Motif-driven Dense Subgraph Discovery in Directed and Labeled Networks

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ABSTRACT
Dense regions in networks are an indicator of interesting and unusual information. However, most existing methods only consider simple, undirected, unweighted networks. Complex networks in the real-world often have rich information though: edges are asymmetrical and nodes/edges have categorical and numerical attributes. Finding dense subgraphs in such networks in accordance with this rich information is an important problem with many applications. Furthermore, most existing algorithms ignore the higher-order relationships (i.e., motifs) among the nodes. Motifs are shown to be helpful for dense subgraph discovery but their wide spectrum in heterogeneous networks makes it challenging to utilize them effectively. In this work, we propose quark decomposition framework to locate dense subgraphs that are rich with a given motif. We focus on networks with directed edges and categorical attributes on nodes/edges. For a given motif, our framework builds subgraphs, called quarks, in varying quality and with hierarchical relations. Our framework is versatile, efficient, and extendible. We discuss the limitations and practical instantiations of our framework as well as the role confusion problem that needs to be considered in directed networks. We give an extensive evaluation of our framework in directed, signed-directed, and node-labeled networks. We consider various motifs and evaluate the quark decomposition using several real-world networks. Results show that quark decomposition performs better than the state-of-the-art techniques. Our framework is also practical and scalable to networks with up to 101M edges.

KEYWORDS
graph motif, k-core, k-truss, dense subgraph discovery

1 INTRODUCTION
Dense regions in networks contain unusual and interesting information [19]. Dense subgraph discovery is shown to be an effective analysis method in many applications across various domains [5, 13, 17, 18, 26, 30, 33]. It is often a good and cheaper proxy for graph clustering because cohesive subgraphs in real-world networks exhibit good cuts [19, 33]. However, most algorithms to find dense subgraphs are designed for simple, undirected, unweighted graphs. In reality, a common characteristic of natural and engineered systems from various domains is that the nodes (entities) and edges (relationships) have rich information associated with them; i.e., networks are heterogeneous [56]. Relationships can be asymmetrical (one-way) and entities/relationships can be associated with categorical and numerical attributes; e.g., length of a road or gender of a person. Finding the dense subgraphs while actively considering the rich information on nodes/edges has various applications, such as entity resolution [64] and link prediction [11].

Furthermore, most dense subgraph discovery algorithms are designed to capture only the first-order relationships. Higher-order structures (i.e., motifs/graphlets) are shown to be the fundamental building blocks in the organization and dynamics of real-world networks such as social and neural networks [2, 24, 40, 43, 44, 54]. Discovering dense subgraphs that have a higher-order structure is imperative in the analysis of those networks. In simple networks, motifs are used to find subgraphs with higher-order structure, which cannot be detected with edge-centric methods [50, 60]. However, it is not clear how to find dense subgraphs with motifs in heterogeneous networks. The spectrum of motifs in heterogeneous networks is wide due to the edge directions and node/edge attributes. Although this variety is particularly effective for the analysis as the structure and dynamics can vary with respect to the type of motif considered [8, 57, 61], handling the diverse nature of heterogeneous networks while watching for the motifs is a challenging problem.

In this work, we introduce the quark decomposition framework to find motif-driven subgraphs in networks with directed edges and categorical attributes on nodes/edges. Our framework builds subgraphs, called quarks, in varying quality and with hierarchical relations. A k-quark is a motif-parameterized subgraph where each node/edge (or small motif) participates in a number of (larger) motifs. The parameter k denotes the extent of participation and not an input: quark decomposition finds non-empty k-quarks for all k values. Our framework is inspired by the peeling approach in simple graphs, namely core, truss, and nucleus decompositions [12, 50, 52], which first locate the outer sparse parts of the graph and then find the inner dense regions. Given the practicality and effectiveness of the peeling techniques, we adapt them to the directed networks with categorical attributes in a principled way. Note that it is beyond nontrivial to consider a generalization since edge directions and node/edge attributes create a wide and diverse spectrum of motifs (see Figure 1 and Figure 6 for examples).

Quark decomposition takes two motifs as parameters, M and N where M ⊂ N, and builds subgraphs where Ms participate in many
Ns. The parameterized formulation enables the discovery of diverse subgraphs and lets a trade-off between quality and practicality. We assign density indicators for each motif \( M \), called quark numbers, to denote how well \( M \) is connected to its neighborhood, which are then used to build the \( k \)-quarks. Quark numbers also relate the \( k \)-quarks with each other using containment relations; the quarks with larger \( k \) are contained in the ones with smaller \( k \).

We use quark decomposition in two broad classes of applications: (1) When the motif of interest is unknown and higher-order organization of the network is asked; (2) When the motif of interest is known and guides the quark decomposition. For (1), we compare quarks obtained with various motifs and analyze the number and quality of resulting quarks in directed and signed-directed networks. In a word-association network, we show that overlapping quarks by the same motif as well as the quarks by different motifs capture diverse contexts of the words. We also analyze the Florida Bay food web and show that quarks obtain consistently better results than the state-of-the-art algorithm for finding groups of compartments with respect to the ground-truth classifications. For (2), we consider the task of finding gender-balanced communities in networks where genders are used as node labels. We focus on the college friendship networks that have low female ratios. We consider clique-based instantiations of the quark decomposition and choose gender-balanced triangles and four-cliques as the motif \( N \). We observe consistently high female ratios when compared to the label-oblivious state-of-the-art methods.

Key contributions in this paper are summarized as follows:

- **Quark decomposition.** We propose a framework to find dense subgraphs according to a given motif in networks with directed edges and categorical attributes on nodes/edges. Quark decomposition is versatile, efficient, and extendible.
- **Limitations and role confusion problem.** We characterize the limitations and practical instantiations of quark decomposition to guide the selection of parameter motifs \( M \) and \( N \). Presence of multiple orbits in some directed motifs results in subgraphs where a node/edge serves in multiple orbits. We call this role confusion and devise role-aware quark numbers as a remedy.
- **Generic peeling algorithm for any motif.** We introduce a generic peeling algorithm that works for any motif pair \( M, N \) to find the quark decomposition. Our algorithm is similar in spirit to the core/truss/nucleus decompositions and can enjoy the optimizations applicable for the peeling algorithms.
- **Extensive evaluation on real-world networks.** We evaluate quark decomposition on three types of heterogeneous networks: directed, signed-directed, and node-labeled. We consider various motifs using several real-world networks. Results show that quark decomposition outperforms the state-of-the-art techniques and is also practically scalable to networks with up to 101M edges.

