION 1. (Reversible path) Given a graph G and its orientation \vec{G} , for a path $s \rightsquigarrow t$ in \vec{G} , if $d_t - d_s \ge 2$, we call that the path $s \rightsquigarrow t$ is a reversible path.

DEFINITION 2. (Egalitarian orientation) Given a graph G and its orientation \vec{G} , if there is no reversible path in \vec{G} , then \vec{G} is an egalitarian orientation.

Intuitively, an egalitarian orientation distributes the indegree of vertices in the most equitable manner, i.e., minimizing the indegree difference between vertices as much as possible. Note that if reverse a reversible path $s \leadsto t, d_s$ increases by 1, d_t decreases by 1, and the indegree of other vertices does not change, making the indegree difference between s and t reduced by 2. When no reversible path exists, the indegree difference can not be reduced anymore. Based on the egalitarian orientation, we give the definition of density decomposition which was originally proposed by Borradailie et al. [14] as follows.

DEFINITION 3. (Density decomposition) [14] Given an undirected graph G = (V, E) and its arbitrary egalitarian orientation $\vec{G} = (V, \vec{E})$, the density decomposition of G is a set of subgraphs, denoted by $\mathcal{R} = \{R_i\}$, where for any non-negative integer i, $R_i \triangleq \{u \in V | d_u(\vec{G}) \geq i \text{ or } u \text{ can reach a vertex } v \text{ with } d_v(\vec{G}) \geq i \}$.

Based on Definition 3, we give a definition of *integral dense* number as follows.

DEFINITION 4. (Integral dense number) For a vertex $u \in R_i \setminus R_{i+1}$, the integral dense number (IDN) of u is defined as $\overline{r}_u = i$.

Example 1. Taking the egalitarian orientation shown in Fig. 1b as an example, only v_5 has the indegree of 3. Consequently, R_3 contains v_5 and all vertices that can reach it, i.e., $R_3 = \{v_1, \ldots, v_7\}$. For vertices v_1, \ldots, v_{14} , their indegrees are larger than or equal to 2, and v_{15} can

reach them, thus R_2 contains all vertices. Besides, R_1 and R_0 also contain all vertices, and for any integer i > 3, $R_i = \emptyset$. For the vertex $u \in R_3 \setminus R_4$, \overline{r}_u is equal to 3, while for the vertex in $u \in R_2 \setminus R_3$, we have $\overline{r}_u = 2$.

Borradaile et al. [14] shown that density decomposition exhibits a hierarchical structure, i.e., for two non-negative integers i>j, R_i is a subset of R_j . Let p be the largest integer such that R_p is non-empty. According to [7, 63], we can derive that p equals the p-seudoarboricity of G, denoted by p(G) or p for brevity. The pseudoarboricity is a classic metric to measure the sparsity of a graph which is defined as the minimum number of edge-disjoint pseudoforests into which the edges of a graph can be divided [48]. Here a pseudoforest is a graph in which each connected component contains at most one cycle. As indicated in [10], the pseudoarboricity of real-world graphs is often very small.

The number of layers of the density decomposition, from $R_0 = V$ to $R_{p+1} = \emptyset$, is p+2, and the number of non-trivial layers, from R_1 to R_p , is p. Note that although a graph G may have many different egalitarian orientations, the density decomposition of G is unique [14]. Furthermore, Borradaile et al. [14] also proved the following result.

LEMMA 1. [14] Given an undirected graph G = (V, E) and its arbitrary egalitarian orientation $\vec{G} = (V, \vec{E})$, if $u \in R_i \setminus R_{i+1}$, then $d_u(\vec{G}) \in \{i, i-1\} = \{\bar{r}_u, \bar{r}_u - 1\}$. Additionally, if $(u, v) \in E$ and $\bar{r}_u < \bar{r}_v$, then in \vec{G} , (u, v) must be oriented as $\langle v, u \rangle$ instead of $\langle u, v \rangle$.

Lemma 1 shows that the indegree of each vertex is either equal to its IDN or IDN minus one, and the edges between layers must be oriented towards the endpoint with the smaller IDN. With these definitions, we formulate our problems as follows.

Problem definition. Given an undirected graph G, our goal is to efficiently compute the density decomposition of G, i.e., computing the IDNs for each vertex in G. For graphs with dynamic updates, we aim to maintain the density decomposition (i.e., maintain the IDNs for all vertex) when an edge is inserted or deleted.

By Definition 3, Definition 4, and Lemma 1, the key to compute the density decomposition is to find an egalitarian orientation. In the following sections, we will develop several more efficient algorithms to tackle this problem.

3 NEW THEORETICAL INSIGHTS

In this section, we first establish an interesting connection between density decomposition and locally-densest subgraph (LDS) decomposition, allowing us to leverage existing highly-optimized LDS decomposition algorithms, such as [42] and [35], for computing density decomposition. Then, we will show a close connection between density decomposition and core decomposition.

3.1 Connection with LDS Decomposition

Let $X \subseteq V$ be a non-empty subset of the vertex set. Define the *induced edges* of X as $E(X) = \{(x, y) \in E | x, y \in X\}$, and the *induced subgraph* of X in G is (X, E(X)). The *density* of X is characterized by $\rho(X) = |E(X)|/|X|$. The subset $X \subseteq V$ that maximizes the density $\rho(X)$ is recognized as the *densest subgraph* of G.

For two vertex subsets X and Y with $X \cap Y = \emptyset$, we define the *cross edges* between X and Y as $E_X(X,Y) = \{(x,y) \in E | x \in X, y \in Y\}$, and the *additional edges* from X with respect to Y as $E_X(X,Y) = E(X) \cup E_X(X,Y)$, representing the increment in edges upon incorporating X into Y. Based on this concept, we define the outer density as follows.

Definition 3.1. **(Outer density)** If $X \cap Y = \emptyset$, the *outer density* of a non-empty vertex set X with respect to a disjoint set Y is defined as $\rho(X,Y) \triangleq |E_{\Delta}(X,Y)|/|X|$. If $X \cap Y \neq \emptyset$, the focus is on the subset of X not included in Y, i.e., $X \setminus Y$. In this case, we define $E_{\times}(X,Y) \triangleq E_{\times}(X \setminus Y,Y)$, $E_{\Delta}(X,Y) \triangleq E_{\Delta}(X \setminus Y,Y)$, and $\rho(X,Y) \triangleq \rho(X \setminus Y,Y)$.

EXAMPLE 2. Consider the graph G in Fig. 1a. The density of B_2 is $\rho(B_2) = 15/7$, which is clearly the densest subgraph of G. The outer density of B_1 with respect to B_2 is calculated as $\rho(B_1, B_2) = \rho(B_1 \setminus B_2, B_2) = (3+5)/4 = 2$.

With these concepts, we give the definition of locally-densest subgraph (LDS) [61] as follows.

DEFINITION 5. **(Locally-densest subgraph)** Given an undirected graph G = (V, E) and a vertex set $W \subseteq V$, W is a locally-densest subgraph if there is no $X \subseteq W$ and Y disjoint from W, such that $\rho(X, W \setminus X) \leq \rho(Y, W)$.

To provide a more intuitive understanding, we rephrase the definition of LDS as follows: W is locally-densest if and only if $W = \emptyset$, or W = V, or $\min_{X \subseteq W} \rho(X, W \setminus X) > \max_{Y \cap W = \emptyset} \rho(Y, W)$. In other words, the minimum-density subgraph X within W still has a higher outer density compared to the maximum-density subgraph Y outside W. This demonstrates that W is "locally"-densest. Tatti [61] proved that all locally-densest subgraphs in a graph exhibit a hierarchical property, i.e., for any two locally-densest subgraphs X and Y, either $X \subseteq Y$ or $Y \subseteq X$. Therefore, locally-densest subgraphs can form a nested chain, which is the LDS decomposition.

DEFINITION 6. **(LDS decomposition)** [61] Given an undirected graph G = (V, E), denote all of its non-empty locally-densest subgraphs by $\{B_0, \ldots, B_k\}$, such that $B_k \subsetneq B_{k-1} \subsetneq B_{k-2} \subsetneq \cdots \subsetneq B_0$. As \emptyset is also an LDS, the LDS decomposition is defined as a set of locally-densest subgraphs, represented as $\mathcal{B} = \{B_{k+1} = \emptyset, B_k, \ldots, B_0\}$.

By the LDS decomposition, we can obtain a density value for each vertex in an LDS which is referred to as the *fractional dense* number

DEFINITION 7. **(Fractional dense number)** For a vertex $u \in B_i \setminus B_{i+1}$, the fractional dense number (FDN) of u is defined as $r_u = \rho(B_i, B_{i+1})$.

Based on these definitions, Tatti [61] further proved several useful properties of the LDS decomposition, as shown in the following lemma

LEMMA 2. For a graph G and its LDS decomposition \mathcal{B} , the following properties hold: (1) the smallest non-empty and largest locally-densest subgraphs, namely B_k and B_0 , are the densest subgraph and V, respectively; (2) B_i exhibits strictly increasing density as i increases, i.e., for two vertices $u \in B_j \setminus B_{j+1}$ and $v \in B_i \setminus B_{i+1}$ with j > i, $r_u = \rho(B_j, B_{j+1}) > \rho(B_i, B_{i+1}) = r_v$ holds.

EXAMPLE 3. Consider the graph shown in Fig. 1a. We can easily check that B_2 , B_1 , and B_0 are locally-densest subgraphs, and $B_{k+1} = B_3 = \emptyset$ is also locally-densest. The set $\{B_3, B_2, B_1, B_0\}$ is the LDS decomposition of G. The FDN r_u equals (1) 15/7 for $u \in B_2 \setminus B_3$; (2) 8/4 for $u \in B_1 \setminus B_2$; and (3) 7/4 for $u \in B_0 \setminus B_1$. These FDNs exhibit strict monotonicity 15/7 > 8/4 > 7/4. It is easy to verify that B_2 is the densest subgraph of G, which is consistent with the result shown in Lemma 2.

Density decomposition and LDS decomposition are studied separately within two different research communities [14, 26, 61]. It

is unclear whether there exists a direct connection between density decomposition and LDS decomposition. In this work, we fill this gap and establish, for the first time, a very close relationship between density decomposition and LDS decomposition, as shown in Theorem 1 and Theorem 2. Before introducing these theorems, we first present an important lemma. Due to the space limits, several proofs are replaced by proof sketches. All the complete proofs can be found in the full version of this paper from https://anonymous.4open.science/r/DD-83C6.

