Abstract
Robustness is a critical measure of the resilience of large networked systems, such as transportation and communication networks. Most prior works focus on the global robustness of a given graph at large, e.g., by measuring its overall vulnerability to external attacks or random failures. In this paper, we turn attention to local robustness and pose a novel problem in the lines of subgraph mining: given a large graph, how can we find its most robust local subgraph (RLS)?

We define a robust subgraph as a subset of nodes with high communicability among them, and formulate the RLS-PROBLEM of finding a subgraph of given size with maximum robustness in the host graph. Our formulation is related to the recently proposed general framework for the densest subgraph problem, however differs from it substantially in that besides the number of edges in the subgraph, robustness also concerns with the placement of edges, i.e., the subgraph topology. We show that the RLS-PROBLEM is NP-hard and propose two heuristic algorithms based on top-down and bottom-up search strategies. Further, we present modifications of our algorithms to handle three practical variants of the RLS-PROBLEM. Experiments on synthetic and real-world graphs demonstrate that we find subgraphs with larger robustness than the densest subgraphs even at lower densities, suggesting that the existing approaches are not suitable for the new problem setting.

1 Introduction
Complex networked systems, such as the Internet, road networks, communication networks, the power grid, etc., are a major part of our modern world. The performance and reliable functioning of complex networks depend on their structural robustness, e.g., their ability to retain functionality in the face of damage to parts of the network. Robustness has been studied in many fields including physics, biology, mathematics, and networking. The research areas include quantifying robustness of a network, studying the response of networks to various attack strategies, manipulating a network to improve its overall robustness, and designing optimally robust networks from scratch.

A vast majority of prior work has focused on the global robustness of graphs at large. On the other hand, research on local robustness is limited to a few works, e.g., on finding robust subgraphs with large spectral radius and identifying critical regions. In this paper, we turn attention to local robustness and pose a novel subgraph mining problem: given a large graph, how can we find its most robust local subgraph of a given size?

Our measure of robustness is the natural connectivity which is based on the reachability of the nodes, also phrased as their “communicability.” As we introduced in prior work, it exhibits several desirable properties; e.g., it captures redundancy by quantifying the count and length of alternative/back-up paths between the nodes. As such, robust subgraphs are intuitively sets of nodes with high communicability among each other. From the practical point of view, they may form the cores of larger communities or constitute the central backbones in large networks, maintaining connectivity and communication at large scale.

While the robust subgraph problem has not been studied before, similar problems have been addressed. Probably the most similar to ours is the densest subgraph problem, aiming to find subgraphs with highest average degree or edge density. However, density is different from robustness; while the former concerns with the number of edges in the subgraph, the topology is also of concern for the latter. We offer the following contributions.

- We formulate a new problem of finding the most robust local subgraph (RLS) in a given graph. While in the line of subgraph mining problems, it has not been studied theoretically before.
- We show that RLS-PROBLEM is NP-hard, and further study its heredity and monotonicity properties.
- We propose two fast heuristic algorithms to solve the RLS-PROBLEM for large graphs: a top-down greedy algorithm that iteratively removes nodes, and a bottom-up approach based on the greedy randomized adaptive search procedure (GRASP).
- We introduce three practical variants of the RLS-PROBLEM, and show how to modify our algorithms to address these problem variants.

We extensively evaluate our methods on both synthetic and real-world graphs. As our RLS-PROBLEM is a new one, we compare to three algorithms (one in [8], two in [39]) for the densest subgraph problem. We find subgraphs with...
higher robustness than the densest subgraphs even at lower densities, demonstrating that the existing algorithms are not applicable for the new problem setting (§5).

2 Background and Preliminaries

2.1 Graph Robustness

Robustness is a critical property of large-scale networks, and thus has been studied in physics, mathematics, computer science, and biology. As a result, there exists a diverse set of robustness measures, e.g., mean shortest paths, efficiency, pairwise connectivity, etc. [11].

In this paper, we adopt a spectral measure of robustness called natural connectivity [11], written as

\[ \bar{\lambda}(G) = \log \left( \frac{1}{n}\sum_{i=1}^{n} e^{\lambda_i} \right) , \]

which can be thought of as the “average eigenvalue” of graph \( G \), where \( \lambda_1 \geq \lambda_2 \geq \ldots \geq \lambda_n \) denote a non-increasing ordering of the eigenvalues of its adjacency matrix \( A \).

Among other desirable properties [7], natural connectivity is interpretable; it is directly related to the subgraph centralities (SC) in the graph. The SC(i) of a node \( i \) is known as its communicability [14], and is based on the “weighted” sum of the number of closed walks that it participates in:

\[ S(G) = \sum_{i=1}^{n} SC(i) = \sum_{i=1}^{n} \sum_{k=0}^{\infty} \frac{(A^k)_{ii}}{k!} , \]

where \( (A^k)_{ii} \) is the number of closed walks of length \( k \) of node \( i \). The \( k! \) scaling ensures that the weighted sum does not diverge, and longer walks count less. \( S(G) \) is also referred to as the Estrada index [14] which strongly correlates with the folding degree of proteins [12].

Noting that \( \sum_{i=1}^{n} \lambda_i = \text{trace}(A) = \sum_{i=1}^{n} \lambda_i^k \) and by Taylor series of the exponential function we can write

\[ S(G) = \sum_{k=0}^{\infty} \frac{1}{k!} \sum_{i=1}^{n} (A^k)_{ii} = \sum_{i=1}^{n} \sum_{k=0}^{\infty} \frac{\lambda_i^k}{k!} = \sum_{i=1}^{n} e^{\lambda_i} . \]

As such, natural connectivity is the normalized Estrada index and quantifies the “average communicability” in \( G \).

2.2 Robustness vs. Density

Graph robustness appears to be related to graph density; however as we show here, there exist key distinctions between them.

Firstly, while density directly uses the number of edges \( e \), such as \( \frac{2e(G)}{|V|^2} \) as in average degree [4, 8, 21] or \( \frac{2e(G)}{|V||V|-1} \) as in edge density [28, 39], robustness follows an indirect route; it quantifies the count and length of paths and uses the graph spectrum. Thus, the objectives of robust and dense subgraph mining problems are distinct.

![Figure 1: Example graphs with the same density but different robustness, due to their distinct graph topology.](image)

Figure 1: Example graphs with the same density but different robustness, due to their distinct graph topology.

More notably, density concerns with the number of edges in the graph and not with the topology. On the other hand, for robustness the placement of edges (i.e., topology) is as much, if not more important. In fact, graphs with the same number of nodes and edges but different topologies are indistinguishable from the density point of view (Figure 1).

To illustrate further, we show in Figure 2 the robustness vs. density of example subgraphs, each of size 50, sampled\(^1\) from a real-world email network (§5 Table 1). While the two properties are correlated, subgraphs with the same density can have a range of different robustness. In fact, among the samples, the densest and the most robust subgraphs are distinct, indicating that one does not always imply the other.

3 Robust Local Subgraphs

3.1 Problem Definition

In their inspiring work [39], Tsourakakis et al. recently defined a general framework for subgraph density functions, which is written as

\[ f_\alpha(S) = g(e(S)) - \alpha h(|S|) , \]

where \( S \subseteq V \) is a set of nodes, \( S \neq \emptyset \), \( e(S) \) is the number of edges in the subgraph induced by \( S \), \( \alpha > 0 \), and \( g \) and \( h \) are any two strictly increasing functions.