2 RELATED WORK

Here we summarize the related works on motif-driven dense subgraphs in heterogeneous networks. Note that there are too many clustering, community detection/search works on heterogeneous networks [15], but our focus is limited to motif-based approaches that find dense subgraphs in directed and labeled networks.

**Peeling approaches on simple networks.** Core decomposition is a simple but effective model to locate the seed regions where dense subgraphs can be found [52]. It makes use of degrees to assign core numbers to nodes. Regularizing the node degrees, that span to a large range, to the core numbers in a smaller range is the key and makes the peeling a fundamental building block in an array of applications [5, 10, 29, 37, 42]. Higher-order variants of the peeling are introduced to take advantage of the triangles and small subgraphs. Truss decomposition leverages triangles [12, 23, 46], nucleus decomposition makes use of small cliques [50, 51], and tip/wing decomposition utilizes butterflies (2,2-bicliques) in bipartite networks [3, 49] to find dense regions in a principled way.

Nucleus decomposition generalizes core and truss decompositions. It finds \( k\)-\( (r,s) \) nucleus, defined as a subgraph where each \( r \)-clique is a part of \( k \) number of \( s \)-cliques where \( r < s \). Those peeling approaches work on undirected simple networks. Here we work on directed networks with attributes on nodes/edges. We compare our methods with nucleus decomposition to highlight the benefit of considering edge directions and node labels.

**Cycle-, flow-trusses.** Regarding the adaptations of core and truss decompositions for directed graphs, Takaguchi and Yoshida [58] introduced cycle- and flow-trusses. Their algorithms work on directed networks with respect to cycle and flow (acyclic) motifs (see Figure 1) and rely on the occurrences of the cycle and flow motifs for each edge. However, they do not consider the bidirectional edges and handle each such edge as two separate unidirectional edges. We compare our work with cycle-flow trusses in Section 5.1.

**Motif-based densest subgraph and clique finding.** There are a few works in the literature that studies the motif-driven densest subgraph problem. For constant size cliques, which can be thought of as motifs in simple undirected graphs, Tsourakakis introduced the \( k \)-clique densest subgraph problem [60] to generalize the classical densest subgraph discovery [20] for \( k \)-cliques (\( k > 2 \)). Analogous to finding the subgraph with the maximum average degree, Tsourakakis proposed to find the subgraphs that have a maximum average triangle (or \( k \)-clique) count per node. More recently, Fang et al. proposed exact and approximate algorithms for the same problem [16] and Hu et al. considered heterogeneous information networks with specific schemas to find maximal motif-cliques [22].

Note that our problem setup is more general and we aim to find multiple subgraphs that are not necessarily perfect cliques but always significantly dense.

**Higher-order motif clustering.** The motif-based graph clustering problem is studied from a spectral perspective in heterogeneous networks. Benson et al. [8] introduced a nice generalized framework for clustering the networks based on the higher-order connectivity patterns. They defined the motif conductance as the ratio of the number of motifs cutting the border between two regions to the number of motif instance endpoints (i.e., nodes) in the subgraph or its complement, whichever is smaller. Since getting the optimal solution for motif conductance is NP-Hard, they proposed an approximate algorithm that finds the near-optimal cluster in a given network. Their method relies on the spectral clustering of motif adjacency matrix whose entry \( i, j \) is the number of motifs where nodes \( i \) and \( j \) co-occur. The set of nodes in the spectral ordering that has the minimum conductance is reported as the optimal higher-order cluster. Recursive bisection method iteratively finds the near-optimal clusters in the complement of the graph and
Induced hypergraph is also known as section hypergraph. **Dense subgraphs.** We call a subgraph dense if it has many motifs (also called motif-based or -driven dense subgraph). Formally, we use average motif degree to quantify the density of a subgraph with respect to a given motif.

**Definition 2.** For a subgraph \( S \) and a motif \( N \), the average motif degree of \( S \) is the number of instances of \( N \) in \( S \) divided by the number of nodes in \( S \).

### 4 Quark Decomposition Framework

We first define the motif hypergraph.

**Definition 3.** Given a graph \( G \) and template motifs \( M \) and \( N \) s.t. \( M \subset N \). Let \( \{ M \} \) and \( \{ N \} \) be the set of instances of \( M \) and \( N \) in \( G \), respectively, and \( f, g \) be bijective functions. Motif hypergraph \( H_G(M, N) = (V_G, E_G) \) is a hypergraph constructed as follows:

- Each instance of \( M \) in \( G \) forms a node \( u \in V_G \); by \( f(M) \) → \( V_G \).
- Each instance of \( N \) in \( G \) forms a hyperedge \( e \in E_G \); by \( g(N) \) → \( E_G \).

Iff \( M \subset N \) in \( G \), then \( f(M) \subset g(N) \) in \( H_G \).

Note that \( H_G(M, N) \) is a \( t \)-uniform hypergraph (i.e., \( s(e) = t \forall e \in E_G \)) where \( t \) is the number of occurrences of \( M \) in \( N \). We also refer to the degree of each node \( u \in H_G \) as the motif degree, denoted by \( d_{H_G}(u) \) (or \( d(u) \) when \( H_G \) is obvious).

We now introduce the notion of \( k \)-quark subgraph.

**Definition 4.** Given a graph \( G \) and template motifs \( M, N \) such that \( M \subset N \), say \( H_G(M, N) \) is the motif hypergraph defined as above.

- For any \( k \in \mathbb{N} \), a \( k \)-quark of \( H_G(M, N) \) is a connected and maximal induced sub-hypergraph \( H|M| \) such that \( \delta_G(S) \geq k \).
- For a node \( u \in H_G(M, N) \) (corresponding to an instance of motif \( M \) in \( G \)), the quark number of \( u \), denoted by \( K(u) \), is the largest value of \( k \) such that \( u \) belongs to a non-empty \( k \)-quark.

We also refer to \( k \)-quark as quark when \( k \) is irrelevant or clear.

**Definition 5.** \( k \)-quarks form a hierarchy by containment.

- Let \( S \) be a \( k \)-quark and \( T \) be a \( k' \)-quark such that \( k' < k \) and \( S \subset T \). \( S \) is the child of \( T \) (and \( T \) is the parent of \( S \)) if there is no \( k \)-quark \( U \) such that \( k' < k < S \subset U \subset T \).
- A \( k \)-quark is leaf (childless) if there is no \( k^+ \)-quark in \( S \) with \( k^+ > k \).
- Maximum quark number of a graph is the largest \( k \) for which there is a non-empty \( k \)-quark.
- Maximum \( k \)-quark is a quark where \( k \) is the maximum quark number in the graph.