Lemma 3. Given an undirected graph G and an orientation \vec{G} , let k be a non-negative integer and T be the vertex set with $T \triangleq \{u \in V | d_u(\vec{G}) \geq k \text{ or } u \text{ can reach a vertex } v \text{ with } d_v(\vec{G}) \geq k \}$. If (1) for all $u \in T$, $d_u(\vec{G}) \geq k - 1$; (2) for all $u \in V \setminus T$, $d_u(\vec{G}) \leq k - 1$; (3) all edges in $E_X(T, V \setminus T)$ point towards $V \setminus T$, then T is a locally-densest subgraph.

PROOF. When T = V or $T = \emptyset$, T is obviously locally-densest. Next, we assume $T \neq V$ and $T \neq \emptyset$.

For all non-empty $X \subseteq T$ and non-empty Y disjoint from T, we

have
$$\rho(Y,T) = \frac{|E_{\times}(Y,T)| + |E(Y)|}{|Y|} \stackrel{(1)}{=} \frac{\sum_{u \in Y} d_u(\vec{G})}{|Y|} \stackrel{(3)}{\leq} k - 1, \rho(X,T \setminus X) = \frac{|E_{\times}(X,T \setminus X)| + |E(X)|}{|X|} \stackrel{(2)}{\geq} \frac{\sum_{u \in X} d_u(\vec{G})}{|X|} \stackrel{(4)}{\geq} k - 1.$$
 In a directed graph, the sum of the indegree for all vertices in

In a directed graph, the sum of the indegree for all vertices in a set equals the number of directed edges ending in that set. This leads to equality (1) and inequality (2), as all edges in $E_{\times}(Y,T)$ are directed towards Y, while there may be edges in $E_{\times}(X,T\setminus X)$ not directed toward X. Further, the inequalities (3) and (4) can be derived by the first two conditions in Lemma 3.

Next, we examine inequalities (2) and (4) to demonstrate their strictness. Concerning (4), if there exists a vertex $u \in X$ with an indegree greater than k-1, then (4) can be refined to a strict inequality. Otherwise, according to the definition of T, at least one edge in $E_X(X,T\setminus X)$ is directed towards $T\setminus X$, thereby (2) can be tightened to a strict inequality. Therefore, inequalities (2) and (4) have at least one strict inequality, and we can derive that $\rho(X,T\setminus X)>k-1\geq \rho(Y,T)$, indicating that T is a locally-densest subgraph.

Based on Lemma 3, we can obtain the following results.

THEOREM 1. Given an undirected graph G and its density decomposition $\mathcal{R} = \{R_i\}$, for all $i = 0, 1, ..., R_i$ is a locally-densest subgraph.

PROOF. In Lemma 3, the set R_k conforms to the definition of T, and, by Lemma 1, all three conditions in Lemma 3 are satisfied. Therefore, $R_k = T$ and thus it is a locally-densest subgraph. \Box

Theorem 2. Given an undirected graph G = (V, E) and a vertex $v \in V$, $\overline{r}_v = \lceil r_v \rceil$ holds.

PROOF. For an arbitrary vertex $v \in V$, suppose $v \in R_i \setminus R_{i+1}$ (i.e., $\overline{r}_v = i$) and $v \in B_j \setminus B_{j+1}$ (i.e., $r_v = \rho(B_j, B_{j+1})$). According to Theorem 1, R_i and R_{i+1} are locally-densest subgraphs, and we denote $B_{u+1} = R_{i+1}$ and $B_l = R_i$. Because of the hierarchy of locally-densest subgraphs, it follows that $B_u \subseteq B_j \subseteq B_l$, implying $\rho(B_u, B_{u+1}) \ge \rho(B_j, B_{j+1}) \ge \rho(B_l, B_{l+1})$ by Lemma 2. In the proof of Lemma 3, consider $T = R_i$ and $X = B_l \setminus B_{l+1}$. We can derive $\rho(B_l \setminus B_{l+1}, R_i \setminus (B_l \setminus B_{l+1})) = \rho(B_l, B_{l+1}) > i - 1$. Let $T = R_{i+1}$ and $Y = B_u \setminus B_{u+1}$, we have $i \ge \rho(B_u \setminus B_{u+1}, R_{i+1}) = \rho(B_u, B_{u+1})$. In conclusion, $i \ge \rho(B_j, B_{j+1}) > i - 1$ holds and thus $\overline{r}_v = i = \lceil \rho(B_j, B_{j+1}) \rceil = \lceil r_v \rceil$.

The above theorems reveal that: (1) every layer of density decomposition is a locally-densest subgraph; and (2) the IDN is the round-up value of the FDN. It is worth noting that the number of locally-densest subgraphs can be extensive and only be bounded by O(n). In contrast, the density decomposition groups locally-densest subgraphs based on the round-up of their densities, resulting in a number of non-trivial layers equaling the pseudoarboricity p, which is typically much smaller than n. This characteristic ensures that the density decomposition is not only computationally efficient but also retains a significant amount of information about the density structure of the graph.

3.2 Connection with Core Decomposition

Here, we reveal the relationship between the density decomposition and the core decomposition. The definitions of k-core and core decomposition are given as follows.

DEFINITION 8. Given an undirected graph G and an integer k, the k-core of G, denoted by C_k , is the maximal subgraph in which every vertex has a degree of at least k. The core decomposition of G is a set of subgraphs, denoted by $C = \{C_k\}$, containing all k-cores of G.

As previously mentioned, core decomposition may not always accurately reflect the density structure of a graph. However, we find that it can serve as an approximation of density decomposition. Below, we present two theorems, one of which demonstrates the 2-approximation relationship of the outer density of these two decompositions. The other one shows that they are sandwiched by each other, also following a factor of 2.

THEOREM 3. Given an undirected graph G = (V, E), for $i = 0, 1, \ldots, p, \rho(C_i, C_{i+1}) \ge \rho(R_i, R_{i+1})/2$.

PROOF. Let \vec{G} be an arbitrary egalitarian orientation of G. For brevity, denote $C_i \setminus C_{i+1}$ as C and $R_i \setminus R_{i+1}$ as R. We have $\rho(C_i, C_{i+1}) = \frac{|E_{\Delta}(C_i, C_{i+1})|}{|C|} \geq \frac{\sum_{u \in C} d_u(C_i)}{2|C|} \geq \frac{i}{2} \geq \frac{\sum_{u \in R} d_u(\vec{G})}{2|R|} = \frac{\rho(R_i, R_{i+1})}{2}$.

THEOREM 4. (Sandwich Theorem) Given an undirected graph G, for any non-negative integer i, we have $C_{2i} \subseteq R_i \subseteq C_i \subseteq R_{\lceil i/2 \rceil}$.

PROOF. To prove $R_i \subseteq C_i$, if $R_i = \emptyset$, then it evidently holds. If $R_i \neq \emptyset$, we show that in the subgraph of G induced by R_i , denoted as $G(R_i)$, every vertex has degree at least i. Let \vec{G} be an arbitrary egalitarian orientation of G. Recall that all edges between R_i and $V \setminus R_i$ are directed toward $V \setminus R_i$ in \vec{G} . For each vertex u in R_i , if $d_u(\vec{G}) \geq i$, then obviously $d_u(G(R_i)) \geq d_u(\vec{G}) \geq i$. On the other hand, if $d_u(\vec{G}) = i - 1$, since $u \in R_i$, u must have an out-edge enabling it to reach a vertex in R_i whose indegree is at least i. Thus $d_u(G(R_i)) \geq d_u(\vec{G}) + 1 \geq i$. In summary, $R_i \subseteq C_i$.

To prove $C_i\subseteq R_{\lceil i/2\rceil}$, if $C_i=\emptyset$, then it evidently holds. If $C_i\neq\emptyset$, let u be an arbitrary vertex in C_i , below we show that $u\in R_{\lceil i/2\rceil}$. Given an arbitrary egalitarian orientation \vec{G} of G, let $S\triangleq \{v|v\in C_i \text{ and } u \text{ can reach } v \text{ in } \vec{G}\}$. Note that all edges belonging to $E_\times(S,C_i\setminus S)$ are directed toward S in \vec{G} . Therefore, we have $\sum_{v\in S}d_v(\vec{G})\geq |E_\times(S,C_i\setminus S)|+|E(S)|$. By the definition of i-core, $|E_\times(S,C_i\setminus S)|+|E(S)|\geq \sum_{v\in S}d_v(C_i)/2\geq i|S|/2$. That is, the sum of indegree of vertices in S is at least i|S|/2. Then by the pigeonhole principle, at least one vertex $t\in S$ has indegree at least $\lceil i/2\rceil$ in \vec{G} . By the definition of S and density decomposition, u can reach t and thus $u\in R_{\lceil i/2\rceil}$.

Since $C_i \subseteq R_{\lceil i/2 \rceil} \Rightarrow C_{2i} \subseteq R_i$, $C_{2i} \subseteq R_i$ also holds. This completes the proof. \Box

Algorithm 1: Path (*G*)

```
Input: An undirected graph G.

Output: An egalitarian orientation \vec{G}.

1 Arbitrarily orient the edges in G to obtain \vec{G};

2 while there is a reversible path s \rightsquigarrow t in \vec{G} do

3 Reverse s \leadsto t;

4 return \vec{G};
```

Algorithm 2: GetLayer(\vec{G}, k)

```
Input: An orientation \vec{G} = (V, \vec{E}) and an integer k.
     Output: The updated \vec{G} and the layer R_k of tje density decomposition.
    V' \leftarrow V \cup \{s, t\}, where s is source and t is sink;
 2 d ← k - 1:
 3 foreach \langle u, v \rangle \in \vec{E} do
      Add arc \langle u, v \rangle to A and let c(u, v) \leftarrow 1;
 5 foreach u, d_u(\vec{G}) < d do
      Add arc \langle s, u \rangle to A and let c(s, u) \leftarrow d - d_u(\vec{G});
 7 foreach u, d_{u}(\vec{G}) > d do
      Add arc \langle u, t \rangle to A and let c(u, t) \leftarrow d_u(\vec{G}) - d;
 9 Compute the maximum flow value f_{max} of (V', A, c);
10 foreach \langle u, v \rangle \in \vec{E} do
                                                               // Copy the residual network to \vec{G}
      if \langle u, v \rangle \in A is saturated then reverse the edge \langle u, v \rangle \in \vec{E};
12 R_k \leftarrow \{u \in V | d_u(\vec{G}) \ge d \text{ or } u \text{ can reach a vertex } v \text{ with } d_v(\vec{G}) \ge d\};
13 return (\vec{G}, R_k);
```

4 ALGORITHMS FOR STATIC GRAPHS

4.1 Existing Algorithms

Existing algorithms for computing the density decomposition. Borradaile [13, 14] proposed the Path algorithm, shown in Algorithm 1, which provides a basic approach to obtain an egalitarian orientation. Since the egalitarian orientation requires that there is no reversible path $s \leadsto t$, the algorithm repeatedly searches for such paths until none can be found. Once the algorithm terminates, an egalitarian orientation can be obtained, and then R_i can be identified layer by layer based on the definition of density decomposition [13]. The time complexity of the Path algorithm is $O(m^2)$ as shown in [13]. Clearly, such a high time complexity of the Path algorithm poses a significant challenge when handling large graphs.