Under this framework, maximizing the average degree of a subgraph [4, 8, 21] corresponds to \( g(x) = h(x) = \log x \) and \( \alpha = 1 \) such that

\[ f(S) = \log \left( \frac{e(|S|)}{|S|} \right) . \]

In order to define our problem, we can relate the objective of our setting to this general framework. Specifically, our objective can be written as

\[ f(S) = \log \left( \sum_{i=1}^{e(S)} \frac{e(S)}{|S|} \right) , \]

which is to maximize the average eigenvalue of a subgraph. Therefore, the objectives of the two problems are distinct, although they both fall under a more general framework [39].

\(^1\) We create subgraphs by snowball sampling: pick a random node and progressively add its neighbors with probability \( p \), and iterate in a depth-first fashion. Connectivity is guaranteed by adding at least one neighbor of each node. We use varying \( p \in (0, 1) \) to control the tree-likeness, and obtain subgraphs with various densities.
In the following we formally define our robust local subgraph mining problem, which is to find the highest robustness subgraph of a certain size (hence the locality) in a given graph, which we call the RLS-PROBLEM.

**Problem 1. (RLS-PROBLEM)** Given a graph \( G = (V, E) \) and an integer \( s \), find a subgraph with nodes \( S^* \subseteq V \) of size \( |S^*| = s \) such that
\[
 f(S^*) = \log \sum_{i=1}^{n} e^{\lambda_i(S^*)} - \log s \geq f(S), \quad \forall S \subseteq V, |S| = s.
\]

\( S^* \) is referred as the most robust \( s \)-subgraph.

One can interpret a robust subgraph as containing a set of nodes having large communicability within the subgraph.

**Theorem 3.1.** The optimal RLS-PROBLEM is NP-Hard.

**Proof.** See Appendix A.

### 3.2 Problem Properties

Certain characteristics of hard combinatorial problems sometimes guide the development of approximation algorithms for those problems. In this work, we study two such characteristics, namely semi-heredity and subgraph monotonicity, for the RLS-PROBLEM.

Problems that exhibit the (semi-)heredity or monotonicity properties often enjoy algorithms that explore the search space in a smart and efficient way. For example cliques exhibit heredity, i.e., all induced subgraphs are also cliques. This is a key property used in successful algorithms for the maximum clique problem, e.g., checking maximality by inclusion is a trivial task and effective pruning strategies can be employed within a branch-and-bound framework. Other algorithms exploit monotonicity to employ “smart node ordering” strategies to find iteratively improving solutions. Such orderings help starting with a promising node and sequentially adding the next node in the order such that the resulting subgraphs all satisfy some desired criteria, like a minimum density, which enables finding large solutions quickly.

**Theorem 3.2.** Robustness \( \bar{\lambda} \) is not semi-hereditary. That is, a graph with \( \bar{\lambda} = \alpha \) and \( s > 1 \) nodes is not always a strict superset of some graph with \( s - 1 \) nodes and \( \bar{\lambda} \geq \alpha \).

**Theorem 3.3.** Robustness \( \bar{\lambda} \) is not subgraph monotonic.

**Proof.** See Appendix B for definitions and proofs.

Alas, robust subgraphs do not exhibit any of these properties. This suggests that our RLS-PROBLEM is likely harder than the maximum clique and densest subgraph problems as, unlike robust subgraphs, (quasi-)cliques are shown to exhibit heredity, i.e., all induced subgraphs are also cliques.

### 3.3 Problem Variants

In Appendix C we introduce three practical variants of our RLS-PROBLEM: finding (i) the most robust subgraph (no size constraint), (ii) top-\( k \) most robust \( s \)-subgraphs, and (iii) the most robust \( s \)-subgraph including a given set of seed nodes. We also show how to adapt our algorithms for the RLS-PROBLEM to these variants.

### 4 Finding Robust Local Subgraphs

Given the hardness of the RLS-PROBLEM, we design two heuristic solutions. The first is GREEDYRLS, a top-down approach that iteratively removes nodes to obtain a subgraph of desired size. This greedy strategy serves as a simple baseline. Our second and proposed solution GRASP-RLS is a bottom-up randomized approach in which we iteratively add nodes to build up our subgraphs. Both solutions order the nodes by their contributions to the robustness.

#### 4.1 Greedy Top-down Search Approach

This approach iteratively and greedily removes the nodes one by one from the given graph \( G = (V, E), |V| = n, |E| = m \), until a subgraph with the desired size \( s \) is reached. At each iteration, the node whose removal results in the maximum robustness of the residual graph is selected for removal.

The removal of a node involves removing the node itself and the edges attached to it from the graph, where the residual graph becomes \( G[V \setminus \{i\}] \). Let \( i \) denote a node to be removed. Let us then write the updated robustness \( \bar{\lambda} \) as
\[
\bar{\lambda} = \log \left( \frac{1}{n-1} \sum_{j=1}^{n-1} e^{\lambda_j + \Delta \lambda_j} \right).
\]

As such, we are interested in identifying the node that maximizes \( \bar{\lambda} \), or equivalently
\[
\max e^{\lambda_1 + \Delta \lambda_1} + e^{\lambda_2 + \Delta \lambda_2} + \ldots + e^{\lambda_{n-1} + \Delta \lambda_{n-1}}
\]
\[
e^{\lambda_1} (e^{\Delta \lambda_1} + e^{\lambda_2 - \lambda_1} \Delta \lambda_2 + \ldots + e^{(\lambda_{n-1} - \lambda_1)} \Delta \lambda_{n-1})
\]
\[
e^{\lambda_1} (e^{\lambda_1} + c_1 e^{\Delta \lambda_2} + \ldots + c_{n-1} e^{\Delta \lambda_{n-1}})
\]

where \( c_j \)‘s denote \( e^{\lambda_j - \lambda_1} \forall j \geq 2 \) and \( c_j \leq 1 \).

#### 4.1.1 Updating the eigen-pairs

When a node is removed from the graph, its spectrum (i.e., the eigen-pairs \( (\lambda_j, u_j) \)) also changes. Recomputing the eigen-values to compute robustness \( \bar{\lambda} \) every time a node is removed is computationally challenging. Therefore, we employ fast update schemes based on the first order matrix perturbation theory.

Let \( \Delta A \) and \( (\Delta \lambda_j, \Delta u_j) \) denote the change in \( A \) and \( (\lambda_j, u_j) \) respectively, where \( \Delta A \) is symmetric. Suppose after the adjustment \( A \) becomes
\[
\tilde{A} = A + \Delta A
\]

where each eigen-pair \( (\tilde{\lambda}_j, \tilde{u}_j) \) is written as
\[
\tilde{\lambda}_j = \lambda_j + \Delta \lambda_j \quad \text{and} \quad \tilde{u}_j = u_j + \Delta u_j
\]

**Lemma 4.1.** Given a perturbation \( \Delta A \) to a matrix \( A \), its eigenvalues can be updated by
\[
\Delta \lambda_j = u_j \Delta A u_j.
\]

**Proof.** See Appendix D.1

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*Robustness of the residual graph can be lower or higher: \( S(G) \) decreases due to monotonicity, but the denominator also shrinks to \( (s - 1) \) at every step.*
Since updating the eigenvalues involves using the eigenvectors, which also change with node removals, we use the following to update the eigenvectors as well.