Quark decomposition is the process of finding the quark numbers and \( k \)-quarks for a given pair of motifs \( M, N \) in a graph \( G \). Leaf \( k \)-quarks are the locally optimal subgraphs; they are surrounded by less dense quarks (with lower \( k \) values) and hence often contain the most interesting information.

If \( G \) is a simple undirected graph where \( M \) is \( r \)-clique and \( N \) is \( s \)-clique (\( r < s \)), then \( k \)-quark is nothing but a \( k-(r,s) \) nucleus [50, 51]. If \( G \) is a simple undirected bipartite graph where \( M \) is edge and \( N \) is 2-biclique, then \( k \)-quark reduces to be a \( k \)-wing [49]. In the quark decomposition, each \( M \) instance is given a quark number and the \( k \)-quarks along with the hierarchical relationships among them can be constructed accordingly. Note that, \( N \) is the motif of interest for which dense regions are to be found. Motif \( M \)
For instance, motifs, shown in Figure 1, orbits of the nodes are denoted with the same topological connectivity patterns. For directed triangle to the automorphism orbits \[44, 47\]. Automorphism orbit (or just hyperedge in the motif hypergraph becomes one, thus it is not \(M\) for the \(\mathcal{K}_i(u)\) for \(1 \leq i \leq b\). \(\mathcal{K}_i(u)\) is the largest value of \(k\) such that the node \(u\) belongs to a non-empty role-aware \(k\)-quark where its orbit is \(i\).

In a role-aware \(k\)-quark, each \(M\) instance participates in at least \(k\) instances and the orbit of the \(M\) instance in each of those participations is the same (note that different \(M\) instances can have different orbits in the quark). Role-aware quark numbers describe the extent of participation as a particular orbit while the quark number indicates the extent of participation as any orbit. When \(M\) is node and \(N\) is cycle or reciprocal (in Figure 1), there is no role

\[\text{raw text}\]
confusion since there is only one node orbit in each. For all the other directed triangles, there is a role confusion. When $M$ is edge, we always consider the orbits of the two nodes in it. In the context of directed triangles, one important point is the distinction between unidirectional and bidirectional edges. In this work, we consider the bidirectional edge as an atomic entity; i.e., not combination of two unidirectional edges, due to the fact that a symmetrical relation has a different semantic than two asymmetrical ones—in a sense we only consider the induced edges. For instance, $in^+$ and $out^+$ do not create any role confusion when $M$ is edge, because the unidirectional edge between the white and black nodes in $in^+$ (or $out^+$) cannot serve as a bidirectional edge (between two black nodes) in an adjacent $in^+$ (or $out^+$) motif (see Figure 1).

4.1.2 Example. Figure 2 illustrates an example on a toy graph. We choose the vanilla edge as the motif $M$ and acyclic triangle as the motif $N$, shown in the top-left. We denote the node orbits in acyclic with different colors; orange shows the node with two outgoing edges, blue is for the node with two incoming edges, and red denotes the node with one incoming and one outgoing edge. Motif hypergraph, $H$, with respect to $G, M, N$ is shown next. As described in Definition 3, we create a motif for each edge in a vanilla edge (bottom set in $H$) and a hyperedge for each acyclic (top set in $H$). If each acyclic and edge in $H$ is formed by the concatenation of the node ids in it, e.g., $H$ denotes the edge from $1$ to $2$. Each hyperedge is connected to three nodes since there are three edges in an acyclic—making $H$ a 3-uniform hypergraph. The degree of each node in $H$ is called the motif degree, e.g., it is $2$ for edge-4-5. Quark numbers of the edges are denoted by gray ($K = 1$) and black ($K = 2$) in the bottom set of $H$; e.g., quark number of edge-4-5 is $2$. Based on those quark numbers, we construct the quarks in the top-right. Two quarks are created; a 1-quark, corresponding to the entire graph $G$, and a 2-quark with nodes $1, 3, 4$, and $5$. We observe role confusions for the thick edges in those quarks, which in turn implies the role confusions on nodes. Regarding the edges, $1-4$, $3-4$, and $3-5$ have role confusions in both quarks; e.g., in 1-quark, 1-4 connects orange to blue in 1-4-6 acyclic but links orange to red in 1-5-4 acyclic. For the nodes, $3$ and $4$ in both quarks have role confusions, e.g., node $3$ is red in the $1-3-5$ acyclic (and in $1-3-4$) while being orange in the $3-4-5$ acyclic. We give the role-aware quarks and quark numbers of the edges at the bottom. If we construct the role-aware quarks by abiding the orbits, we get three role-aware 1-quarks. Note that the only edges that have multiple non-zero $K_i$ are the ones that have role confusion in the top-right.

4.2 Algorithms

Here we discuss our peeling algorithms, first for quark decomposition (Definition 4) and then for role-aware quarks (Definition 6).

4.2.1 Quark decomposition. Algorithm 1 outlines the quark decomposition. Here we assume the motif $M$ in Definition 4 to be node or edge for simplicity. Note that larger structures can be considered as well. Also, we avoid constructing the actual hypergraph of motifs since it requires enumerating all the $N$s which will have a significant space cost. Instead, we discover those motifs for each node/edge as needed, similar to the space-efficient approach in nucleus decomposition [50]. There are three different phases in the quark decomposition; motif degree counting, peeling to find quark numbers, and constructing the subgraphs and the hierarchy.

Motif degree counting. Line 1 corresponds to counting the occurrences of $N$ instances per each node or edge. There are some studies [39, 43, 66] that leverage certain commonalities among motifs to avoid redundant computations. Adapting those studies for per node/edge motif counting to improve runtime is possible but out of scope for this work. Also, simultaneously counting the motif degrees for multiple motifs would speed up the workflow when the motif of interest is unknown to the user and multiple options need to be investigated. Note that if the label set of the input motif $N$ is smaller than the label set of the input graph $G$, we can filter the graph to only keep the label set of $N$, e.g., if a triangle of female students is $N$ for an undirected graph where genders are the node labels, only the induced graph of female nodes can be considered.