Existing algorithms for LDS decomposition. As shown in Theorem 2, each IDN is a round-up value of FDN. Thus, we can obtain the density decomposition by computing the LDS decomposition first and then rounding up the FDN values. Tatti [61] proposed a network flow algorithm to compute LDS decomposition with a time complexity of $O(n^2m)$. Then, Danisch et al. [26] proposed the current state-of-the-art FW algorithm. The algorithm initially employs convex optimization to obtain an approximate LDS decomposition and then invokes Tatti's algorithm to calculate exact LDS decompositions. Subsequently, Harb et al. [35] introduced a new approximate algorithm based on the proximal gradient method, which can be integrated into the FW algorithm, resulting in the Fista algorithm. Both of these two highly-optimized algorithms can significantly improve the practical performance, but the worstcase time complexity remains $O(n^2m)$. As a consequence, these LDS-based algorithms may still perform poorly on large graphs due to their high worst-case time complexities, as confirmed in our experiments.

4.2 A Re-orientation Network Flow Algorithm

As discussed above, existing algorithms for density decomposition suffer from the high complexity issue. To overcome this problem, we develop a novel algorithm based on a so-called re-orientation network flow technique. The re-orientation network flow technique was originally devised to compute the pseudoarboricity of a graph [7]. In this paper, we discover that such a powerful network flow technique can also be used to compute the density decomposition. Below, we introduce the concept of the re-orientation network.

DEFINITION 9. (Re-orientation network) [7] Given an orientation $\vec{G} = (V, \vec{E})$ and an integer d, the re-orientation network is a weighted network with an additional source vertex s and sink vertex t where the weight of each edge denotes the capacity of the edge. Specifically, the re-orientation network with parameter d is defined as $(V \cup \{s,t\},A,c)$ where $(1) \langle u,v \rangle \in A, c(u,v) = 1$, if $\langle u,v \rangle \in E$; $(2) \langle s,u \rangle \in A, c(s,u) = d - d_u(\vec{G})$, if $d_u(\vec{G}) < d$; and $(3) \langle u,t \rangle \in A, c(u,t) = d_u(\vec{G}) - d$, if $d_u(\vec{G}) > d$.

By Definition 9, the re-orientation network uses a parameter d to separate vertices based on their indegree. The source s is connected to vertices with an indegree less than d, while the sink t is linked to vertices with indegree greater than d. Based on this, we can perform a maximum flow computation on this re-orientation network. When the maximum flow algorithm terminates, there does not exist a path from s to t (i.e., no augmentation path exists in the residual network). That is to say, in this case, the vertices with indegree less than d cannot reach the vertices with indegree larger than d. Let T be the set of vertices that can reach t. When the maximum flow algorithm terminates, only the vertices in T have degrees larger than d. As a consequence, the set T satisfies the three conditions in Lemma 3, indicating that T is a locally-densest subgraph. Below, we further prove that the set T defined in Lemma 3 is exactly R_{d+1} .

LEMMA 4. The subgraph T defined in Lemma 3 is R_k .

PROOF. By the proof of Lemma 3, when $T \neq \emptyset$ and $T \neq V$, for all $X \subseteq T$ and Y disjoint from T, we have $\rho(X, T \setminus X) > k-1 \ge \rho(Y, T)$. Suppose that T is the locally-densest subgraph B_i . Then, according to the above inequalities, $\rho(B_i \setminus B_{i+1}, B_{i+1}) > k-1 \ge \rho(B_{i-1} \setminus B_i, B_i)$. Thus, the FDN of vertices in T is strictly greater than k-1, while the FDN of vertices in $V \setminus T$ is less than or equal to k-1. By Lemma 2, the round-up FDN is equal to IDN. As a result, all vertices in T have IDN no less than t, and all vertices in $t \setminus T$ have IDN less than t, which proves $t \in T$ and $t \in T$ are $t \in T$. Only the inequality $t \in T$ is a property of $t \in T$. It has a property of $t \in T$ in the inequality $t \in T$ is the lemma can be easily established.

By Lemma 4, we devise the re-orientation network flow algorithm, namely GetLayer, which is outlined in Algorithm 2. Below, we show that each invocation of GetLayer(\vec{G}, k) can correctly output one layer of density decomposition, i.e., R_k .

Theorem 5. Algorithm 2 correctly outputs R_k .

PROOF. First, an important observation is that in the output orientation \vec{G} , there is no path $s \rightsquigarrow t$ such that $d_s(\vec{G}) < k-1$ and $d_t(\vec{G}) > k-1$. If such a path exists, then the flow value can be increased by filling this path. As the flow reaches its maximum value, such paths no longer exist.

To avoid confusion, we use T to denote the output R_k of the GetLayer algorithm. Next, we prove that T equals the layer R_k in the density decomposition. T contains all vertices with indegree greater than k-1 or reachable to a vertex with indegree greater

Algorithm 3: Flow(G)

```
Input: An undirected graph G. Output: Density decomposition \mathcal{R} = \{R_i\}.

1 Invoke a 2-approximation algorithm to calculate an approximate orientation \vec{G};

2 foreach k = 0, 1, 2, \dots do

3 \begin{pmatrix} (\vec{G}, R_k) \leftarrow \text{GetLayer}(\vec{G}, k); \\ \text{if } R_k = \emptyset \text{ then break}; \end{pmatrix}

5 \vec{G} \leftarrow \text{the induced subgraph of } R_k \text{ in } \vec{G}; // pruning strategy

6 return \mathcal{R} = \{R_i\};
```

than k-1. Since vertices with indegree less than k-1 cannot reach vertices with indegree greater than k-1, they are not in T. As a result, the vertices in T all have indegree greater than or equal to k-1, the vertices in $V\setminus T$ have indegree less than or equal to k-1, and the edges in $E_{\times}(T,V\setminus T)$ all point to $V\setminus T$. By Lemma 4, $T=R_k$.

The time complexity of Algorithm 2 depends on computing the maximum flow in the re-orientation network. It was shown that the re-orientation network maximum flow algorithm can be implemented with time complexity of $O(m^{3/2})$ and space complexity of O(m) [10], and thus Algorithm 2 has the same complexity.

4.3 The Proposed Flow Algorithm

As shown in Theorem 5, each invocation of the GetLayer algorithm outputs one layer of density decomposition. Consequently, simply invoking GetLayer p+2 times can yield all layers from $R_0=V$ to $R_{p+1}=\emptyset$. Based on this, we propose the Flow algorithm for density decomposition, as shown in Algorithm 3.

Algorithm 3 first invokes an existing linear-time 2-approximation algorithm to obtain an orientation [28] (Line 1). Note that with such a 2-approximation initial orientation, the maximum flow computation can be faster than that with arbitrary initial orientation, because this approach is expected to reduce the number of augmentation paths in the maximum flow computation as indicated in [10]. Then, the Flow algorithm iteratively performs GetLayer from the bottom layer R_0 to yield each layer one by one (Lines 2-5). Once R_k is computed, the algorithm can directly assign \vec{G} as the induced subgraph of R_k in \vec{G} by discarding the vertices in $V \setminus R_k$ (Line 5). Since the indegree of vertices in $V \setminus R_k$ is no larger than k-1, they cannot reach vertices in R_k . In the following 'foreach' loop, the vertices in $V \setminus R_k$ are all connected to the sink t, but the flow cannot enter $V \setminus R_k$ from R_k . Thus, Flow only needs to compute R_{k+1} within R_k in the subsequent computation, which can significantly improve the efficiency. When k equals p + 1, R_k becomes an empty set, indicating that all layers of the density decomposition are obtained, thus the Flow algorithm terminates. The correctness of Flow can be guaranteed by Theorem 5. Below, we analyze the complexity of Flow(G).

THEOREM 6. The worst-case time and space complexity of Algorithm 3 is $O(p \cdot m^{3/2})$ and O(m+n) respectively.

PROOF. Algorithm 3 invokes the GetLayer algorithm p+2 times, and each invocation consumes $O(m^{3/2})$ time. Therefore, the total time complexity of $\mathsf{Flow}(G)$ is $O(p \cdot m^{3/2})$. For the space complexity, it is easy to see that the algorithm only consumes O(m) space. \square

Note that since $p \le \sqrt{m}$ [20], the worst-case time complexity of Algorithm 3 is lower than that of Algorithm 1. Furthermore, p is often small in real-world graphs [10], thus the practical performance of Algorithm 3 is typically much better than the worst-case time

Algorithm 4: Flow++ (*G*)

```
Input: An undirected graph G.
     Output: The egalitarian orientation \vec{G} and density decomposition \{R_i\}
    Invoke a 2-approximation algorithm to obtain an approximate orientation \vec{G} and a
        2-approximation of pseudoarboricity \bar{p};
 2 \quad R_{\bar{p}+1} \leftarrow \emptyset; R_0 \leftarrow V;
 3 \hat{\text{Divide}}(R_{\bar{p}+1}, R_0);
 4 return (\vec{G}, \{R_i\});
    Function Divide(R_u, R_l)
| if u - l \le 1 or R_u = R_l then return;
             k_u \leftarrow u, k_l \leftarrow l;
            while k_u > k_l do
 | k \leftarrow \lfloor (k_u + k_l + 1)/2 \rfloor;
                    (\vec{G}, R_k) \leftarrow \text{GetLayer}(\vec{G}, k);
10
                    if |E_{\Delta}(R_l, R_k)| < |E_{\Delta}(R_l, R_u)|/2 then k_l \leftarrow k;
11
12
                    else k_u \leftarrow k - 1;
13
             \mathsf{Divide}(R_k,R_l);
            Divide(R_u, R_{k+1});
```

complexity suggests. Indeed, as shown in our experiments, Algorithm 3 is around two orders of magnitude faster than Algorithm 1 on most datasets.