**Lemma 4.2.** Given a perturbation $\Delta A$ to a matrix $A$, its eigenvectors can be updated by

$$\Delta u_j = \sum_{i=1}^{n} \left( u_i^T \Delta A u_i \right) \lambda_j - \lambda_i u_i.$$  

**Proof.** See Appendix D.2

### 4.1.2 Node selection for removal

By using Lemma 4.1 we can write the effect of perturbing $A$ with the removal of a node $i$ on the eigenvectors as

$$\Delta \lambda_j = u_i^T \Delta A u_i = -2u_{i,v} \sum_{v \in N(i)} u_{v,j}$$

where $\Delta A(i,v) = \Delta A(v,i) = -1$, for $v \in N(i)$, and 0 elsewhere, and $N(i)$ denotes the set of neighbors of $i$. Thus, at each step we choose the node $i \in V$ that maximizes

$$(4.7) e^{\lambda_i} \left( -2u_{i,v} \sum_{v \in N(i)} u_{v,1} + \ldots + e_{n-1} \lambda_i - 2u_{i,v} \sum_{v \in N(i)} u_{v,n-1} \right)$$

We remark that it is infeasible to compute all the $n$ eigenvalues of a graph with $n$ nodes, for very large $n$. Thanks to the skewed spectrum of real-world graphs [15], we can rely on the observation that only the top few eigenvalues have large magnitudes. This implies that the $c_j$ terms in Eq. (4.3) and also Eq. (4.7) become much smaller for increasing $j$ and can be ignored. Therefore, we use the top $f$ eigenvalues to approximate the robustness of a graph. In the past, the skewed property of the spectrum has also been exploited to approximate triangle counts in large graphs [38].

The outline of the GreedyRLS algorithm, its complexity analysis, and its adaptations for the RLS-PROBLEM variants (3.3) can be found in Appendix E.

### 4.2 Greedy Randomized Adaptive Search Procedure (GRASP) Approach

The top-down approach makes a greedy decision at every step. If the desired subgraphs are small, however, this incurs many greedy decisions, especially on large graphs where the number of greedy steps $(n - s)$ would be excessive. Since the RLS-PROBLEM does not exhibit monotonicity or semi-heredity properties (3.2), taking large number of greedy steps can yield poor performance. Therefore, we propose a bottom-up approach that performs local operations to build up solutions from scratch.

Our local approach is based on a meta-heuristic called GRASP [16] for solving combinatorial optimization problems. A GRASP, or greedy randomized adaptive search procedure, is a multi-start or iterative process, in which each iteration consists of two phases: (i) a construction phase, in which an initial feasible solution is produced, and (ii) a local search phase, in which a better solution with higher objective value in the neighborhood of the constructed solution is sought. The best overall solution becomes the final result.

The pseudo-code in Algorithm 1 shows the general GRASP for maximization, where $T_{\text{max}}$ iterations are done. For maximizing our objective, we use $f : S \rightarrow \mathbb{R} \equiv \lambda$, i.e., the robustness function as given in Eq. (2.1). We next describe the details of our two GRASP phases.

**Algorithm 1** GRASP-RLS

**Input:** Graph $G = (V, E)$, $T_{\text{max}}$, $f(\cdot)$, $\alpha(\cdot)$, integer $s$

**Output:** Subset of nodes $S^* \subseteq V$, $|S^*| = s$

1: $f^* = -\infty$, $S^* = \emptyset$
2: for $z = 1, 2, \ldots, T_{\text{max}}$ do
3:  $S \leftarrow$ GRASP-RLS-CONSTRUCTION($G$, $f(\cdot)$, $\alpha(\cdot)$, $s$)
4:  $S' \leftarrow$ GRASP-RLS-LOCALSEARCH($G$, $f(\cdot)$, $\alpha(\cdot)$, $s$)
5:  if $f(S') > f^*$ then $S^* \leftarrow S$, $f^* = f(S)$
6: end for
7: return $S^*$

#### 4.2.1 Construction

In the construction phase, a feasible seed solution is iteratively constructed, one node at a time. At each iteration, the choice of the next node to be added is determined by ordering all candidate nodes $C$ in a restricted candidate list, called $RCL$, with respect to a greedy function $g : C \rightarrow \mathbb{R}$, and randomly choosing one of the candidates in the list. Candidate set in the first iteration is set to $V$ and in later iterations it contains the nodes in the neighborhood $N(S)$ of the current solution $S$. The size of $RCL$ is determined by a real parameter $\beta \in [0, 1]$, which controls the amount of greediness and randomness. $\beta = 1$ corresponds to a purely greedy construction, while $\beta = 0$ produces a purely random one. Algorithm 2 describes our construction phase.

**Algorithm 2** GRASP-RLS-CONSTRUCTION

**Input:** Graph $G = (V, E)$, $g(\cdot)$, integer $s$

**Output:** Subset of nodes $S \subseteq V$

1: $S \leftarrow \emptyset$, $C \leftarrow V$
2: while $|S| < s$ do
3:  Evaluate $g(v)$ for all $v \in C$
4:  $\bar{c} \leftarrow \max_{v \in C} g(v)$, $c \leftarrow \min_{v \in C} g(v)$
5:  Select $\beta \in [0, 1]$ using a strategy
6:  $RCL \leftarrow \{ v \in C \mid g(v) \geq c + \beta (\bar{c} - c) \}$
7:  Select a vertex $r$ from $RCL$ at random
8:  $S := S \cup \{ r \}$, $C \leftarrow N(S) \setminus S$
9: end while
10: return $S$

**Selecting $g(\cdot)$:** We aim to include locally dense nodes in our seed solutions. Therefore, in the first iteration of the construction we use $g(v) = \frac{t(v)}{d(v)}$, where $t(v)$ denotes the number of local triangles of $v$, and $d(v)$ is its degree. Initially the candidate set $C$ is equal to the node set $V$, thus we approximate the local triangle counts for speed [38]. In later iterations we use $g(v) = \Delta \lambda_v$; the difference in robustness when a candidate node is added to the current subgraph.

**Selecting $\beta$:** Setting $\beta = 1$ is purely greedy and produces
the same seed subgraph in every GRASP iteration. To incorporate randomness while staying close to the greedy best-first selection, we choose \( \beta \in [0.8, 1] \) uniformly at random at every step. This produces high quality solutions in the presence of large variance in constructed solutions [16].

4.2.2 Local Search A solution generated by GRASP-RLS-CONSTRUCTION is a preliminary one and may not necessarily have the best robustness. Thus, it is almost always beneficial to apply a local refinement procedure to each constructed solution. A local search algorithm works in an iterative fashion by successively replacing the current solution with a better one in the neighborhood of the current solution. It terminates when no better solution can be found. We describe our local search phase in Algorithm 3.

As the RLS-PROBLEM asks for a subgraph of size \( s \), the local search takes as input an \( s \)-subgraph generated by construction and searches for a better solution around it by “swapping” nodes in and out. Ultimately it finds a locally optimal subgraph of size upper bounded by \((s+1)\).