Peeling to find quark numbers. Lines 3 to 10 assigns a quark number for each node/edge. The classical core decomposition implementation [7] makes use of the bucket data structure to keep track of the nodes with the minimum degree. We also use this approach to watch the nodes/edges with the minimum motif degree. Initially, all the nodes/edges are marked as unprocessed (line 2), which will come handy to ensure each $N$ instance is processed only once. In each iteration, an unprocessed node/edge with the minimum motif degree is chosen (line 3). The motif degree of the chosen node/edge is set to be its quark number ($K$) (line 4). Then, the $N$ instances that contain the chosen node/edge are found and processed in lines 5 to 9. In each such instance of $N$, we make sure that the other nodes/edges (the neighbors of the chosen node/edge) are unprocessed (line 6); this is done to ensure that each $N$ instance is examined only once. Then, the neighbor nodes/edges in each $N$ instance are checked (line 7) and their motif degree is decremented if larger than the motif degree of the chosen node/edge (lines 8 and 9). At the end of the iteration, the chosen node/edge is marked as processed (line 10). For any $M, N$ motif pair, two basic procedures are necessary and sufficient to instantiate the quark decomposition;

• Finding the $N$ instances that contain a given $M$ instance (line 5),
• Finding the $M$ instances in a given $N$ instance (lines 6 and 7).

Constructing the subgraphs and the hierarchy. It is also possible to construct the subgraphs and build hierarchy among $k$-quarks during the peeling operation, as noted in magenta lines after lines 6 and 10. Subgraph and hierarchy construction during the peeling process is introduced in [48] and can be adapted to the quark decomposition. In line 6, the $N$ nodes/edges that are in the same $N$ instance with the chosen node/edge are checked. If those nodes/edges are already processed (i.e., assigned a quark number), we can build non-maximal $k$-quarks during the peeling process. At the end, those non-maximal $k$-quarks are converted to the real (maximal) $k$-quarks with a light-weight post-processing that uses the union-find data structure (following line 10).

Theorem 1. Given a graph $G = (V, E)$ and motif $N$, Algorithm 1 finds the quark numbers, $K(u)$, of all $u \in V$ (or $e \in E$).

Proof. We give the proof for the node case without loss of generality (i.e., it is similar for the edge). As noted in Section 3, nodes $u, v \in V$ are neighbors if they participate in a common motif $N$. $K(u) = k$ indicates that there are at least $k$ instances of $N$ which contain $u$ and in each such $N$, $u$ has a neighbor node $v$ s.t. 
We continue the traversal until no such node
with motifs due to the motif-based traversal, and (3)
Time and space complexity. Algorithm 1
(Definition 4), contradiction. □
the role-aware quark decomposition; again, for simplicity, we as-
consider

 computes the number of
its motif degree is equal to
We consider three baselines in our comparisons.
that a node/edge has in N (line 1). In line 2, we count the orbit
degree for each node/edge and for all orbits 1,...,|B|. Orbit degree
d, of a node u (or edge e) is the number of N instances that contain
de such that the orbit of u (e) is (Definition 6). Each orbit degree of a
edge is processed separately. We also keep |B| arrays to keep track of the processed
(u, o) or (e, o) tuples (e is the orbit of node u, or edge e) (line 3). In lines 4-13, we
process all the orbit degrees in non-decreasing order. Role-aware
quark number is assigned for the chosen node/edge (line 3) and we
find the neighbors of the node/edge in each N to adjust their
orbit degrees (lines 6 to 12). At the end, we return role-aware quark
numbers for each node; |K_i(·)| for 1 ≤ i ≤ |B|.

5 EXPERIMENTS
We evaluate our framework on three types of networks and motifs therein; directed (Section 5.1), signed-directed (Section 5.2), and
node-labeled (Section 5.3) networks. We implement quark decom-
positions for various motifs in each type and evaluate the resulting
subgraphs. All experiments are performed on a Linux operating system
(v. 4.12.14-150.52) running on a machine with Intel(R) Xeon(R) CPU E5-2698 v3 processor at 2.30GHz with 64 GB DDR3 1866 MHz
memory. Algorithms are implemented in C++ and compiled using gcc 6.1.0 at the -O2 level. The code is available at http://sariyuce.
com/quark_decomposition.tar. For each network type, we discuss
the set of motifs used and present the results. We compare
quark decomposition to the state-of-the-art methods and highlight
anecdotal examples to stress the contrast between our method and
others. We also present the runtime performance of quark decom-
positions and other state-of-the-art methods.

Baselines. We consider three baselines in our comparisons.
• Motif clustering (MC) [8]. Set of higher-order clustering algo-
rithms by Benson et al. (see Section 2 for details). We consider
three versions; (1) MC-SINGLE: Algorithm that gives a single
subgraph with near-optimal motif conductance. (2) MC-REC-
BISECTION: Recursive bisection algorithm that iteratively finds
multiple clusters (starting with the optimal) until the cluster size
gets too small (less than 10 nodes) or quality degrades too much
(conductance goes above 0.5). (3) MC-k-MEANS: k-means algorithm
that is run on the motif adjacency matrix – number of clusters (k) must be specified for this version.

\begin{algorithm}
\caption{Role-aware Quark Decom. (G (V, E), motif N)}
1 Let B be the set of orbits that a node/edge has in N
2 Compute d_u (orbit deg) \forall orbits o=1,...,|B|, \forall u \in V \lor d_e(e)
3 Mark every triple (u, o) as unprocessed for o=1,...,|B| \lor \langle e, o \rangle
4 for each unprocessed (u, o) \lor (e, o) with min. orbit degree d_u do
5 \quad \mathcal{K}_u(u) \leftarrow d_u(u) \lor \mathcal{K}_u(e) \leftarrow d_e(e)
6 \quad for each motif N s.t. u \in N (or e \in N) do
7 \quad \quad if any node v \in N (or edge f \in E) then continue
8 \quad \quad // can also find subgraphs & hierarchy
9 \quad \quad for each node v (≠ u) \subset N (or edge f (≠ e)) do
10 \quad \quad \quad if d(v) > d(u) (or d(f) > d(e)) then
11 \quad \quad \quad \quad d(v) \leftarrow d(v) - 1 \lor d(f) \leftarrow d(f) - 1
12 \quad \quad \quad Mark u (or e) as processed
13 \quad \quad // Optional post-processing to build the hierarchy
14 return array \mathcal{K}(·)
\end{algorithm}
We also list the number of motifs for each directed triangle (see Figure 1) and maximum quark numbers in each quark decomposition. Largest motif count and quark number for each network are shown in bold (for quark numbers, \( M \) is a vanilla edge, \( N \) is a triangle from Figure 1).