4.4 The Proposed Flow++ Algorithm

In this subsection, we proposed an improved algorithm, called Flow++. By using a newly-developed divide-and-conquer technique, Flow++ improves the time complexity of Flow from $O(p \cdot m^{3/2})$ to $O(\log p \cdot m^{3/2})$. The key idea of Flow++ is that it partitions the graph into almost equal parts at each step and performs maximum flow computation recursively in each part. Recall that in Flow, we compute R_i within R_{i-1} instead of the whole graph G due to the hierarchical structure of density decomposition. Similar to this strategy, an improved strategy is to compute R_i between R_u and R_l , where u > i > l, which can intuitively further reduce the data scale. To construct such R_{μ} and R_{l} , a divide-and-conquer manner can be employed to recursively compute all layers of density decomposition. Specifically, the recursive function, denoted as $Divide(R_u, R_l)$, is designed to find all R_i for u > i > l. It first selects a parameter kwhere u > k > l and divides the graph into two parts: $R_l \setminus R_k$ and $R_{k+1} \setminus R_u$. Then, Divide (R_k, R_l) and Divide (R_u, R_{k+1}) are separately invoked for deeper recursions.

Equipped with such a divide-and-conquer approach, the improved algorithm Flow++ is proposed to compute the density decomposition, which is outlined in Algorithm 4. Specifically, Algorithm 4 starts by invoking an approximate algorithm to calculate a 2-approximate pseudoarboricity $\bar{p} \geq p$ (Line 1). Then, we can obtain two trivial layers: $R_{\bar{p}+1}=\emptyset$ and $R_0=V$ (Line 2). The recursion begins with these two initial layers by calling $Divide(R_{\bar{p}+1}, R_0)$ (Line 3). During this invocation, a parameter k is selected to divide the graph (Lines 7-13). We use a binary search (Lines 7-13) to find the maximum k satisfying $|E_{\Delta}(R_l, R_k)| < |E_{\Delta}(R_l, R_u)|/2$. Thus, for k+1, we have $|E_{\Delta}(R_l,R_{k+1})| \ge |E_{\Delta}(R_l,R_u)|/2$, suggesting that $|E_{\Delta}(R_{k+1}, R_u)| = |E_{\Delta}(R_l, R_u)| - |E_{\Delta}(R_l, R_{k+1})| < |E_{\Delta}(R_l, R_u)|/2.$ Therefore, both the edges $E_{\Delta}(R_l, R_k)$ and the edges $E_{\Delta}(R_{k+1}, R_u)$ are less than half of the edges $E_{\Delta}(R_l, R_u)$. This ensures the number of edges being divided by 2 in each depth of recursion. When $u-l \le 1$ or $R_u = R_l$, there are no further subdivided layers between R_u and R_l , thus Divide can return directly. The correctness of the Flow++ algorithm can be easily derived by induction because in each recursion we can identify a R_k and no layer will be missed by the divide-and-conquer procedure. Thanks to our divide-andconquer technique, the graph size is halved with each recursion.

As a result, the computation costs exponentially decrease for the higher-level layers. This is why our Flow++ algorithm can achieve a lower time complexity compared to the Flow algorithm.

The remaining challenge in the above divide-and-conquer algorithm lies in implementing the computation within $R_l \setminus R_u$. We find that we can slightly modify the re-orientation network flow algorithm to achieve this goal. Specifically, we need to consider not only the subgraph induced by $R_l \setminus R_u$ in \vec{G} but also the cross edges $E_{\times}(R_l, R_u)$. In GetLayer, arcs from the source and sink to vertices in only $R_l \setminus R_u$ are connected (the other vertices are pruned). However, for computing the indegree of the vertices in $R_l \setminus R_u$, the edges in $E_{\times}(R_l, R_u)$ are also considered. The reasons why these modifications of GetLayer can ensure the correctness are as follows. Because R_I and R_{II} are either previously computed by the GetLayer algorithm or are the trivial subgraphs \emptyset or V. According to the properties of GetLayer, the edges in $E_{\times}(V \setminus R_u, R_u)$ all point towards $V \setminus R_u$, and the edges in $E_{\times}(V \setminus R_l, R_l)$ all point towards $V \setminus R_l$. This indicates that at this point when the GetLayer algorithm is called again for further computation, there will be no flow from $V \setminus R_u$ to R_u , and no flow from $V \setminus R_l$ to R_l . As a result, we can safely ignore $V \setminus R_l$ and R_u , which do not have any flow passing through them, and only connect the vertices in $R_l \setminus R_u$ to the source and sink.

Clearly, with this divide-and-conquer implementation, the size of the network can be bounded by $E_{\Delta}(R_l,R_u)$ in each recursion. Thus, by [10], the time complexity of GetLayer on subgraph $R_l \setminus R_u$ is $O(|E_{\Delta}(R_l,R_u)|^{3/2})$. Based on this, we can derive the total time and space complexity of Flow++ as follows.

THEOREM 7. The time complexity of Algorithm 4 is $O(m^{3/2} \cdot \log p)$, and the space complexity is $O(p \cdot n)$.

PROOF. The time complexity of Divide (R_l,R_u) consists of two parts: the complexity of the binary search procedure in each recursion and the complexity of two deeper recursions Divide (R_k,R_l) and Divide (R_u,R_{k+1}) . Since the binary search range is O(p), it only requires $O(\log p)$ rounds of binary search. Therefore, the time complexity of the binary search in Divide (R_u,R_l) is $O(|E_{\Delta}(R_l,R_u)|^{3/2} \cdot \log p)$. For deeper recursions, the binary search in Divide ensures that k is the largest integer satisfying $|E_{\Delta}(R_l,R_k)| < |E_{\Delta}(R_l,R_u)|/2$, which guarantees that both $|E_{\Delta}(R_l,R_k)|$ and $|E_{\Delta}(R_{k+1},R_u)|$ are less than $|E_{\Delta}(R_l,R_u)|/2$. Thus, the data scale is divided by 2 in each recursive layer. According to the master theorem for divide-and-conquer algorithms [5], the time complexity of deeper recursions in Divide (R_u,R_l) is no higher than the time complexity of the binary search. Hence, it is easy to show that the total time complexity of Flow++ is $O(m^{3/2} \cdot \log p)$.

For the space complexity, GetLayer takes O(m) space, and each recursion depth needs to store information for R_u and R_l , occupying O(n) space. Since the number of layers does not exceed O(p), the space complexity for storing R_u and R_l is $O(p \cdot n)$. Because $p \ge m/n \Rightarrow O(m) \subseteq O(p \cdot n)$ [48], the space complexity of Flow++ is $O(p \cdot n)$.

Note that since p is small in real-world graphs, $\log_2 p$ is a very small constant. For example, for most real-world graphs where $p \leq 1024$, we have $\log_2 p \leq 10$. Consequently, the complexity of our algorithm is near to $O(m^{3/2})$, making it highly efficient for processing real-world graphs. Indeed, as shown in our experiments, Flow++ is extremely efficient, and it can further achieve a significant speedup compared to the Flow algorithm.

5 DYNAMIC MAINTENANCE ALGORITHMS

When the graph is updated by an edge insertion or deletion, a straightforward algorithm to maintain the density decomposition is to invoke the Flow++ algorithm to re-compute the density decomposition from scratch. Clearly, such an approach is costly. To improve the efficiency, in this section, we develop several novel algorithms to maintain the density decomposition when the graph is updated by an edge insertion or deletion. To avoid confusion, we assume that the notations, including IDN \bar{r} , indegree d, and R_i , all denote their values before edge updated unless otherwise specified.

5.1 Density Decomposition Update Theorem

In this subsection, we establish a density decomposition update theorem, based on which we can devise the maintenance algorithms.

Theorem 8. (Density decomposition update theorem) When inserting (resp. deleting) an edge (u,v) in the undirected graph G, assuming $\overline{r}_v \leq \overline{r}_u$, only the IDNs of vertices whose IDN equals \overline{r}_v , i.e., vertices in $R_{\overline{r}_v} \setminus R_{\overline{r}_v+1}$, may change, and it can only increase by 1 (resp. decrease by 1).

PROOF. We first analyze the case of edge insertion. Prior to the insertion operation, based on Theorem 1 and Lemma 3, we can conclude that the inequality $\rho(X,R_k\setminus X)>k-1$ holds for any R_k and non-empty $X\subseteq R_k$. The subsequent insertion of (u,v) does not decrease $\rho(X,R_k\setminus X)$, thus $\rho(X,R_k\setminus X)>k-1$ is satisfied. As a result, the FDNs of vertices in R_k remain larger than k-1. By Theorem 2, the IDNs are no less than k, indicating that inserting (u,v) does not decrease the IDNs.

Next, we show that only the IDNs of vertices within $R_{\overline{r}_n} \setminus R_{\overline{r}_n+1}$ may increase by at most 1. Let \vec{G} be an egalitarian orientation of G before the insertion of (u, v). For any non-negative integer $k \neq 0$ $\overline{r}_v + 1$, suppose that we directly insert $\langle u, v \rangle$ into \vec{G} . The condition $\overline{r}_v \leq \overline{r}_u$ ensures that the edges in $E_{\times}(R_k, V \setminus R_k)$ still point to $V \setminus R_k$. Moreover, the insertion of $\langle u, v \rangle$ only alters the indegree of v, increasing by 1. Therefore, the indegree of vertices in R_k continues to be no less than k-1, and the indegree of vertices in $V \setminus R_k$ remains no larger than k-1. The analysis confirms that R_k obeys the three conditions in Lemma 3 and Lemma 4, and thus we can conclude that R_k remains unchanged after inserting (u, v). However, when $k = \overline{r}_v + 1$, the indegree of the vertex $x \in V \setminus R_k$ may reach k. Since the IDN of each vertex does not decrease, it can be derived that after inserting (u, v), only vertices in $R_{\overline{r}_n} \setminus R_{\overline{r}_n+1}$ might be merged into $R_{\overline{r}_v+1} \setminus R_{\overline{r}_v+2}$, leading to an increment of 1 in the IDN.