Algorithm 3 GRASP-RLS-LOCALSEARCH

**Input:** Graph \( G = (V,E) \), \( S \), integer \( s \)

**Output:** Subset of nodes \( S' \subseteq V, |S'| = s \)

1: \( more \leftarrow \) TRUE, \( S' \leftarrow S \)
2: \( while \ more \ do \)
3: \( if \ \exists v \in S \) such that \( \lambda(S\setminus \{v\}) \geq \lambda(S) \) then
4: \( S := S\setminus \{v\} \) s.t. \( v^* := \max_{\bar{v} \in N(S) \setminus S} \lambda(S\setminus \{\bar{v}\}) \)
5: \( if \ |S| = s \) then \( S' \leftarrow S \) end if
6: \( else \)
7: \( more \leftarrow \) FALSE
8: \( end if \)
9: \( add \leftarrow \) TRUE
10: \( while \ add \ and \ |S| \leq s \ do \)
11: \( if \ \exists \bar{v} \in N(S) \setminus S \) s.t. \( \lambda(S \cup \{\bar{v}\}) > \lambda(S) \) then
12: \( S := S \cup \{v^*\}, v^* := \max_{\bar{v} \in N(S) \setminus S} \lambda(S \cup \{\bar{v}\}) \)
13: \( more \leftarrow \) TRUE
14: \( if \ |S| = s \) then \( S' \leftarrow S \) end if
15: \( else \)
16: \( add \leftarrow \) FALSE
17: \( end if \)
18: \( end while \)
19: \( return \ S' \)

The local search is guaranteed to terminate, as the objective value (i.e., subgraph robustness) improves with every iteration and it is upper-bounded by the robustness of the \((s+1)\)-clique. We provide the complexity analysis and the GRASP-RLS algorithm variants in Appendix [F].

5 Evaluation

We evaluate our methods extensively on numerous synthetic and real-world graphs. Our real graphs, as in Table 1, come from various domains, including biological, email, Internet AS backbone, P2P, collaboration, and the Web.

Our work is in the general lines of subgraph mining, however with a new objective based on robustness. The closest to our setting is the densest subgraph mining. Therefore, we compare our results to dense subgraphs found by Charikar’s algorithm [8] (which we refer to as Charikar), as well as by Tsourakakis et al.’s two algorithms [39] (which we refer to as Greedy and LS for local search following the convention in their work). We remark that the objectives used in those works are distinct; namely, average degree and edge-surplus, respectively, and are also different from ours.

We first evaluate the accuracy of the algorithms against ground truth. To do so, we create synthetic graphs and inject a clique in each graph. Note that a clique is both the densest and the most robust subgraph of a certain size. Therefore, the algorithms are compared on the same grounds.

Table 2 provides precision, recall, and subgraph size \(|S|\) averaged over ten Erdős-Rényi random graphs, with \( n = 3000 \) nodes and \( p = \{0.5, 0.1, 0.008\} \), in which we inject a clique of size 30. \( p \)'s are selected to capture very dense, medium-dense, and sparse graphs. We notice that while all methods perform sufficiently well for sparse graphs with \( p = 0.008 \), accuracy of GRASP-RLS is superior to the competing methods at all densities.

We also compare the algorithms on the Chung-Lu random power-law graphs [9], with \( n = 3000 \) and power-law exponent varying from 2.2 to 3.1 as observed in real graphs (larger exponent implies a sparser graph), in which we inject a clique of 15 nodes. We run the LS algorithm seeded with one of the nodes of the clique as previously done in [39], while GRASP-RLS is not favored by such a selection. Precision and recall curves averaged over ten graphs are given in Figure 3. We note that while the accuracies of all methods improve by the increasing exponent as the task becomes easier.

### Table 1: Real-world graphs. \( \delta \): edge density, \( \lambda \): robustness

| Dataset            | \( n = |V| \) | \( m = |E| \) | \( \delta \) | \( \lambda \) |
|--------------------|--------------|--------------|------------|------------|
| Jazz               | 198          | 2742         | 0.1406     | 34.74      |
| Clegans N.         | 297          | 2148         | 0.0489     | 21.32      |
| Email              | 1133         | 5451         | 0.0085     | 13.74      |
| Oregon-A           | 7352         | 15665        | 0.0005     | 42.29      |
| Oregon-B           | 10860        | 23409        | 0.0004     | 47.54      |
| Oregon-C           | 13947        | 30584        | 0.0003     | 52.10      |
| P2P-GnutellaA      | 6301         | 20777        | 0.0010     | 19.62      |
| P2P-GnutellaB      | 8114         | 26013        | 0.0008     | 19.45      |
| P2P-GnutellaC      | 8717         | 31525        | 0.0008     | 13.35      |
| P2P-GnutellaD      | 8846         | 31839        | 0.0008     | 14.46      |
| P2P-GnutellaE      | 10876        | 39994        | 0.0007     | 7.83       |
| DBLP               | 317080       | 1049866      | 2.09 \times 10^{-5} | 103.18 |
| Web-Google         | 875713       | 4322051      | 1.13 \times 10^{-5} | 99.36   |
Table 2: Precision & Recall (avg.) for our GRASP-RLS, Charikar [8], Greedy & LS [39] on ten ER graphs.

| ER parameters | GRASP-RLS | GreedyRLS | Charikar [8] | Greedy [39] | Local Search [39] |
|---------------|-----------|-----------|--------------|-------------|------------------|
| n  | p  | | | | |
| 3000 | 0.5 | 30 | 0.97 | 30 | 0.02 | 3000 | 0.01 | 1 | 3000 | 0.01 | 1 | 3000 | 0.01 | 1 |
| 3000 | 0.1 | 30 | 1 | 30 | 0.95 | 3000 | 0.01 | 1 | 29.60 | 0.99 | 0.97 | 20.63 | 0.37 | 0.35 |
| 3000 | 0.008 | 30 | 1 | 30 | 0.99 | 30 | 1 | 1 | 30 | 1 | 1 | 28.23 | 0.94 | 0.93 |

Figure 3: Precision & Recall for our GRASP-RLS, Charikar [8], Greedy & Local Search [39] vs. exponent of power-law graphs.

GRASP-RLS remains superior with robust performance at all host graph densities.

Clique sizes are both the densest and the most robust subgraphs, however, it is expected that the algorithms will find different subgraphs in general due to their distinct objectives. To understand their differences, we turn to real world graphs and compare the robust and dense subgraphs based on three main criteria: (a) robustness $\bar{\lambda}$ as in Equ. (2.1), (b) triangle density $t(S)/(S^3/3)$, and (c) edge density $e(S)/(S^2)$.

Table 3 shows results on our largest graphs from each category. Note that the three algorithms we compare to try to find the densest subgraph without a size restriction. Thus, each one obtains a subgraph of a different size. To make the robust subgraphs (R) comparable to the densest subgraphs (DS), we find subgraphs of size $s$ equal to the ones found by Charikar, Greedy, and LS, respectively noted as $s_{Ch}$, $s_{Gr}$, and $s_{LS}$. As such, we compare to the best results achieved by each of the densest subgraph algorithms.