### Table 1: Directed datasets from various domains.

| Domain     | \(|V|\) | \(|E|\) | \(|E_u|\) | \(|E_d|\) | # motifs | maximum quark numbers |
|------------|--------|--------|--------|--------|----------|-----------------------|
| Foodweb    | 128    | 2.1K   | 31     | 2.0K   | cycle    | 70 7.9K 91 80 212 75 0 |
| EAT        | 23.2K  | 325.0K | 20.1K  | 284.8K | cycle+   | 1 8 1 1 1 1 0         |
| email-EuAll| 265.0K | 419.0K | 54.5K  | 310.0K | out+     | 1 5 3 3 2 3 3         |
| cit-HepPh  | 34.5K  | 421.5K | 65     | 420.2K | cycle+   | 1 4 7 4 2 6 11        |
| Slashdot   | 77.4K  | 828.2K | 359.0K | 110.2K | cycle++  | 1 3 6 3 2 3 3         |
| web-ND     | 325.7K | 1.5M   | 379.6K | 710.5K | cycle+   | 15 14 11 2 3 148       |
| amazon     | 403.4K | 944.0K | 1.5M   | 944.0K | cycle++  | 1 5 6 4 2 4 9          |
| wiki-Talk  | 2.4M   | 5.0M   | 361.8K | 4.3M   | cycle+   | 17 12 7 7 6 18 18      |
| soc-pokec  | 1.6M   | 30.6M  | 8.5M   | 14.0M  | cycle+   | 2 25 9 5 3 7 18        |
| liveJournal| 4.8M   | 68.5M  | 25.6M  | 17.2M  | cycle+   | 7 133 98 89 27 65 247  |
| en-wiki    | 4.2M   | 101.3M | 9.4M   | 82.6M  | cycle+   | 7 28 22 24 5 18 29      |

#### Takaguchi and Yoshida (TY) [58].
Cycle-truss and flow-truss algorithms (see Section 2 for details).

#### (r,s) nucleus [50].
Nucleus decomposition to find hierarchical dense subgraphs in undirected networks (see Section 2).

### Metrics

We consider three metrics to measure the quality of the subgraphs. We also show anecdotal examples when feasible.

#### Motif conductance. Edge conductance is adapted for motifs in [8]. Motif conductance of a subgraph is defined as the ratio of the number of motif instances cut (i.e., motifs in the boundary) to the number of motif instance end points in the subgraph (i.e., nodes participating in the motif). The lower values are better.

#### Average motif degree. Conductance metric is known to have a bias toward giving better results for smaller numbers of clusters [4, 35]. As an alternative, we consider the average motif degree in each subgraph, as given in Definition 2. In edge-based clustering literature, the densest subgraph of a graph is defined as the one with the largest average degree [20, 60]. Here we adapt this measure for the motif-based subgraphs and simply consider the number of motifs per node. The higher values are better.

#### Edge density. We also consider the ratio of edges over all possible in a subgraph \([|E|/\binom{|V|}{2})\]. We use this metric in Section 5.3 for undirected networks. The higher values are better.

### 5.1 Directed networks

#### Datasets

We consider several directed networks from various domains in our experiments: Florida Bay food web (Foodweb), word associations (EAT), emails (email-EuAll), citations (cit-HepPh), online social networks (slashdot, soc-pokec, livejournal, wiki-Talk), web networks (web-ND, wiki-Talk), and product co-purchasing relations (amazon). All networks (except EAT [28]) are obtained from SNAP [34]. Table 1 gives several statistics, including the motif counts and maximum quark numbers.

#### Motifs

We instantiate the quark decomposition for directed networks by considering the edge and triangle motifs (Figure 1), corresponding to the \( M \) and \( N \) in Definition 4, respectively. Note that considering edge as motif \( M \) is more advantageous than node. Since the edges are assigned quark numbers, \( k \)-quarks can overlap with each other. Also, role confusion does not happen for \( out+ \) and \( in+ \) as explained in Section 4.1.1. We also incorporate the reciprocity by considering the undirectional and bidirectional edges separately, rather than treating each bidirectional edge as two undirectional edges. This is because directed networks often have a significant percentage of bidirectional edges (also observed in Table 1) and those need to be treated differently, as discussed in [41, 53]. We use the vanilla edge as the motif \( M \) and each of the seven directed triangles (Figure 1) as the motif \( N \).

We first discuss the motif counts and quarks for directed triangles. Then we compare quarks with baselines using the EAT and Foodweb networks. We finish by comparing the runtimes.

#### 5.1.1 Motif counts and subgraphs. Table 1 lists the motif counts and maximum quark numbers for each motif. cycle is often the least common motif. Networks with significant fraction of bidirectional edges tend to contain reciprocal motifs the most. Note that the most frequent motif does not always yield the highest quark number. In particular, reciprocals are often concentrated in a small region, thus yield highly-dense subgraphs. For example, en-wiki has 5.9M reciprocal and 163.3M acyclic motifs but the maximum quark numbers for those motifs are 29 and 26, respectively. This is because the maximum \( k \)-quark has 465 edges in the reciprocal case but has 9461 edges in acyclic. We also observe that the number of quarks are independent of the motif counts. For example, the most prevalent motif in EAT network is, by far, acyclic. However, acyclic yields only 447 \( k \)-quarks while the cycle+ locates 2826 subgraphs. Overall, the abundance of a particular motif does not imply the existence of dense subgraphs containing that motif.

#### 5.1.2 Comparison with previous methods. We compare quark decomposition against the three baselines listed above. We use EAT network, a collection of word association norms where the nodes are English words and an edge \((u,v)\) implies that human subjects consider the word \( u \) when they are shown the word \( u \) as stimulus.

We first compare the quality of subgraphs given by the quark decomposition and MC algorithms [8] using motif conductance and average motif degree metrics. For quark decomposition, we only consider the \( k \)-quarks with at least 10 nodes for MC algorithms, we consider MC-SINGLE and MC-REC-BOISECTION. Note that we aim to compare the ‘best’ clusters reported by the two algorithms and do not intend any comparison with respect to the number of clusters reported. Figure 3 gives the comparison for reciprocal, \( out+ \), acyclic, and \( cycle+ \) motifs (other motifs show similar results and omitted). For each subgraph, we report the size as well as the quality metric; number of motifs per node in the top row and the
motif conductance in the bottom row. MC variants often give large subgraphs, always with very low conductance, as expected. MC-Single (optimal cluster) has more than 1000 nodes for out+, in+, acyclic, and cycle++. For cycle+, however, it has only five nodes. Quarks are often small, most in the range of 10-100 nodes, and have higher conductance scores. For out+ and cycle++, some quarks yield comparable conductance scores with MC-Rec-Bisection. We also observe that a quark for acyclic has a better conductance than MC-Single (which is the near-optimal as shown in [8]). Regarding the average motif degrees (top row), quarks perform significantly better in all the motifs (note that y-axis is in log-scale). By definition of the quark (the connectivity constraint in particular), if the size is $n$, the number of motifs is at least $n - 2$ (for $k = 1$, a single motif is a valid subgraph and can be extended by a new node that creates a new motif, keeping the motif count $n - 2$). This ensures a lower bound, $\frac{n - 2}{n}$ (close to 1), for average motif degree in quarks. In general, quarks tend to be smaller in size when compared to MC results, have consistently higher average motif degrees, and comparable conductance scores for some motifs. Overall, the top-down partitioning approach in the motif clustering is likely to result in larger subgraphs in varying quality whereas the bottom-up computation in quark decomposition yields smaller subgraphs with larger average motif degrees.