For edge deletion, using a similar argument, we can easily derive that: (1) the removal of (u,v) does not increase the IDN of vertices; (2) for any non-negative integer $k \neq \overline{r}_v$, R_k remains unchanged; (3) only the vertices in $R_{\overline{r}_v} \setminus R_{\overline{r}_v+1}$ might be merged into $R_{\overline{r}_v-1} \setminus R_{\overline{r}_v}$, resulting in a decrement of 1 in the IDN. This completes the proof.

Theorem 8 indicates that maintaining density decomposition is highly localized. That is, we only need to consider the vertices with IDN equal to \bar{r}_v . Moreover, given that the density decomposition is defined based on an egalitarian orientation, the key to updating the density decomposition lies in maintaining the egalitarian orientation, building upon which we develop several efficient density decomposition maintenance algorithms in the following.

Algorithm 5: Insert $(\vec{G}, \overline{r}, (u, v))$

```
Input: Egalitarian orientation \vec{G}, the IDN of all vertices \overline{r}, and the edge (u,v) to be inserted.

Output: The updated egalitarian orientation \vec{G} and IDN \overline{r}.

1 Suppose \overline{r}_v \leq \overline{r}_u, otherwise swap the input edge (u,v);

2 if d_v = \overline{r}_v - 1 then \vec{G} \leftarrow \vec{G} \cup \langle u,v \rangle;

3 else //d_v = \overline{r}_v

4 | \vec{G} \leftarrow \vec{G} \cup \langle u,v \rangle;

5 if there is a reversible path s \leadsto v, where d_s = \overline{r}_v - 1 then

6 | Reverse the path;

7 else | \vec{r}_v \leftarrow \vec{r}_w + 1 \rangle;

9 | foreach w \in \{w | \overline{r}_w = \overline{r}_v \text{ and } w \text{ can reach } v\} do \overline{r}_w \leftarrow \overline{r}_w + 1;

10 return (\vec{G}, \overline{r}):
```

5.2 The Proposed Insertion Algorithm

Based on Theorem 8, we propose an insertion algorithm, called Insert, to handle the case of edge insertion. The detailed implementation of our algorithm is shown in Algorithm 5. The main idea of Insert is as follows. By Theorem 8, Insert needs to consider three different cases: (1) $d_v = \overline{r}_v - 1$ (Line 2); (2) $d_v = \overline{r}_v$ and there exists $s \leadsto v$ (Lines 3-6); (3) $d_v = \overline{r}_v$ and there is no $s \leadsto v$ (Lines 3-4 and Lines 7-9). In case (1), the Insert algorithm directly inserts $\langle u, v \rangle$. Following this operation, \vec{G} is still an egalitarian orientation, and the IDNs do not need to be updated. In case (2), Insert also inserts $\langle u, v \rangle$, but a reversible path $s \rightsquigarrow v$ is created, making \vec{G} non-egalitarian. Consequently, Insert promptly reverses this path, restoring \vec{G} to an egalitarian state. At this point, the IDNs also do not change. Note that for case (2), we can perform Breadth-First Search (BFS) in the reversed graph from the vertex v to find the reversible path $s \rightsquigarrow v$ (Lines 5-6). In case (3), where no reversible path is found, \vec{G} remains an egalitarian orientation, but the IDNs of some vertices need to be increased by 1 according to Theorem 8. Similarly, in this case, we can also invoke a reverse BFS algorithm from v to identify the set $\{w|\overline{r}_w=\overline{r}_v \text{ and } w \text{ can reach } v\}$ (Line 8). Below, we prove that Insert correctly maintains the density decomposition (i.e., maintaining all IDNs for all vertices).

Theorem 9. Algorithm 5 correctly maintains an egalitarian orientation and all IDNs for all vertices.

PROOF. We prove the theorem by considering three cases, namely: (1) $d_v = \overline{r}_v - 1$; (2) $d_v = \overline{r}_v$ and there exists $s \leadsto v$; (3) $d_v = \overline{r}_v$ and there is no $s \leadsto v$.

We first prove that Insert correctly updates the egalitarian orientation, i.e., proving that the output orientation has no reversible path. For case (1), since $d_v+1=\overline{r}_v\leq\overline{r}_u$, even if we directly insert $\langle u,v\rangle$ into \vec{G} , \vec{G} still satisfies the properties in Lemma 1, and it is clear that \vec{G} has no reversible path. In the case of (2), because for any k, the vertices in $V\setminus R_k$ cannot reach the vertices in R_k , combining that all vertices on the path $s\leadsto v$ can reach v, all these vertices also must all be in $R_{\overline{r}_v}\setminus R_{\overline{r}_v+1}$. Reversing the path $s\leadsto v$ reduces d_v by 1, and the situation becomes the same as case (1). In case (3), there is no reversible path ending at v, and because of $\overline{r}_v\leq\overline{r}_u$ and Lemma 1, there is also no reversible path ending at other vertices.

Next, we prove that Insert correctly maintains the IDNs of vertices. As the output \vec{G} is an egalitarian orientation, we can directly analyze the indegree and reachability of vertices. According to Theorem 8, only vertices in $R_{\overline{r}_v} \setminus R_{\overline{r}_v+1}$ need to be considered. Based on the proof of egalitarian orientation, it can be deduced that in the updated \vec{G} , vertices in $R_{\overline{r}_v} \setminus R_{\overline{r}_v+1}$ cannot reach the vertices in $R_{\overline{r}_v} \setminus R_{\overline{r}_v+1}$ for cases (1) and (2). Thus, the indegree of vertices in $R_{\overline{r}_v} \setminus R_{\overline{r}_v+1}$

Algorithm 6: Delete $(\vec{G}, \overline{r}, (u, v))$

```
Input: Egalitarian orientation \vec{G}, the IDN of all vertices \vec{r}, and the edge (u,v) need to be deleted.

Output: The updated egalitarian orientation \vec{G} and IDN \vec{r}.

1 Suppose (u,v) is oriented as (u,v), otherwise swap the input edge (u,v);

if d_v = \vec{r}_v = 1 then

There must be a reversible path v \leadsto t, where d_t = \vec{r}_v;

4 Reverse the path;

5 \vec{G} \leftarrow \vec{G} - \langle u,v \rangle;

6 Let \vec{r}_0 \leftarrow \vec{r}_v;

7 Let S \leftarrow \{w|\vec{r}_w = \vec{r}_0, and d_w = \vec{r}_0 or w can each an \vec{r}_0-indegree vertex};

8 forall w where \vec{r}_w = \vec{r}_0 and w \notin S do

9 \vec{r}_w \leftarrow \vec{r}_0 = 1;

return (\vec{G}, \vec{r});
```

does not increase to \overline{r}_v+1 , meaning that the IDN remains \overline{r}_v . In case (3), only the indegree of vertex v in $R_{\overline{r}_v}\setminus R_{\overline{r}_v+1}$ increases to \overline{r}_v+1 . Therefore, the vertices in $R_{\overline{r}_v}\setminus R_{\overline{r}_v+1}$ that can reach vertex v, as well as vertex v itself, have their IDNs increased to \overline{r}_v+1 . \square

Below, we analyze the time complexity of Algorithm 5 as follows.

Theorem 10. The worst-case time complexity of Algorithm 5 is $O(\sum_{u \in R_{\overline{r}_n} \setminus R_{\overline{r}_n+1}} d_u(G))$.

PROOF. Finding the path $s \leadsto v$ and obtaining the set $\{w | \overline{r}_w = \overline{r}_v \text{ can reach } v\}$ both can be accomplished using the reverse Breadth-First Search (BFS) algorithm once, and it only needs to search for vertices in $R_{\overline{r}_v} \setminus R_{\overline{r}_v+1}$. Therefore, the time complexity of Insert is $O(\sum_{u \in R_{\overline{r}_v} \setminus R_{\overline{r}_v+1}} d_u(G)) \subseteq O(m)$.

As shown in Theorem 10, the time complexity of Algorithm 5 is clearly bounded by O(m). Furthermore, its complexity only depends on the size of the subgraph where the vertices' IDNs might be updated, indicating that the algorithm is local and only needs to explore a small portion of the graph. Indeed, as shown in our experiments, Algorithm 5 can be 5-7 orders of magnitude faster than the baseline algorithm that relies on re-computation.

5.3 The Proposed Deletion Algorithm

In this subsection, we proposed the Delete algorithm to maintain the density decomposition for handling edge deletion. The detailed implementation of Delete is outlined in Algorithm 6. Suppose that the deleted edge (u,v) is oriented as $\langle u,v\rangle$ (Line 1). According to Lemma 1, the inequality $\bar{r}_v \leq \bar{r}_u$ holds. The Delete algorithm addresses two cases for maintaining the egalitarian orientation: (1) $d_v = \bar{r}_v$ (not satisfying the 'if' condition of Line 2); (2) $d_v = \bar{r}_v - 1$ (satisfying the 'if' condition). In case (1), Delete directly deletes $\langle u,v\rangle$ (Line 5), and \bar{G} is still an egalitarian orientation. In case (2), by the definition of density decomposition, a reversible path $v \leadsto t$ must exist (Line 3). This ensures that Delete does not encounter the scenario of being unable to find a reversible path, as opposed to Insert. After reversing $v \leadsto t$, case (2) transforms into case (1). Then, $\langle u,v\rangle$ can be deleted and \bar{G} is restored to the egalitarian orientation.

With the egalitarian orientation, the Delete algorithm starts maintaining the IDNs of vertices. Unfortunately, unlike Insert, which can update IDNs directly, there is no direct method to update IDNs in the deletion case. Therefore, Delete needs to perform Lines 6-9 to update IDNs. Since \bar{r}_v may change, to avoid ambiguity, we use \bar{r}_0 to denote the value of \bar{r}_v before updating (Line 6). By the definition of density decomposition, Delete employs a single BFS algorithm to determine the set of vertices S that do not require IDN updates (Line 7). It then decreases the IDNs of vertices that were originally equal to \bar{r}_0 but are not in the set S (Lines 8-9). Finally,

the Delete algorithm terminates, outputting the egalitarian orientation and the IDNs of all vertices. The following theorem shows the correctness of Algorithm 6.

Theorem 11. Algorithm 6 correctly updates the egalitarian orientation and the IDNs of all vertices.