We notice that densest subgraphs found by Greedy and LS are often substantially smaller than those found by Charikar, and also have higher edge density, which is the same observation as in [39]. On the other hand, robust subgraphs have higher robustness than densest subgraphs, even at lower densities. This shows that high density does not always imply high robustness, and vice versa, illustrating the differences in the two problem settings.

Thus far, we also note that GRASP-RLS consistently outperforms GreedyRLS, suggesting that the proposed bottom-up search is superior to the greedy top-down search.

Figure 4 shows the relative difference in robustness of GRASP-RLS subgraphs over again, the best results obtained by Charikar, Greedy, and LS on all of our real graphs. We achieve a wide range of improvements depending on the graph, while the difference is always positive. The improvements with respect to the LS results are the most pronounced.

Comparisons in Table 3 and Figure 4 are for subgraphs at sizes where best results are obtained for each of the three densest subgraph algorithms. Our algorithms, on the other hand, accept a subgraph size input $s$. Thus, we next compare the competing methods at varying output sizes. Charikar and Greedy are both top-down methods, in which the lowest degree node is removed at each step and the best subgraph (best average degree or edge surplus, respectively) is output among all graphs created along the way. We modify these so that we pull out the subgraphs when size $s$ is reached during the course of their run.

Figure 5 shows that our GRASP-RLS produces subgraphs with higher robustness at varying sizes on two example graphs (similar results on others). This also shows that the densest subgraph approaches are not directly applicable to our problem.

Experiments thus far illustrate that we find subgraphs with robustness higher than the densest subgraphs. These are relative results. To show that the subgraphs we find are in fact robust, we next quantify the magnitude of the robustness values we achieve through significance tests.

Local search by [39] finds locally optimal subgraphs, which are not guaranteed to grow to a given size $s$. Thus, we omit comparison to LS subgraphs at varying sizes. Figure 4 shows that improvements over LS subgraphs are already substantially large.
Table 3: Comparison of robust and densest subgraphs. Ch: Charikar [8], Gr: Greedy [39], Ls: Local search [39].

| Data   | Method          | robustness $\lambda[S]$ | triangle density $\Delta[S]$ | edge density $\delta[S]$ |
|--------|-----------------|--------------------------|-----------------------------|--------------------------|
|        | Ch Gr Ls        | Ch Gr Ls                 | Ch Gr Ls                    | Ch Gr Ls                 |
| Email  | DS (271, 12, 13)| 13.58 8.51 4.96          | 0.0009 1.0000 0.2237        | 0.0600 1.0000 0.5897     |
| Oreg-C | DS (386, 22, 4) | 8.81 6.40 0.86           | 9.77E-06 0.0 0.0            | 0.0306 0.4372 0.6667     |
| P2P-E  | DS (240, 105, 18)| 52.15 47.62 10.20        | 0.0266 0.2160 0.4178        | 0.2274 0.4759 0.7254     |

Given a subgraph that GRASP-RLS finds, we bootstrap $B = 1000$ new subgraphs by rewiring its edges at random. We compute an empirical $p$-value for each subgraph by dividing the number of randomly rewired subgraphs that have larger robustness by $B$. The $p$-value essentially captures the probability that we would be able to obtain a subgraph with robustness greater than what we find by chance if we were to create a topology with the same number of nodes and edges at random (note that all such subgraphs would have the same edge density). Thus a low $p$-value implies that, among the same density topologies, the one we find is in fact robust with high probability.

Figure 6 shows that the subgraphs we find on almost all real graphs are significantly robust at 0.05. For cases with large $p$-values, it is possible to obtain higher robustness subgraphs with rewiring. For example, P2P-E is a graph where all the robust subgraphs (also the dense subgraphs) found contain very few or no triangles (see Table 3). Therefore, rewiring edges that short-cut longer cycles they contain help improve their robustness. We remark that large $p$-values indicate that the found subgraphs are not significantly robust, but does not imply our algorithms are unable to find robust subgraphs. That is because the rewired more robust subgraphs do not necessarily exist in the original graph $G$, and it is likely that $G$ does not contain any subgraph with robustness that is statistically significant.

Next we analyze the performance of our GRASP-RLS approach in more detail. Recall that GRASP-RLS-CONSTRUCTION quickly builds a subgraph which GRASP-RLS-LOCALSEARCH uses to improve over to obtain a better result. In Figure 7 we show the robustness of subgraphs obtained after construction and after local search on two example graphs for $s = 50$ and $T_{max} = 300$. We notice that most of the GRASP-RLS iterations find a high robustness subgraph right at construction. In most other cases, local search is able to improve over construction results significantly. In fact, the most robust outcome on Oregon-A (Figure 7 left) is obtained when construction builds a subgraph with robustness around $\lambda = 6$, which the local search improves over $\lambda = 20$.

We next perform several case studies on the DBLP co-authorship network to qualitatively analyze our subgraphs. Here, we use the seeded variant of our problem (Appendix C). Christos Faloutsos is a prolific researcher with various interests. In Figure 8(a), we invoke his interest in databases when used with Rakesh Agrawal as seeds, as Agrawal is an expert in this field. Later in (b), we invoke his interest in data mining when we use Jure Leskovec as the second seed, who is a rising star in the field. Likewise in (c) and (d) we find robust subgraphs around other selected prominent researchers in data mining and databases. In (d) we show how our subgraphs change with varying size. Specifically, we find a clique that the seeds J. Widom and J. Ullman belong to, for $s=10$. The subgraph of $s=15$, while no longer a clique, remains stable in which other researchers like R. Motwani and H. Garcia-Molina are included.
Given the local search characteristics of GRASP-RLS, its complexity is linear in host graph size, as theoretically shown in Appendix F.1. Figure 9 also illustrates the linear scalability w.r.t. input graph size empirically.

Figure 9: Scalability of GRASP-RLS by graph size \( m \) and subgraph size \( s \) (run time avg’ed over 10 runs, bars: 25%-75%).

6 Related Work

Robustness is a critical property of networked systems. Thus, it has been studied extensively in various fields including physics, biology, mathematics, and networking. One of the early studies in measuring graph robustness shows that scale-free graphs are robust to random failures but vulnerable to intentional carefully-planned attacks [11]. This observation has stimulated studies on the response of networks to various attack strategies [6, 7, 10, 13, 23, 36]. Other works look at how to design networks that are optimal with respect to some survivability criteria [17, 20, 31, 29]. A vast body of these works focuses on global robustness of graphs at large.

With respect to research on local robustness, Trajanovski et al. aim to spot critical regions in a graph and the destruction of which would cause the biggest harm to the network [37]. Similar works aim to identify the critical nodes and links of a network [18, 25, 32, 34]. These works try to spot vulnerability points in the network, whereas our objective is somewhat orthogonal: identify robust regions. Closest to ours, Andersen et al. consider spectral radius as an objective criterion and propose algorithms for identifying small robust subgraphs with large spectral radius [2].

While having major distinctions as we illustrated in this work, robust subgraphs are related to dense subgraphs, which have been studied extensively. Finding the largest clique in a graph, well-known to be NP-complete [19], is also shown to be hard to approximate [22].