Next we compare quark decomposition with TY [58]. For EAT network, TY reports maximum cycle-truss number of 3 and maximum flow-truss number of 10. For each maximum truss subgraph, we checked the quarks that are the most similar. Figure 4 presents the results for cycle-truss, flow-truss, and their corresponding quarks with size and average motif degree information. For cycle- and flow-truss, we calculate the induced cycle and acyclic motif degrees (i.e., bidirectional edges are not included). Next to each quark, we denote the size of its intersection with the truss. Various types of quarks are able to obtain almost all the nodes in those trusses. 70 of 77 nodes in cycle-truss are obtained with 15 quarks and all of the 45 nodes in flow-truss are given in 15 other quarks. This verifies the artificial over-representation of cycle- and flow-trusses due to the non-induced nature. Overall, treating the bidirectional edges as atomic units enables finding diverse subgraphs while correctly capturing the semantics of pairwise relationships.

Lastly, we compare quarks with (2, 3) nucleus decomposition [50], which ignores edge directions. We observe that incorporating the edge directions results in more diverse subgraphs. The number of subgraphs (of any quality) obtained by each quark decomposition is significantly larger than what nucleus decomposition yields. As an anecdotal example, we show a subgraph found by the nucleus decomposition in Figure 5. The direction-oblivious subgraph contains words related to astronomy and space. Quark decompositions capture several diverse contexts related to those words. Thanks to the overlapping quarks, in+ finds two subgraphs that contain space and stars: one in astronomy theme (uranus, venus, ...) and another in the religious context (god, eternity, ...). Out+ yields a subgraph in the air-flight context (sky, aircraft, wing, ...).
We also recognize that multiple meanings of the homonym words are reflected in various k-quarks. For instance, lie is reported in two subgraphs by cycle++: one is about incorrectness (falsehood, untruth, ...) and the other is about staying at rest in the horizontal position (couch, rest, ...). Overall, quark decompositions by various motifs can locate diverse contexts for a given word thanks to the motif-aware approach and overlapping nature of quarks.

5.1.3 Analysis of Florida Bay food web. Here we analyze the structure of Florida Bay food web network (Foodweb) where the nodes are the compartments (i.e., organisms, species) and the edges are the directed carbon exchanges (i.e., $u \rightarrow v$ if $v$ eats $u$). Benson et al. showed that high-quality clusters (i.e., with low conductance) by MC only exist for out+, which implies that the organization of compartments is better described with out+ (as opposed to the common belief that acyclic is the key motif) [8]. They also show that the 4 clusters by MC-K-MEANS for out+ reflect the ground-truth subgroup classifications better than the state-of-the-art clustering algorithms such as spectral edge clustering (with k-means and recursive bisection) [63], InfoMap [45], and Louvain method [9].

We first compare the quarks for out+ with MC-K-MEANS. We find 7 quarks for out+, so we consider MC-K-MEANS with 7 clusters as well as with 4 clusters. The nodes that appear in multiple quarks are only considered to be part of their largest quark (other choices give similar results). Table 2 presents the results for two ground-truth classifications given in [8, 62] by four metrics: Adjusted Rand Index (ARI), F1 score, Normalized Mutual Information (NMI), Purity [38]. Quarks clearly outperform MC-K-MEANS variants in both classifications by all the metrics. One particular difference is that MC-K-MEANS considers some macroinvertebrates like predatory crabs among the benthic predators of eels and toadfish whereas quark decomposition finds all macroinvertebrates in the same subgraph. We believe the main reason is that MC-K-MEANS considers motif counts from the node-perspective while quark decomposition is based on the edges and their motif counts.

We also consider acyclic and use role-aware quark decomposition (Algorithm 2) to determine the roles of the compartments in the resulting quarks. For an acyclic formed by $u \rightarrow v, u \rightarrow w$, and $v \rightarrow w$, we define $u$ as the prey orbit, $v$ as the balancer orbit, and $w$ as the predator orbit. The maximum quark obtained by QuarkDec (Algorithm 1) and RA-QuarkDec (Algorithm 2) is the same and contains 48 compartments. RA-QuarkDec assigns three quark numbers for each edge, corresponding to the three edge orbits in acyclic (each is a union of its nodes’ orbits). We determine the role profile for each node based on the quark numbers of its outgoing and incoming edges. The compartments that are dominantly predator are the birds including predatory ducks, big herons & egrets, greeb, and more. Dominantly prey compartments include clon goby, four types of zooplankton microfauna, and seven macroinvertebrates (including pink and herbivorous shrimps). Lastly, the ones that have the balancer role are the fishes, such as (bay) anchovy, sardines, and mojarra. Note that it is not possible to understand the roles of the compartments by QuarkDec since each edge has a single quark number. For instance, the average quark numbers of incoming and outgoing edges for predatory ducks and code goby are very close, which tells nothing about their roles.

5.1.4 Runtime performance. We measure the runtime for quark decomposition and MC-SINGLE on all directed networks. Table 3 lists the results for large networks. MC-SINGLE (denoted M) gives only one near-optimal cluster and quark decomposition (denoted Q) finds all the quark numbers. Quark decomposition is faster than MC-SINGLE for all motifs in web-ND, amazon, soc-pokec, and liveJournal. For some configurations, such as in+ in liveJournal, we observe up to 10x speedup. For en-wiki, however, MC-SINGLE is fastest for all motifs. Note that MC-Single runtime includes motif adjacency construction and spectral clustering to find one cluster. In order to find more clusters, spectral clustering needs to be run again. However, the spectral clustering takes 36% of the total time for en-wiki (on avg.), hence obtaining 10 clusters will increase the runtime by 4x. All in all, although MC-SINGLE finds only one cluster, quark decomposition is faster for most networks and motifs, and a better choice especially when multiple subgraphs are targeted.