PROOF. To prove the theorem, we need to consider two cases: (1) $d_v = \overline{r}_v$; (2) $d_v = \overline{r}_v - 1$. We first prove that Delete correctly updates the egalitarian orientation. In case (1), Delete directly deletes $\langle u, v \rangle$. For vertex v, before the edge deletion, it is impossible for v to reach any vertex with an indegree no less than \bar{r}_v + 1. Therefore, after deleting the edge, a reversible path starting from v cannot exist. For other vertices, the deletion operation does not expand their reachability to other vertices, thus no reversible path from them appears. For case (2), we can easily derive that all vertices in the path $v \rightsquigarrow t$ are also in $R_{\overline{r}_n} \setminus R_{\overline{r}_n+1}$. Therefore, after reversing $v \rightsquigarrow t$, \vec{G} remains an egalitarian orientation, and case (2) becomes (1). Thus, the proof is established. Then, we show the correctness of Delete for updating IDNs which is straightforward. By Theorem 8, only the vertices having IDNs of \overline{r}_0 need to be updated, while the set Scontains all vertices whose IDNs remain unchanged. Therefore, the IDNs of vertices can be correctly updated by Delete.

Theorem 12. The worst-case time complexity of Algorithm 6 is $O(\sum_{u \in R_{\overline{r}_n} \backslash R_{\overline{r}_{n+1}}} d_u(G))$.

PROOF. Similar to Insert, all computation in Delete can be carried out within $R_{\overline{r}_v} \setminus R_{\overline{r}_v+1}$. Furthermore, the search for S can be done by performing the BFS algorithm starting from all vertices with indegree equal to \overline{r}_0 in $R_{\overline{r}_v} \setminus R_{\overline{r}_v+1}$. Therefore, the time complexity of the Delete algorithm is the same as Insert, which is $O(\sum_{u \in R_{\overline{r}_v} \setminus R_{\overline{r}_v+1}} d_u(G)) \subseteq O(m)$.

Similar to the Insert algorithm, the Delete algorithm is also local, and the worst-case time complexity is clearly bounded by O(m). Our experiments also show that Delete is very efficient and can be 3-5 orders of magnitude faster than the baseline which is based on re-computation.

5.4 An Improved Deletion Algorithm

Despite exhibiting the same time complexity, in our experiments, the Delete algorithm runs around three orders of magnitude slower than Insert. The main issue is, to update the IDNs of vertices, Insert performs BFS only within $\{w|\bar{r}_w=\bar{r}_v \text{ and } w \text{ can reach } v\}\subseteq R_{\bar{r}_v}\setminus R_{\bar{r}_v+1}$, whereas Delete necessitates considering the entire $R_{\bar{r}_v}\setminus R_{\bar{r}_v+1}$. In this subsection, we devise an improved algorithm, called Delete++ (Algorithm 7), to further enhance the performance of the deletion algorithm.

The rationale and intuition behind Delete++ are as follows. If the IDN of a vertex w decreases due to the deletion of $\langle u,v\rangle$, by Definition 3, w must have been capable of reaching v prior to the deletion. Otherwise, w would have possessed the ability to reach another vertex with an \overline{r}_v -indegree distinct from v before the deletion, thereby preventing its IDN from decreasing. Therefore, we can calculate the set $P\subseteq R_{\overline{r}_v}\setminus R_{\overline{r}_v+1}$, which includes the vertices capable of reaching v (Line 2). The vertices where the IDNs may be reduced are specifically within the set P, allowing us to narrow down the scope for updating the IDN to P rather than $R_{\overline{r}_v}\setminus R_{\overline{r}_v+1}$.

Subsequently, we turn our focus to updating the IDNs of the vertices in P. Recall that the Delete algorithm identifies all vertices capable of reaching the vertices with \bar{r}_v -indegree, and then reduces the IDNs of vertices other than these vertices. Such an approach,

Algorithm 7: Delete++ $(\vec{G}, \bar{r}, (u, v))$

```
Input: Egalitarian orientation \vec{G}, the IDN of all vertices \overline{r}, and the edge (u,v) to be deleted.

Output: The updated egalitarian orientation \vec{G} and IDN \overline{r}.

1 Perform Lines 1-4 of Delete;

2 Let P \leftarrow \{w | w \text{ can reach } v\} \cap (R_{\overline{r}_U} \setminus R_{\overline{r}_U+1});

3 \vec{G} \leftarrow \vec{G} - \langle u, v \rangle;

4 foreach s \in P and d_s = \overline{r}_v - 1 do

5 | if the SCC containing s has not been computed then

6 | Let S \leftarrow \{s\} \cup \{w | s \text{ can reach } w\} \cap (R_{\overline{r}_U} \setminus R_{\overline{r}_U+1});

7 | Compute all SCCs in the set S, while recording whether they can reach a \overline{r}_O-indegree vertex;

8 | if the SCC containing s cannot reach a vertex with \overline{r}_U indegree then

9 | \overline{r}_S \leftarrow \overline{r}_S - 1;
```

however, cannot be directly applied to update the IDN within P. This is because, if a vertex in P cannot reach \overline{r}_v -indegree vertices within P, it does not necessarily imply that its IDN needs to be decreased since it might still have the possibility of reaching \overline{r}_v -indegree vertices outside of P.

A basic idea is performing a BFS individually on each vertex in P to determine its reachability to vertices with an indegree of \bar{r}_v . But it is impractical due to massive redundant re-computations. The key to solving this problem lies in reducing the number of BFS operations instead of "individually" searching for each vertex. To achieve this, we leverage a classic concept of Strongly Connected Component [60] (SCC). Once whether a vertex s can reach \bar{r}_v -indegree vertices is confirmed, vertices within the same SCC as s also share the same reachability, eliminating the need to invoke the BFS algorithm on these vertices again.

In the following, we show the correctness of the Delete++ algorithm. Since Line 1 and Line 3 of Delete++ is the same as the method for maintaining egalitarian orientation in Delete, Delete++ can correctly update egalitarian orientation. Next, we demonstrate that Delete++ can correctly update the IDNs of vertices.

Theorem 13. Algorithm 7 correctly updates the IDNs of vertices.

PROOF. First, an important observation is that in an egalitarian orientation \vec{G} , vertices within the same SCC share identical IDNs. According to the definition of density decomposition, the IDN of a vertex depends on its reachability in the egalitarian orientation. Given that vertices within the same SCC share an identical set of reachable vertices, their IDNs are identical.

In accordance with Theorem 8, the necessity to update IDNs is confined to vertices in $R_{\overline{r}_v} \setminus R_{\overline{r}_v+1}$. In Line 2 of Algorithm 7, the vertices with (\overline{r}_v-1) -indegree in the set $(R_{\overline{r}_v} \setminus R_{\overline{r}_v+1}) \setminus P$ are guaranteed to reach another vertex having an \overline{r}_v -indegree distinct from v. It further follows that the deletion of $\langle u,v \rangle$ will not affect their IDNs. Thus, the key consideration lies in determining whether there is a necessity to reduce the IDNs of (\overline{r}_v-1) -indegree vertices in set P. In the 'foreach' loop, consider each (\overline{r}_v-1) -indegree vertex s in P one by one. By Definition 3, the IDN of vertex s ought to decrease if and only if there are no vertices with an \overline{r}_v -indegree in S. This condition is equivalent to the SCC containing s being unable to reach a \overline{r}_v -indegree vertex. Therefore, the IDN of vertex s can be updated correctly.

We employ the linear-time Tarjan algorithm [60] to compute SCCs, and thus the complexity of Delete++ is equivalent to that of Delete. However, in contrast to Delete, which accounts for the entire set $R_{\overline{r}_n} \setminus R_{\overline{r}_n+1}$, Delete++ focuses solely on the neighborhood

P of vertex *v*. Consequently, the actual running time of Delete++ proves notably faster than that of Delete in our experiments.

5.5 Discussions

As shown in Theorems 10 and 11, both our insertion and deletion algorithms only need to traverse a small portion of the graph and their worst-case time complexity can clearly be bounded by O(m), revealing the advantage of relatively-easy maintenance of density decomposition. In contrast, the LDS decomposition appears to be challenging to maintain exactly. To our knowledge, there does not exist an efficient algorithm to exactly maintain the LDS decomposition, except for re-computation from scratch. The potential reasons are summarized as follows.

First, for the LDS decomposition, we find that the insertion or deletion of an edge may generate many new layers, resulting in high updating costs. For example, consider a graph G consists of three subgraphs, A, B, and C, where A is connected to both B and C, but B and C are not interconnected. Assume that $\rho(A) < \rho(B,A) < \rho(C,A)$, and the entire graph $A \cup B \cup C$ forms the densest subgraph. Now, we insert an edge within A, leading to $\rho(B,A) < \rho(C,A) < \rho(A)$, and A becomes the new densest subgraph. Given that B and C are not connected to each other, A, B, and C are assigned to different layers in the LDS decomposition, indicating the division of one layer into three layers. As a result, to maintain it, we need to identify all these divided layers, which is clearly costly, because identifying any one layer of the LDS decomposition may need to invoke the parameterized network flow technique once [34].

Second, whenever an edge is inserted or deleted, the outer density must be changed, and therefore the FDNs also need to be updated. In contrast, IDN is a round-up value of FDN, thus it is updated infrequently. Indeed, updating a single edge in a graph certainly changes the FDN, but it typically has a small impact on the round-up values of FDNs (i.e., IDNs). For example, as shown in the Insert algorithm, IDN only needs to change during insertion when $d_v = \overline{r}_v$ and no reversible path can be found, further confirming that IDN changes are infrequent. This result suggests that maintaining the LDS decomposition is often much more difficult than maintaining the density decomposition, further highlighting the superiority of density decomposition.

6 EXPERIMENTS

Different algorithms. For static graphs, we compare our proposed algorithms Flow (Algorithm 3) and Flow++ (Algorithm 4) with three baselines: Path (Algorithm 1) [14], FW [26], and Fista [35]. Here, Flow, Flow++, and Path are implemented by us. FW [26] and Fista [35] are the state-of-the-art algorithms for LDS decomposition. By our results established in Section 3, these algorithms can also be used to compute the density decomposition, thus we also include them as baselines. For dynamic graphs, we implement three proposed algorithms Insert (Algorithm 5), Delete (Algorithm 6), and Delete++ (Algorithm 7). Since there is no algorithm that can maintain density decomposition, we use the Flow++ algorithm for re-computing the density decomposition as a baseline.