A relaxation of the clique problem is the densest subgraph problem. Goldberg [21] and Charikar [8] designed exact poly-time and \( \frac{1}{2} \)-approximate linear-time solutions to this problem, respectively, where density is defined as the average degree. This problem is shown to become NP-hard when the size of the subgraph is restricted [5]. Most recently, Tsourakakis et al. [39] also proposed fast heuristic solutions, where they define density as edge surplus; the difference between number of edges and \( \alpha \) fraction of maximum edges, for user-specified constant \( \alpha > 0 \). Likewise, Pei et al. study detecting quasi-cliques in multi-graphs [30]. Other definitions include \( k \)-cores, \( k \)-plexes, and \( k \)-clubs, etc. [26].

Dense subgraph discovery is related to finding clusters in graphs, however with major distinctions. Most importantly, dense subgraph discovery has to do with absolute density where there exists a preset threshold for what is sufficiently dense. On the other hand, graph clustering concerns with relative density measures where density of one region is compared to another. Moreover, not all clustering objectives are based on density and not all types of dense subgraphs can be found by clustering algorithms [26].

In summary, while similarities among them exist, discovery of critical regions, robust subgraphs, cliques, densest subgraphs, and clusters are substantially distinct graph mining problems, for which different algorithms can be applied. To the best of our knowledge, our work is the first to consider identifying robust local subgraphs in large graphs.

7 Conclusion

We introduced the RLS-PROBLEM of finding the most robust local subgraph of a given size in large graphs, as well as its three practical variants. While our work bears similarity to densest subgraph mining, it differs from it in its objective; robustness emphasizes subgraph topology more than edge density. We showed that our problem is NP-hard and that it does not exhibit semi-heredity or subgraph monotonicity properties. We designed two heuristic algorithms based on top-down and bottom-up search strategies, and showed how we can adapt them to address the problem variants.

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*We use nine Oregon graphs with various sizes [4], the largest three of which are listed in Table 4. Running time is averaged over \( T_{\text{ RSA }} \) iterations as one can run each pair of construction followed by local search phases completely in parallel.
We found that our bottom-up strategy provides consistently superior results, scales linearly with input graph size, and finds subgraphs with significant robustness. Experiments on synthetic and real graphs showed that our subgraphs are of higher robustness than densest subgraphs even at lower densities, which illustrates the novelty of our problem setting.

Our research sets off several future directions, including the hardness analysis for the RGS-PROBLEM, exploration of new robustness measures with desirable properties, and the design of efficient algorithms for those new objectives.

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Appendix

A NP-Hardness Proof of RLS-PROBLEM

The decision version of the RLS-PROBLEM is as follows.

**P1.** (robust s-subgraph problem RLS-PROBLEM) Is there a subgraph S in graph G with |S| = s nodes and robustness \( \bar{\lambda}(S) \geq \alpha \) for some \( \alpha \geq 0 \)?

In order to show that P1 is NP-Hard, we reduce from the NP-Complete s-clique problem P2 \([19]\).

**P2.** (s-clique problem CL) Does graph G contain a clique of size s?

**Proof.** (of Theorem \([3.2]\))

It is easy to see that P1 is in NP, since given a graph G we can guess the subgraph with s nodes and compute its robustness in polynomial time.

In the reduction, we convert the instances as follows. An instance of CL is a graph \( G = (V, E) \) and an integer s. We pass \( G, s, \) and \( \alpha = \bar{\lambda}(C_s) \) to RLS-PROBLEM, where \( C_s \) is a clique of size s. We show that a yes instance of CL maps to a yes instance of RLS-PROBLEM, and vice versa.

First assume \( C \) is a yes instance of CL, i.e., there exists a clique of size s in G. Clearly the same is also a yes instance of RLS-PROBLEM as \( \bar{\lambda}(C_s) \geq \alpha \).

Next assume \( S \) is a yes instance of RLS-PROBLEM, thus \( \bar{\lambda}(S) \geq \bar{\lambda}(C_s) \). The proof is by contradiction. Assume \( S \) is a subgraph with s nodes that is not a clique. As such, it should have one or more missing edges from \( C_s \). Let us denote by \( W_k = \text{trace}(A_{C_s}^k) \) the number of closed walks of length \( k \) in \( C_s \). Deleting an edge from \( C_s \), \( W_k \) will certainly not increase, and in some cases (e.g., for \( k = 2 \)) will strictly decrease. As such, any s-subgraph \( S' \) of \( C_s \) with missing edges will have \( \bar{\lambda}(S') < \bar{\lambda}(C_s) \), which is a contradiction to our assumption that \( S \) is a yes instance of the RLS-PROBLEM. Thus, S should be an s-clique and also a yes instance of CL. □

B Properties of Robust Subgraphs

In this section we study two properties of the RLS-PROBLEM, namely semi-heredity (a.k.a. quasi-inheritance) and subgraph monotonicity. Analysis shows that our objective formulation does not exhibit neither of these properties.

B.1 Semi-heredity:

It is easy to identify \( \alpha \)-robust graphs (i.e., \( \lambda = \alpha \)) that contain subsets of nodes that induce subgraphs with robustness less than \( \alpha \). As such, robust subgraphs do not display heredity. Here, we study a weaker version of heredity called semi-heredity or quasi-inheritance.

**Definition 1. (Semi-heredity)** Given any graph \( G = (V, E) \) satisfying a property \( p \), if there exists some \( v \in V \) such that \( G - v \equiv G[V \setminus \{v\}] \) also has property \( p \), \( p \) is called a semi-hereditary property.

**Proof.** (of Theorem \([3.2]\))

The proof is by counter example. In particular, robustness of cliques is not semi-hereditary. Without loss of generality, let \( C_k \) be a k-clique. Then, \( \bar{\lambda}(C_k) = \ln \left( \frac{1}{k} (e^{k-1} + (k - 1)^{1/2}) \right) \). Any subgraph of \( C_k \) with \( k - 1 \) nodes is also a clique having strictly lower robustness, for \( k > 1 \), i.e.,

\[
\frac{1}{k-1} \left( e^{k-2} + (k - 2)^{1/2} \right) \leq \frac{1}{k} \left( e^{k-1} + (k - 1)^{1/2} \right)
\]

\[
ke^{k-2} + \frac{k(k-2)}{e} \leq (k-1)e^{k-1} + \frac{(k-1)^2}{e}
\]

\[
ke^{k-1} + k^2 - 2k \leq (k-1)e^k + (k^2 - 2k + 1)
\]

\[
ke^{k-1} \leq (k-1)e^k + 1
\]

where the inequality is sharp only for \( k = 1 \). Thus, for \( \alpha = \bar{\lambda}(C_k) \), there exists no \( v \) such that \( C_k - v \) is at least \( \alpha \)-robust. □

B.2 Subgraph monotonicity:

As we defined in \([2.1]\) our robustness measure can be written in terms of subgraph centrality as \( \bar{\lambda}(G) = \log \left( \frac{1}{n} S(G) \right) \).

As \( S(G) \) is the total number of weighted closed walks, \( \bar{\lambda} \) is strictly monotonic with respect to edge additions and deletions. However, monotonicity is not directly obvious for node modifications due to the \( \frac{1}{n} \) factor in the definition, which changes with the graph size.