### Table 3: Runtimes for large directed networks (sec). Best result in each motif column is in bold.

| Network | cycle++ | acyclic | out+ | in+ | cycle+ |
|---------|---------|---------|------|-----|--------|
| web-ND | 3.37    | 1.26    | 0.62 | 0.63 | 0.55   |
| amzn    | 7.40    | 1.29    | 2.53 | 1.52 | 1.18   |
| wiki    | 28.91   | 11.62   | 10.91| 16.41| 20.58  |
| soc-p   | 23.62   | 66.91   | 37.01| 119.34| 48.91  |
| live-w  | 57.42   | 180.04  | 118.11| 113.59| 122.54 |
| en-w    | 900.1   | 774.1   | 684.1| 799.1| 677.4  |

### Table 2: Foodweb classification results. Best in each row is in bold.

| Class 1 | Metric     | Quarks (7 subgraphs) | MC-K-MEANS w/ 4 clusters | MC-K-MEANS w/ 7 clusters |
|---------|------------|----------------------|--------------------------|--------------------------|
| ARI     | 0.3627     | 0.3005               | 0.1485                   |                          |
| F1      | 0.4869     | 0.4574               | 0.3794                   |                          |
| NMI     | 0.5415     | 0.5040               | 0.4843                   |                          |
| Purity  | 0.5968     | 0.5645               | 0.5161                   |                          |

| Class 2 | Metric     | Quarks (7 subgraphs) | MC-K-MEANS w/ 4 clusters | MC-K-MEANS w/ 7 clusters |
|---------|------------|----------------------|--------------------------|--------------------------|
| ARI     | 0.3816     | 0.3265               | 0.1871                   |                          |
| F1      | 0.5675     | 0.5380               | 0.4601                   |                          |
| NMI     | 0.5206     | 0.4822               | 0.4309                   |                          |
| Purity  | 0.6452     | 0.6129               | 0.5645                   |                          |
and *reddit-title* consider the positive and negative interactions among users who belong to different subreddits [31] We considered the last interactions in the datasets. We also consider *epinions*, a who-trust-whom social network [21], and *slashdot* which contains the self-tagged friend/foe relationships [32]. All are obtained from SNAP [34]. Table 4 gives the number of positive and negative edges, motif counts, and maximum quark numbers.

**Motifs.** We use edge and triangle motifs, corresponding to the $M$ and $N$ in Definition 4, respectively. This also ensures that quarks can overlap with each other. There is no bidirectional edge and there are twelve possible triangle motifs in total, as shown in Figure 6; four cycle motifs since there is a single orbit and eight acyclic motifs where each $++$ and $--$ appears in three different ways. We use the *vanilla edge* as $M$ and each of the twelve triangles (Figure 6) as $N$.

5.2.1 Motif counts and quark numbers. Acyclic variants are significantly more common than the cycles in all networks. Among the cycle variants, $++$ is the most prevalent in *reddit* networks and $--$ is the least common in all. This is also coherent with the structural balance theory [14], which states that the triangles with an odd number of negative links are rare. However, *cycle++* is more common than the other cycles in the *epinions* network. This might be due to hierarchical status among the nodes; the lower status nodes are likely to trust the ones with higher status but the reverse is not true. The ratio of balanced triangles is 0.8 for *reddit* status nodes are likely to trust the ones with higher status but the might be due to hierarchical status among the nodes; the lower cycles structural balance theory [14], which states that the triangles with $N$ and $M$ in Definition 4, respectively. This also ensures that quarks appear in three different ways. We use the *vanilla edge* as $M$ and each of the twelve triangles (Figure 6) as $N$.

5.2.2 Comparison with MC. We compare the $k$-quarks with the *MC-SINGLE* and *MC-Rec-Bisection* for *acyclic+++* motif in Figure 7. Some quarks are able to obtain very low conductance scores, close to MC results. For the average motif degrees, quarks significantly outperform MC-Rec-Bisection. In particular, one quark has 284 nodes with average motif degree of 185.2.

One of the quarks by *acyclic+++* has 21 subreddits about rappers/singers, such as *kanye* and *kendricklamar*. The ones which praised the others but have not received much praise (the white node in *acyclic* in Figure 1) are *boogaloized*, *runtheflurious*, and *charlieputh*. The last two are a young rapper duo and a new Canadian singer, respectively, with 6.6K and 279 members. On the other hand, the subreddits that got praised by the others but have not reciprocated (the black node in *acyclic*) are *theweeknd*, *frankocoon*, and *kidcudi*. Those are experienced ones (active since 2010, 2005, and 2003) with tens of thousands of members in their subreddits.

5.3 Node-labeled networks

**Datasets.** Here we consider node-labeled undirected networks. We use the Facebook100 dataset that contains the complete Facebook networks of 100 American colleges from a single-day snapshot in September 2005 [59]. Each node has multiple labels, here we only consider the genders of the nodes (there are only two available in the dataset; female and male) and use quark decomposition to find subgraphs that have balanced gender ratios, i.e., close number of females and males. Excluding the female-only institutions, the overall average female ratio is 48.5 and there are 57 networks with less than 50% female. We choose 18 networks with the lowest female ratio (all have < 45%), Table 5 gives a partial list.

**Motifs.** We instantiate the quark decomposition in five ways, where $F/M$ denotes the female/male nodes: (1) $M$ is vanilla edge and $N$ is triangle in the following two forms: $FMM$ and $FMF$; and (2) $M$ is vanilla triangle, $N$ is four-clique in the following three forms: $FMM$, $FFM$, and $FFF$. Also, there is no role confusion for any variant since the graph is undirected and node labels ensure that an edge cannot serve in different roles in its triangles in (1) (likewise for (2)).