All algorithms are implemented using C++ and compiled with the GCC compiler, employing the O3 optimization. All experiments are conducted on a PC operating a Linux system, equipped with a 2.2GHz AMD 3990X 64-Core CPU and 128GB memory.

Datasets. We use 4 medium graphs and 4 large graphs in our experiments which are downloaded from the Network Repository [50] and the Koblenz Network Collection (http://konect.cc/). The detailed information is provided in Table 1.

Table 1: Datasets statistics

Name	Type	n	m	
DBLP	co-authorship network	317,081	1,049,866	
Citeseer	citation network	384,414	1,736,145	
Yahoo	lexical network	653,261	2,931,698	
Skitter	internet	1,694,617	11,094,209	
Weibo	social network	58,655,850	261,321,033	
UKlink	web graph	18,483,187	261,787,258	
Twitter	social network	20,826,113	294,585,816	
Wiki	web graph	13,593,033	334,591,525	

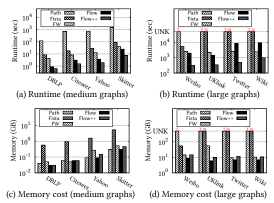


Figure 2: Results of density decomposition on static graphs.

6.1 Performance studies

Exp-1: Runtime of various density decomposition algorithms.

Fig. 2a and Fig. 2b depict the results of different algorithms for computing the density decomposition on medium graphs and large graphs. In Fig. 2b, there are two cases designated as 'UNK' (unknown) for runtime: (1) the algorithm's runtime exceeds our time constraint of 50,000 seconds, labeled as 'T'; (2) the algorithm's memory usage exceeds 128GB, labeled as 'M'. From Fig. 2a and Fig. 2b, Path unsurprisingly exhibits the longest execution time, and even fails to complete the computation within the time constraint on all large graphs. In contrast, our proposed algorithms, Flow and Flow++, not only successfully compute the density decomposition for all datasets but are also at least 1 and 2 orders of magnitude faster than Path respectively, exhibiting significant superiority. For instance, on Citeseer, the algorithms Path, Flow, and Flow++ consume 712.02, 2.5, and 1.56 seconds, respectively, indicating that Flow and Flow++ can be 284x and 456x faster than Path. In addition, Flow++ shows remarkable enhancements, particularly on large graphs, where efficiency is improved by about an order of magnitude compared to Flow. These results confirm our theoretical analysis in Section 2.

When comparing the two LDS decomposition algorithms, we find that FW outperforms Fista over all datasets. The reason could be that Fista was originally designed as an approximate LDS decomposition algorithm, relying on a peeling technique [35], and it may not be well-suited for exact LDS decomposition. Compared to the LDS decomposition algorithms, our best algorithm Flow++ is consistently faster on all datasets, and can still achieve one order of magnitude speedup on large datasets. For instance, on Weibo, the runtime of Flow++ and FW are 190.42 seconds and 2,575.96 seconds respectively, representing a 13x acceleration. This result further demonstrates the high efficiency of our Flow++ algorithm. Exp-2: Memory overheads of different algorithms. The mem-

exp-2: Memory overheads of different algorithms. The memory usages of different algorithms are shown in Fig. 2c and Fig. 2d. Both the Path and Flow algorithms have lower memory costs, which are consistent with the theoretical analysis of the space complexity as O(m). The algorithms Flow++, FW, and Fista all involve a

Table 2: Number of layers of different decompositions "DD" denotes density decomposition. "LDD" represents locally-densest subgraph decomposition.

deno	tes density dec	ompositio	n. "LDD" re	presents locally	-densest s	ubgraph deco	omposition
	Datasets	DD	LDS	Datasets	DD	LDS	
	DBLP	59	1,088	Weibo	168	5,609	
	Citeseer	16	1,435	UKlink	474	40,875	
	Yahoo	26	1,376	Twitter	840	11,949	
	Skitter	02	3 /103	W/iki	602	23 203	

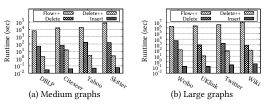


Figure 3: Runtime of various maintenance algorithms

recursive procedure, which requires auxiliary space usage. Among them, Flow++ needs to store each layer R_i , and since the number of layers is small as shown in Table 2, the memory cost of Flow++ remains modest. On all datasets, Flow++ requires no more than 15GB of memory. In the case of FW and Fista, their memory costs depend to a large extent on the effectiveness of the approximation algorithms that they employ. The approximation algorithm in FW is more effective, thus it often consumes less memory than Fista. Nevertheless, on UKlink, FW algorithm uses significantly higher memory compared to Flow and Flow++, indicating lower robustness in terms of memory cost for the FW algorithm. These results confirm that the memory usage of our algorithms is acceptable with efficient computational performance.

Exp-3: Number of layers of different decompositions. The number of layers of density decomposition and LDS decomposition are presented in Table 2. For density decomposition, the number of layers is equal to the pseudoarboricity plus 2, which is typically small in real-world graphs [40]. However, the LDS decomposition yields a considerably larger number of layers, reaching 40,875 layers on UKlink. This multitude of layers often leads to high computation time. Moreover, such a fine partition of layers may inevitably lead to unnecessary separation of tightly connected subgraphs, thus reducing the performance of identifying dense subgraphs for practical applications. These results confirm our analysis in Sections 1 and 3.

Exp-4: Runtime of the various maintenance algorithms. We randomly delete and insert 10,000 edges from each graph dataset to evaluate different maintenance algorithms. The total time required to process these 10,000 deleted or inserted edges is shown in Fig. 3. For the baseline algorithm, since both insertion and deletion require invoking Flow++ to recompute, the runtime for deletion and insertion is the same, thus we only plot one bar in Fig. 3. As can be seen, both Delete and Delete++ are around 3-5 orders of magnitude faster than the baseline algorithm (Flow++), respectively. Furthermore, Delete++ significantly outperforms Delete on all datasets as expected. For example, on UKlink, Delete consumes 943.62 seconds to maintain the density decomposition, while Delete++ only takes 10.98 seconds, achieving an 85x acceleration. These results confirm our analysis in Section 5. For edge insertion, Insert is substantially faster than Flow++, improving by about 5-7 orders of magnitude. When applied to large graphs such as Wiki with hundreds of millions of edges, handling the insertion of 10,000 edges only requires 0.4 seconds. These results further demonstrate the high efficiency of our algorithm. Additionally, it is worth noting that the runtime of our insertion algorithm is often orders of magnitude faster than that of the deletion algorithms. This significant efficiency stems

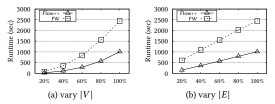


Figure 4: Scalability test results on Wiki.

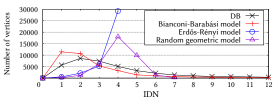


Figure 5: The IDN distribution of DB.

from the fact that (as analyzed in Section 5) updating the IDNs during insertion is much easier compared to the deletion case.

Exp-5: Scalability test. In this experiment, we evaluate the scalability of Flow++ and FW, which are the best algorithms for density decomposition and LDS decomposition respectively. We randomly sample the vertex set V and edge set E to generate 8 subgraphs for each dataset to conduct the scalability test. The results on the Wiki dataset are shown in Fig. 4, and similar results can also be observed on the other datasets. In Fig. 4, we can see that Flow++ outperforms FW at all data scales, consistent with previous observations. Furthermore, the runtime of FW exhibits a rapid increase as the data scale grows, whereas the runtime of Flow++ increases more smoothly. These results demonstrate the high scalability of the Flow++ algorithm.

Exp-6: The IDN distribution. We use a subgraph DB of DBLP, consisting of 37,177 vertices and 131,715 edges, to illustrate the IDN distribution. DB is constructed from the DBLP by extracting authors who have published at least one paper in a conference related to databases and data mining, along with the collaborations between them. For comparison, various synthetic graphs are meticulously generated to match DB's scale in terms of vertices and edges using different models. Specifically, the models include Erdős-Rényi model (ER) [29], power-law Bianconi -Barabási model (BB) [9], random geometric model (RG) [30], power-law Barabási-Albert model (BA) [2], and small-world Watts-Strogatz model (WS) [65]. The IDN distributions of DB and these synthetic graphs are depicted in Fig. 5, and the results on other datasets are consistent. Given that the pseudoarboricity of DB and the BB graph are 19 and 40, we only draw distributions with IDNs no larger than 12 in Fig. 5 for clarity. When IDN exceeds 12, their distributions still exhibit a gradual decline, ultimately converging to zero. In addition, in the BA graph and the WS graph, all vertices share the same IDN. This phenomenon is not explicitly depicted in Fig. 5 and will be elucidated later.

As shown in Fig. 5, similar to the degree distribution, the IDN distribution of DB is highly uneven. The number of vertices is higher when IDN is not greater than 2 but exhibits a declining trend when IDN exceeds 2. This suggests that represented by DB, real-world graphs are typically sparse with small dense regions. For the ER graph, which has a lower pseudoarboricity of 4, most vertices cluster at the top layer of the density decomposition, and its IDN distribution monotonically increases. This differs significantly from real-world graphs that adhere to the power-law principle. In the case of the BB graph, it can be observed that the IDN distribution

closely resembles that of DB. However, its pseudoarboricity is 40, significantly higher than DB's pseudoarboricity 19. Thus, the graph generated by this model can effectively reflect the density structure of real-world graphs, but its densest region could be excessively dense. Regarding the RG graph, the IDN distribution also exhibits a trend of initially increasing and then decreasing. But its pseudoarboricity is only 7, and more than half of the vertices have IDN greater than half of the pseudoarboricity. This indicates that most vertices are located in denser regions within the RG graph, which does not align with real-world graphs. For both the BA graph and WS graph, all vertices have the same IDN of 4. This is because for almost all subgraph $H \subsetneq G$, $\rho(H,G) \approx 4$ holds, resulting in a highly uniform density throughout the graph. These results indicate that the BB graph can relatively better match the density characteristics of real-world graphs.