**Definition 2. (Subgraph monotonicity)** An objective function (in our case, robustness) \( R \) is subgraph monotonic if for any subgraph \( g = (V', E') \) of \( G = (V, E) \), \( V' \subseteq V \) and \( E' \subseteq E \), \( R(g) \leq R(G) \).

**Proof.** (of Theorem \([3.3]\))

Assume we start with any graph \( G \) with robustness \( \bar{\lambda}(G) \). Next, we want to find a graph \( S \) with as large robustness as \( \bar{\lambda}(G) \) but that contains the minimum possible number of nodes \( V_{\min} \). Such a smallest subgraph is in fact a clique, with the largest eigenvalue \( (V_{\min} - 1) \) and the rest of the \( (V_{\min} - 1) \) eigenvalues equal to \(-1\). To obtain the exact value of \( V_{\min} \), we need to solve the following

\[
\bar{\lambda}(G) = \log \left( \frac{1}{V_{\min}} - e^{V_{\min} - 1} + (V_{\min} - 1)e^{-1} \right)
\]

which, however, is a transcendental equation and is often solved using numerical methods. To obtain a solution quickly, we calculate a linear regression over \( (\bar{\lambda}(G), V_{\min}) \) samples. We show a sample simulation in Figure \([10]\) for \( V_{\min} \) 1 to 100 where the regression equation is

\[\text{Any subgraph } g(C) \text{ of a } k \text{-clique } C \text{ has strictly lower robustness } \bar{\lambda}. \text{ This is true when } g(C) \text{ also contains } k \text{ nodes, due to monotonicity of } S(G) \text{ to edge removals (see Appendix A). Any smaller clique has strictly lower robustness, see proof for semi-heredity.}\]
In this section, we describe three practical variants of the original RLS-P problem.

**Problem 2.** (RGS-P) Given a graph $G = (V, E)$, find a subgraph $S^* \subseteq V$ such that

$$f(S^*) = \max_{S \subseteq V} f(S).$$

$S^*$ is referred as the most robust subgraph.

Another variant involves finding the top $k$ most robust $s$-subgraphs in a graph, which we call the kRLS-P problem.

**Problem 3.** (kRLS-P) Given a graph $G = (V, E)$, and two integers $s$ and $k$, find $k$ subgraphs $S = S_1^*, \ldots, S_k^*$, each of size $|S_i^*| = s$, $1 \leq i \leq k$ such that

$$f(S_1^*) \geq f(S_2^*) \geq \ldots \geq f(S_k^*) \geq f(S), \ \forall S \subseteq V, |S| = s.$$

$S$ is referred as the top-$k$ most robust $s$-subgraphs.

In the third and final variant, the goal is to find the most robust $s$-subgraph that contains a set of user-given seed nodes, called the Seeded-RLS-P problem.

**Problem 4.** (Seeded-RLS-P) Given a graph $G = (V, E)$, an integer $s$, and a set of seed nodes $U$, $|U| \leq s$, find a subgraph $U \subseteq S^* \subseteq V$ of size $|S^*| = s$ such that

$$f(S^*) = \max_{U \subseteq S^* \subseteq V, |S^*| = s} f(S).$$

$S^*$ is referred as the most robust seeded $s$-subgraph.

It is easy to see that when $k = 1$ and $U = \emptyset$, the kRLS-P, the RGS-P and the Seeded-RLS-P problem respectively reduce to the RLS-P problem, and thus can easily be shown to be also NP-hard. A formal proof of hardness for the RGS-P problem, however, is nontrivial and remains an interesting open problem for future research.

### D Updating the Eigen-pairs

#### D.1 Updating the Eigen-values

**Proof.** (of Lemma 4.1)

Using the relation $A\mathbf{u}_j = \lambda_j \mathbf{u}_j$ between the eigenvalues and eigenvectors, we can write the updated relation as

$$(A + \Delta A)(\mathbf{u}_j + \Delta \mathbf{u}_j) = (\lambda_j + \Delta \lambda_j)(\mathbf{u}_j + \Delta \mathbf{u}_j)$$

Expanding the above, we get

$$A\mathbf{u}_j + \Delta A\mathbf{u}_j + A\Delta \mathbf{u}_j + \Delta A\Delta \mathbf{u}_j = \lambda_j \mathbf{u}_j + \Delta \lambda_j \mathbf{u}_j + \lambda_j \Delta \mathbf{u}_j + \Delta \lambda_j \Delta \mathbf{u}_j$$

(4.8)

By concentrating on first-order approximation, we assume that all high-order perturbation terms are negligible, including $A\Delta \mathbf{u}_j$ and $\Delta \lambda_j \mathbf{u}_j$. Further, by using the fact that $A\mathbf{u}_j = \lambda_j \mathbf{u}_j$ (i.e., canceling these terms) we obtain

(4.9)

Next we multiply both sides by $\mathbf{u}_j'$ and by symmetry of $A$ and orthonormal property of its eigenvectors we get Eq. (4.4), which concludes the proof.

#### D.2 Updating the Eigen-vectors

**Proof.** (of Lemma 4.2)

Using the orthogonality property of the eigenvectors, we can write the change $\Delta \mathbf{u}_j$ of eigenvector $\mathbf{u}_j$ as a linear combination of the original eigenvectors:
\[ (4.10) \quad \Delta u_j = \sum_{i=1}^{n} \alpha_{ij} u_i \]

where \( \alpha_{ij} \)'s are small constants that we aim to determine. Using Equ. (4.10) in Equ. (4.9) we obtain
\[ \Delta A u_j + A \sum_{i=1}^{n} \alpha_{ij} u_i = \Delta \lambda_j u_j + \lambda_j \sum_{i=1}^{n} \alpha_{ij} u_i \]
which is equivalent to
\[ \Delta A u_j + \sum_{i=1}^{n} \lambda_i \alpha_{ij} u_i = \Delta \lambda_j u_j + \lambda_j \sum_{i=1}^{n} \alpha_{ij} u_i \]
Multiplying both sides of the above by \( u_k \)'s, \( k \neq j \), we get
\[ u_k' \Delta A u_j + \lambda_k \alpha_{kj} = \lambda_j \alpha_{kj} \]
Therefore,
\[ (4.11) \quad \alpha_{kj} = \frac{u_k' \Delta A u_j}{\lambda_j - \lambda_k} \]
for \( k \neq j \). To obtain \( \alpha_{jj} \) we use the following derivation.
\[ u_j' u_j = 1 \Rightarrow (u_j + \Delta u_j)'(u_j + \Delta u_j) = 1 \]
\[ \Rightarrow 1 + 2u_j' \Delta u_j + \| \Delta u_j \|^2 = 1 \]
After we discard the high-order term, and substitute \( \Delta u_j \) with Equ. (4.10) we get
\[ 1 + 2\alpha_{jj} + \sum_{i=1}^{n} \alpha_{ij}^2 = 1 \]
We note that for a slightly better approximation, one can choose not to ignore the high-order term which is equal to
\[ \| \Delta u_j \|^2 = \sum_{i=1}^{n} \alpha_{ij}^2. \]
Thus, one can compute \( \alpha_{jj} \) as
\[ 1 + 2\alpha_{jj} + \sum_{i=1}^{n} \alpha_{ij}^2 = 1 \]
\[ \Rightarrow 1 + 2\alpha_{jj} + \sum_{i=1,i \neq j}^{n} \alpha_{ij}^2 = 1 \]
\[ \Rightarrow (1 + \alpha_{jj})^2 + \sum_{i=1,i \neq j}^{n} \alpha_{ij}^2 = 1 \]
\[ \Rightarrow \alpha_{jj} = \sqrt{1 - \sum_{i=1,i \neq j}^{n} \alpha_{ij}^2} - 1 \]
Using the \( \alpha_{ij} \)'s as given by Equ. (4.11) and \( \alpha_{jj} = 0 \), we can see that \( \Delta u_j \) in Equ. (4.10) is equal to Equ. (4.5).