5.3.1 Finding gender balanced subgraphs. Algorithmic fairness is one of the most important problems in today’s automated

| Table 4: Signed datasets. $|V|$, $|E|$, $|E_+|$, and $|E_-|$ are the number of nodes, edges, positive, and negative edges. Motif counts (see Figure 6) and corresponding maximum quark numbers are also shown. |
|---|---|---|---|---|---|
| **red-body** | **red-title** | **epinions** | **slashdot** |
| $|V|$ | 34.7K | 52.9K | 125.8K | 74.3K |
| $|E|$ | 110.8K | 205.5K | 581.6K | 420.5K |
| $|E_+|$ | 102.5K | 188.7K | 465.4K | 313.1K |
| $|E_-|$ | 8.3K | 16.8K | 116.3K | 109.2K |

**Figure 6: Signed directed triangle motifs.**

**Figure 7: Comparison of quarks and MC [8] for *acyclic+++* in *reddit-body* network with respect to the number of motifs per node (left); higher is better) and motif conductance (right; lower is better). Subgraphs with at least 10 nodes are shown. Each subgraph is denoted by a point; size is on the x-axis and metric is on the y-axis.**
Table 5: Node-labeled graphs. Statistics and avg. of the female ratios in nuclei and quarks are given. $V_f$ denotes the set of female nodes. (2,3)n and (3,4)n are the nucleus decompositions.

| Network    | $|V|$ | $|E|$ | $V_f$ | edge, triangle, 2,3-nuclei | Quarks | triangle, 4-clique |
|------------|-----|-----|------|-------------------------------|--------|-------------------|
| Rich67     | 3.7K| 81.9K| 25%  | 23.0% 45.0% 50.0% 24.5% 40.0% 45.0% 51.6% | FMM    | FFM              |
| Caltech36  | 769 | 16.7K| 30%  | 39.4% 46.0% 52.0% 38.5% 43.1% 50.2% 52.8% | FMM    | FFM              |
| Carnegie49 | 6.6K| 250.0K| 37%  | 32.6% 49.0% 52.5% 38.5% 43.5% 49.5% 54.9% | FMM    | FFM              |
| MIT8       | 6.4K| 251.3K| 37%  | 38.8% 48.0% 52.1% 42.0% 44.3% 50.3% 53.9% | FMM    | FFM              |
| Stanford3  | 11.6K| 568.3K| 40%  | 46.8% 48.1% 49.0% 44.1% 45.4% 49.2% 55.4% | FMM    | FFM              |
| Cornell5   | 18.7K| 790.8K| 44%  | 44.3% 47.6% 51.8% 45.6% 47.7% 54.9% | FFM    | FFM              |
| Penn94     | 41.6K| 1.8M  | 44%  | 49.7% 48.4% 51.4% 42.1% 44.8% 49.8% 55.8% | FMM    | FFM              |
| UPenn7     | 14.9K| 686.5K| 44%  | 37.3% 48.8% 51.1% 46.4% 45.1% 50.4% 55.4% | FMM    | FFM              |
| Average of 18 networks | 40%  | 42.5% 48.2% 51.5% 44.1% 44.4% 49.7% 54.7% | FMM    | FFM              |

World [27]. Algorithms can amplify the implicit bias in the data, particularly based on the protected attributes like gender, race, ethnicity, and this can lead to unwanted consequences in criminal justice system, hiring, credit scoring, and more [6]. The bias in the network data is more complicated; regarding the gender attribute, for example, the problem is not only the imbalanced gender distribution but also how each gender category is connected to the other categories. There are a few studies that analyze the implications on information diffusion [25, 55]. In this context, the community structure in the network plays an important role and algorithms that do not actively consider the protected attributes are likely to fail getting fair results. Algorithms that can find balanced communities even when there is an imbalance in the input network are essential. Here we use quark decomposition to find subgraphs with balanced gender ratios. As explained above, we set the input motif $N$ in ways to reflect the characteristics of gender balanced subgraphs.

Table 5 gives the female ratios in quarks in comparison to the label-oblivious nucleus decomposition algorithms [50]. For each network, we find the leaf quarks (Definition 5) and nuclei with at least 10 nodes, calculate the female ratio in each quark or nucleus, and then take the average of those ratios. We also show the average ratios across all the 18 datasets at the bottom. The average female ratio of all the networks is 40.3%. (2,3)-nuclei have a bit better number, 42.5%. Quarks for FMM get 48.2% and the ratios are consistently good for all the networks, varying between 45% – 51.3%. This significant jump from (2,3)-nuclei is due to the fact that each edge has to participate in a number of FMM triangles. Quarks for FFM give an even better ratio, 51.5%. (3,4)-nuclei are larger in number and also denser than (2,3)-nuclei. Its average female ratio (44.1%) is also a bit better than (2,3)-nuclei. Quarks for FMM are very close to (3,4)-nuclei but more consistent. FMM achieves 49.7% and FFM gets the best: 54.7%. Note that results get better for the motifs with larger female ratio: $\text{FFFM} < \text{FFM} < \text{FMM} < \text{FF} < \text{FFMM}$.

Quarks cannot provide a good theoretical lower bound for the female ratio since there is no size constraint in the quark definition. For instance, a 1-quark for FFM can possibly be formed by a pair of connected female nodes and male nodes that are connected to both females; the female ratio would be $2/(n + 2)$ in this case. But in practice, quarks with female-dominant motifs yield dense subgraphs with high female ratios. Even with FMM, which implies a 1/3 ratio (smaller than the average female ratio of the datasets), quarks can obtain better results than the label-oblivious (2,3)-nuclei.

Figure 8: Comparison of the female ratios in subgraphs obtained by quark (FFFM) and (3,4)-nucleus decompositions for UPenn7. For each subgraph, size is given on the x-axis, female ratio is shown on the y-axis, and the edge density is color-coded.

We also show all the subgraphs with size, edge density, and female ratio information obtained by quark and nucleus decompositions in UPenn7 network. Figure 8 gives (3,4)-nuclei and quarks for FFM on UPenn7 network. Quarks are consistently gender balanced when compared to the nuclei; no quark with less than 25% female ratio exists. Note that there is a bit degradation in the number and density of the quarks for FFM: 216 subgraphs with 0.88 avg. edge density, compared to the 230 (3,4) nuclei with avg. density 0.94. Given the consistently high female ratios, we believe that this is an affordable loss in quality.

6 DISCUSSION

Quark decomposition offers a principled approach for motif-driven dense subgraph discovery in heterogeneous networks by successfully regularizing the motif degrees to quark numbers. Our evaluation shows that the $k$-quarks can find dense subgraphs according to a given motif. Role-aware variant solves the role confusion problem by creating multiple quark numbers for each motif $M$. Overall, quark decomposition is versatile, efficient, and extendible.

For future work, it would be interesting to investigate the other byproducts of the quark decomposition, such as hierarchy structure. Our initial results show limited success; detailed and meaningful hierarchies are rare for the most motifs. Theoretical and empirical analysis of the impact of the input motifs, $M$, $N$, on the hierarchy structure would be interesting. Also, adapting the quark decomposition for numerical attributes on nodes/edges would be promising.

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