6.2 Case Studies

Harry Potter characters relationship network. Here we conduct a case study on the Harry Potter characters network¹, denoted by HPC. We evaluate the effectiveness of three graph decompositions, including density decomposition, LDS decomposition, and core decomposition. The IDN, FDN, and core numbers of all characters in HPC are illustrated in Fig. 7, with vertex sizes corresponding to their degrees. From Fig. 6a and Fig. 6b, we can see that the density decomposition and LDS decomposition on HPC exhibit a high similarity, with the only distinction being that in the LDS decomposition, Quirinus Quirrell, one of the four characters with an IDN of 4, is separated from the other three characters. This result indicates that the density decomposition can well approximate the LDS decomposition. Moreover, in this case study, there seems no need to excessively differentiate the characters in fine detail. Whether a character with FDN of 4.00 or 3.33 can both be considered as similar important secondary characters. This result suggests that a moderately coarse-grained partition achieved by density decomposition often suffices, eliminating the need for an excessively fine-grained partition. Additionally, decomposing the graph into finer layers generally requires more computational time. On HPC, invoking Flow++ for density decomposition takes 0.048ms, while performing FW to compute LDS decomposition consumes 0.918ms. In summary, density decomposition can achieve similar results as the LDS decomposition, while significantly reducing computational time and the number of redundant dense subgraph layers.

When comparing the density decomposition (Fig. 6a) and the core decomposition (Fig. 6c), a noticeable disparity emerges. The most notable difference is that the top layer of density decomposition is divided into four layers in the core decomposition. The main character, Harry Potter, is not included in the top layer (i.e., 10-core), which also exhibits a clearly lower density in comparison to the other two density-aware decomposition methods. These results confirm our analysis in Sections 1 and 3.

DBLP collaboration subgraph DB. We also perform a case study on the DB dataset (with 37,177 vertices and 131,715 edges), which is a subgraph of the DBLP co-authorship network that only contains database researchers and their co-authorships. We depict DB's four subgraphs $C_{35} \subseteq R_{18} \subseteq C_{18}$ and R_{19} as shown in Fig. 7a. The full version of the figure, with the author represented by each vertex, is available at https://anonymous.4open.science/r/DD-83C6.

As shown by our proposed Sandwich Theorem (Theorem 4), C_{35} , R_{18} , and C_{18} show a hierarchical structure, with their numbers of

vertices and edges being $|C_{35}|=72$, $|E(C_{35})|=1$, 260, $|R_{18}|=131$, $|E(R_{18})|=2$, 339, $|C_{18}|=529$, and $|E(C_{18})|=7$, 398. Compared to the original graph with |V|=37, 177 $\gg |C_{18}|$ vertices, R_{18} is tightly sandwiched between C_{35} and C_{18} . Furthermore, we illustrate the subgraph R_{19} with the dashed line in Fig. 7a. It can be observed that R_{18} and R_{19} are adjacent layers, but $C_{35} \subseteq R_{18}$ and $C_{35} \nsubseteq R_{19}$, indicating the tightness of $C_i \subseteq R_{\lceil i/2 \rceil}$ in the Sandwich Theorem. Moreover, within DB, C_{35} and R_{19} serve as the top non-empty layer for core decomposition and density decomposition, respectively. It can be seen that C_{35} fails to separate two loosely-connected communities, whereas R_{19} contains a densely-connected community. Their respective densities are $\rho(C_{35})=17.5<\rho(R_{19})=18.3$, and R_{19} is the densest subgraph. This again highlights the weakness of the core decomposition in capturing dense structures, particularly when contrasted with the density-aware decomposition.

To further confirm the unnecessary fine separation generated by LDS decomposition, we present the subgraph R_{14} of DB in Fig. 7b, using a single large vertex to represent R_{15} for clarity. In density decomposition, vertices in $R_{14} \setminus R_{15}$ are grouped into the same layer, while LDS decomposition divides them into three different layers with FDNs of 13.58, 13.50, and 13.28, respectively. These three layers have nearly the same outer density and form a densely connected community, suggesting that they should not be partitioned into three distinct layers. Furthermore, the four vertices with FDNs of 13.50 are connected with each other by only four edges, indicating a loose connection. It is evident that there is no need to separate these four vertices into an individual layer. All these observations suggest that LDS decomposition may overly separate the dense subgraph layers, while density decomposition, which places $R_{14} \setminus R_{15}$ in the same layer, is more reasonable.

Outer density comparison. In this case study, we evaluate the ability of core decomposition, LDS decomposition, and density decomposition to reflect the density of subgraphs. For measuring the density of a specific layer R_i in a decomposition, utilizing the outer density $\rho(R_i, R_{i+1})$ is considered superior to the alternatives $\rho(R_i \setminus R_{i+1})$ or $\rho(R_i)$. This is because: (1) the computation of $\rho(R_i \setminus R_{i+1})$ disregards the contributions of $E_{\times}(R_i, R_{i+1})$, which represents the closeness of the connection between R_i and R_{i+1} . Contrarily, $\rho(R_i, R_{i+1})$ incorporates these edges into the computation; (2) $\rho(R_i)$ considers the entirety of R_i instead of $R_i \setminus R_{i+1}$. However, our goal is to measure the density of R_i while excluding the influence of R_{i+1} . When i is small, $\rho(R_i)$ typically fails to highlight the impact of $R_i \setminus R_{i+1}$ on the density of R_i .

Fig. 8 depicts the outer densities of subgraphs computed through different decompositions on HPC and DBLP. In Fig. 8, each point along the horizontal axis signifies the number of vertices contained in a layer, and the vertical axis denotes the outer density of that layer. For example, the point of C_9 in Fig. 8a represents the 9-core in HPC, and its horizontal and vertical axes are $|C_9|$ and $\rho(C_9, C_{10})$ respectively. From Fig. 8a, we can see that on HPC, the line chart for core decomposition starts from 5 and goes up to 9 initially because of $\rho(C_{10}, \emptyset) = 5$ and $\rho(C_9, C_{10}) = 9$. This increase indicates that the lower layer (i.e., C_9) is even denser than the higher layer (i.e., C_{10}), suggesting that core decomposition does not perform well in extracting high-density subgraphs. In contrast, density decomposition guarantees a monotonically decreasing trend in the outer density of each layer, ensuring that higher layers are denser than lower ones. Similar results are presented for DBLP, as shown in Fig. 8b, where, for clarity, only the layers with fewer than 1,000 vertices are plotted. Clearly, the outer densities of C_{64} and C_{34} are higher than those of C_{65} and C_{35} , again indicating that core decomposition cannot stably obtain denser graphs. For density decomposition and

¹Data source: https://github.com/efekarakus/potter-network, from which we extract a subgraph containing only main characters and main relationships for clarity.

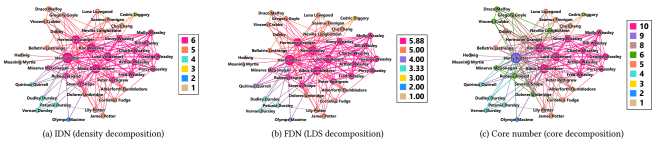


Figure 6: Dense subgraph decompositions of Harry Potter character relationship network HPC.

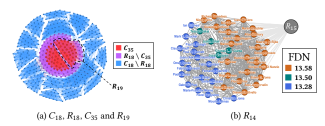


Figure 7: Decompositions of DB network.

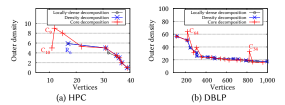


Figure 8: Outer density of different decompositions.

LDS decomposition, as expected, their line charts overlap perfectly. This confirms that density decomposition can well approximate LDS decomposition while significantly reducing the number of redundant layers.

7 RELATED WORK

Densest subgraph discovery. Our work is close to the problem of finding the densest subgraph in a graph. A well-known algorithm for computing the densest subgraph is based on a parameterized network flow technique [34] which needs to invoke the maximum flow algorithm for $O(\log n)$ times. To improve the efficiency, Danisch et al. proposed a convex programming approach to identify the densest subgraph which is considered as the state-of-the-art [26]. Several approximate algorithms have also been devised to compute the densest subgraph, such as the linear-time 2-approximation algorithm [17] O(m+n) and the $(1+\epsilon)$ -approximation iterative algorithm [12, 18], with near-linear time for each iteration. Recently, the techniques for finding the densest subgraph were also extended to the higher-order case [36, 59, 62, 66] as well as the directed graph case [43, 44]. Additionally, there are several approximation algorithms available for maintaining the densest subgraph [8, 47, 56], but none of them can achieve exact maintenance of the densest subgraph. Moreover, all the techniques mentioned above are mainly designed for solving the densest subgraph problem and cannot be directly applied to compute the density decomposition.

Cohesive subgraph decomposition. Except for the density decomposition, there also exist several other cohesive subgraph decompositions that aim to decompose a graph into a set of nested

cohesive subgraphs. Notable examples include the k-core decomposition [4, 41, 52], k-truss decomposition [22, 37, 64], nucleus decomposition [53-55, 58], k-edge connected subgraph decomposition [15, 16], locally densest subgraph (LDS) decomposition [49, 61], distance-generalized core decomposition [11, 25] and colorful h-star k-core decomposition [32, 33]. The core decomposition [4, 41, 52] and their variants [11, 25, 32, 33] is a degree-aware decomposition which iteratively decomposes the graph based on the degrees of the vertices. The truss decomposition is a triangle-aware decomposition that is based on the number of triangles in which an edge participates [22, 37, 64]. The nucleus decomposition can be considered as a higher-order core or truss decomposition which is based on the number of cliques in which a smaller clique participates [53–55, 58]. The k-edge connected subgraph decomposition is a connectivity-aware decomposition that aims to decompose the graph into a set of highly connected subgraphs [15, 16]. The LDS decomposition, however, is a density-aware decomposition that can fully capture the density of the decomposed subgraphs [49, 61]. Note that the density decomposition studied in this paper is also a density-aware decomposition. In this paper, we reveal an interesting inclusion relationship between the density decomposition and LDS decomposition, and we also propose several novel and more efficient algorithms to compute the density decomposition on both large static and dynamic graphs.

8 CONCLUSION

In this paper, we conduct an in-depth investigation of density decomposition on both static and dynamic graphs. First, we establish an inclusion relationship between density decomposition and locally-densest subgraph decomposition, as well as the approximate relationship between density decomposition and core decomposition. For the computation of density decomposition, we develop the Flow++ algorithm using carefully designed network flow and divide-and-conquer techniques, achieving a notable reduction in time complexity to $O(m^{3/2}\log p)$. To handle dynamic graphs, we prove a density decomposition update theorem, which shows that the maintenance of density decomposition is highly localized. Building upon this result, we develop the insertion and deletion algorithms, i.e., Insert, Delete, and Delete++, all with linear time complexity. Extensive experiments on 8 real-world datasets demonstrate the efficiency, scalability, and effectiveness of our solutions.

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