E Greedy Top-down Search for RLS-PROBLEM

E.1 GreedyRLS Algorithm See pseudo-code in Algorithm [E]

Algorithm GreedyRLS

Input: Graph \( G = (V, E) \), its adj. matrix \( A \), integer \( s \)
Output: Subset of nodes \( S^* \subseteq V \), \( |S^*| = s \)
1: Compute top \( t \) eigen-pairs \((\lambda_j, u_j)\) of \( A \), \( 1 \leq j \leq t \)
2: \( S_n \leftarrow V \); \( \lambda(S_n) = \bar{\lambda}(G) \)
3: for \( z = n \) down to \( s + 1 \) do
4: Select node \( i \) out of \( \forall i \in S_z \) that maximizes Equ. (4.7) for top \( t \) eigen-pairs of \( G[S_z] \), i.e.
\[ f = \max_{i \in S_z} c_1 \left( e^{-2u_{i1}} \sum_{v \in N(i)} u_{v1} + \ldots + c_te^{-2u_{it}} \sum_{v \in N(i)} u_{vt} \right) \]
where \( c_1 = e^{-\lambda_1} \) and \( c_j = e^{-\lambda_j - \lambda_1} \) for \( 2 \leq j \leq t \)
5: \( S_{z-1} := S_z \setminus \{i\}; \bar{\lambda}(S_{z-1}) = \log \frac{f_{z-1}}{f_z} \)
6: Update \( A; A(i,i) = 0 \) and \( A(i,i) = 0 \)
7: if \( z = \frac{t}{2}, \frac{t}{4}, \frac{t}{8}, \ldots \) then
8: Compute top \( t \) eigen-pairs \((\lambda_j, u_j)\) of \( A \), \( 1 \leq j \leq t \)
9: else
10: Update top \( t \) eigenvalues of \( A \) by Equ. (4.4)
11: Update top \( t \) eigenvectors of \( A \) by Equ. (4.5)
12: end if
13: end for
14: return \( S^* \leftarrow S_{z=s} \)

E.2 Complexity analysis Algorithm [E] has three main components: (a) computing top \( t \) eigenpairs (L1): \( O(nt + mt + nt^2) \), (b) computing Equ. (4.7) scores for all nodes using top \( t \) eigen-pairs (L4): \( O(mt) \sum_i \sum_{d=t} \sum_i = 2mt \), and (c) updating \( t \) eigenvalues: \( O(d) \) & updating \( t \) eigenvectors: \( O(nt^2) \) (using Equ. (4.6) & updating) when a node is removed (L10 & L11, respectively).
Step (a) is executed only once, whereas (b) and (c) are performed at every step for \( n - s \) i.e., \( O(n) \) steps in general for small constant \( s \). Performing (b) for all nodes at every iteration thus takes \( O(mtnt) \). Moreover, performing (c) iteratively for all nodes requires \( \sum_i \sum_{d=t} \sum_i = 2mt \), i.e., \( O(mt) \) for eigenvalues and \( \sum_i \sum_{i=1}^{n} i, i^2, O(t^2n^2) \) for eigenvectors. Therefore, the overall complexity becomes \( O(\max(mtnt, t^2n^2)) \).

As we no longer would have small perturbations to the adjacency matrix over many iterations, updating the eigenpairs at all steps would yield had approximations. As such, we recompute the eigen-pairs at every \( \frac{n}{2}, \frac{n}{4}, \frac{n}{8}, \ldots \) steps. Performing recomputes less frequently in early iterations is reasonable, as early nodes are likely the peripheral ones that do not affect the eigen-pairs much, for which updates would suffice. When perturbations accumulate over iterations and especially when we get closer to the solution, it becomes
beneficial to recompute the eigen-pairs more frequently.

In fact, in a greedier version one can drop the eigen-pair updates (L10-11), so as to only perform $O(\log n)$ recomputes (L8), in which case the complexity becomes $O(\max(tm \log n, t^2n \log n))$.

E.3 Algorithm Variants To adapt our GreedyRLS algorithm for the $k$RLS-PROBLEM, we can find one subgraph at a time, remove all its nodes from the graph, and continue until we find $k$ subgraphs or end up with an empty graph. This way we generate node-disjoint robust subgraphs. One can also create edge-disjoint subgraphs if instead of removing the nodes of each found subgraph, one removes the edges among the nodes of the subgraph.

For the Seeded-RLS-PROBLEM, we can condition to never remove nodes $u \in U$ that belong to the seed set.

GreedyRLS algorithm is particularly suitable for the RGS-PROBLEM, where we can iterate for $z = n, \ldots, V_{\min}$ record the robustness $\lambda(S_z)$ for each subgraph at each step (Alg. L5), and return the subgraph with the maximum robustness among all the $S_z$’s.

F Greedy Randomized Adaptive Search Procedure (GRASP) for RLS-PROBLEM

F.1 Complexity analysis The size of subgraphs $|S|$ obtained during local search is $O(s)$. Computing their top $t$ eigen-pairs takes $O(s^2t + st^2)$, where we use $e(|S|) = O(s^2)$ as robust subgraphs are often dense. To find the best improving node (L12), all nodes in the neighborhood $N(S) \setminus S$ are evaluated, with worst-case size $O(n)$. As such, each expansion costs $O(ns^2t + nst^2)$. With deletions incorporated (L3-4), the number of expansions can be arbitrarily large [24], however assuming $O(s)$ expansions are done, overall complexity becomes $O(ns^3t + ns^2t^2)$. If all $t = |S|$ eigen-pairs are computed, the complexity is quadruple in $s$ and linear in $n$, which is feasible for small $s$. Otherwise, we exploit eigen-pair updates as in GreedyRLS to reduce computation.

F.2 Algorithm Variants Adapting GRASP-RLS for the $k$RLS-PROBLEM can be easily done by returning the best $k$ (out of $T_{\max} \geq k$) distinct subgraphs computed during the GRASP iterations in Algorithm 1. These subgraphs are likely to overlap, although one can incorporate constraints as to the extent of allowed overlap.

For the Seeded-RLS-PROBLEM, we can initialize set $S$ with the seed nodes $U$ in construction (Alg. L1) while, during the local search phase, we never discard a node $u \in U$ from $S$ (Alg. L4).

Finally, for the RGS-PROBLEM, we can waive the size constraint in the expansion step of local search (Alg. L10).

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$V_{\min}$ denotes the minimum number of nodes a clique $C$ with robustness at least as large as the full graph’s robustness would contain. Any subgraph of the clique $C$ has lower robustness (see Appendix B) and hence would not qualify as the most robust subgraph.